

Premature Ejaculation

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Guideline on the Pharmacologic Management of Premature Ejaculation

Acknowledgements and Disclaimers: AUA Guideline on the Management of Premature Ejaculation (PE)

This document was written by the Erectile Dysfunction Guideline Update Panel of the American Urological Association Education and Research, Inc., which was created in 1999. The Practice Guidelines Committee (PGC) of the AUA selected the committee chairs. Panel members were selected by the chairs. Membership of the committee included urologists with specific expertise on this disorder. The mission of the committee was to develop recommendations, that are analysis-based or consensus-based, depending on panel processes and available data, for optimal clinical practices in the diagnosis and treatment of premature ejaculation. This document was submitted for peer review to 57 urologists and other health care professionals. After the final revisions were made, based upon the peer review process, the document was submitted to, and approved by the PGC and the Board of Directors of the AUA. Funding of the committee was provided by the AUA. Committee members received no remuneration for their work. Each member of the committee provided a conflict of interest disclosure to the AUA.

This report is intended to provide medical practitioners with a consensus of principles and strategies for the treatment of premature ejaculation. The report is based on current professional literature, clinical experience and expert opinion. It does not establish a fixed set of rules or define the legal standard of care and it does not pre-empt physician judgment in individual cases. The medical therapies currently employed in the management of PE have not been approved by the U.S. Food and Drug Administration (FDA) for this specific indication. Thus, doses and dosing regimens may deviate from that employed for FDA-approved indications, and this difference should be considered in the risk-versus-benefit assessment. Physician judgment must take into account variations in resources and in patient needs and preferences.

I. Introduction

The three major forms of male sexual dysfunction are ejaculatory dysfunction, erectile dysfunction (ED), and decreased libido (hypoactive sexual desire disorder). While survey findings vary considerably, most epidemiological studies suggest that premature ejaculation (PE) (Although the terms *early ejaculation* and *rapid ejaculation* recently have been suggested as more accurate descriptions of this disorder, to prevent confusion, the common name *premature ejaculation* will be used throughout this document.) may be the most common male sexual disorder. Data from the National Health and Social Life Survey have revealed a prevalence of 21% in men ages 18 to 59 in the United States¹. Using various definitions, other studies report prevalences ranging from less than 5% 2 to greater than 30% 3,4,5.

A universally accepted definition of PE has yet to be established. Masters and Johnson (1970) ⁶ proposed one of the earliest definitions that focused on the inability to delay ejaculation long enough for the woman to achieve orgasm fifty percent of the time, assuming that PE is the sole cause of the female anorgasmia. Kaplan (1974) ⁷ first suggested that PE is primarily a problem of voluntary control over timing of ejaculation, a concept on which the current definition is based. The American Psychiatric Association's Diagnostic and Statistical Manual of Mental Disorders (4th ed., Text Revision) (DSM-IV TR) (2000) ⁸ defines PE with an added emphasis on the emotional and interpersonal impact of ejaculation that occurs earlier than the male desires. Premature ejaculation has been subclassified into two forms: a primary (lifelong) form that begins when a male first becomes sexually active and a secondary (acquired) form ^{9, 10}. The present guidelines and recommendations are based on the following definition, which assumes the absence of partner sexual dysfunction:

Premature ejaculation is ejaculation that occurs sooner than desired, either before or shortly after penetration, causing distress to either one or both partners.

The exact etiology of PE is unknown. Psychological/behavioristic and biogenic etiologies have been proposed. Consequently, the treatment of PE has encompassed psychological, behavioral, and pharmacologic interventions. Current treatments are largely based upon logical solutions (decreasing sensory input), behavior modification therapies, and observations of drug side effects (those with serotonin reuptake inhibiting activity). This guideline will address only pharmacologic therapies, as other therapies are not routinely prescribed by our target audience.

To facilitate informed treatment decisions by physicians and their patients, recommendations on the use of medications currently available in the United States are provided. The majority of the recommendations contained herein are based on a consensus of expert opinion following review of the literature. In some cases, expert consensus is supplemented with a focused review of the limited data. This guideline does not preempt physician judgment in individual cases.

Variations in patient subpopulations, physician experience, and available resources necessarily will influence choice of clinical strategy. Adherence to the recommendations presented in this document cannot assure a successful treatment outcome.

For ease of review, the recommendations are bolded and followed by supporting text. The evidence supporting the recommendations is summarized in Appendices 1 to 3.

II. Methods

The Erectile Dysfunction Guideline Update Panel (hereafter the Panel) of the American Urological Association (AUA) was convened in April 2000 at the request of the AUA Board of Directors. The Practice Guidelines Committee of the AUA selected the Panel Co-chairmen, and the full Panel roster was assembled by invitation to experts in the field. The Panel evaluated several topics for possible guideline development. Premature ejaculation was selected because of its high prevalence and the availability of a defining body of literature.

Using the MEDLINE® database with MeSH headings related to ejaculatory dysfunction, initial literature searches were performed limiting papers to reports of human studies published in English-language journals between 1966 and January 2001. Only a small number of articles provided outcomes data on PE. Additional studies were identified from references cited in these articles and from recommendations of individual Panel members. The MEDLINE search was last updated in October 2002. Even after the final literature search was completed, however, the Panel continued to scrutinize key references that were identified up until the peer review process.

From a review of abstracts, the Panel chairs selected articles with potentially usable information. Selected papers were reviewed in detail, and relevant data on efficacy and adverse events were extracted and listed in evidence tables (see Appendix 1). Only papers with outcomes data that were relevant to PE, involving pharmacologic treatments generally available in the United States, were included in the evidence tables. If the study was seriously flawed, the article was not considered. Summary tables of adverse event rates and effects of various treatments on latency were created to supplement the data captured in the evidence tables (see Appendices 2 and 3). A

complete list of the 51 references that met all inclusion criteria is available in Appendices 4 and 5. The full Panel reviewed the evidence and summary tables at successive meetings.

Three major limitations were encountered in the evaluation of the evidence that precluded the ability to combine outcomes data and to perform study outcomes comparisons:

- The lack of standardization in studying PE. Clinical trials employ a variety of definitions, entry criteria, physiological measurements and psychometric instruments for evaluation.
- The lack of agreement in quantifying the amount of stimulation that patients experienced. Time to ejaculation is a function of many factors, not the least of which is the nature of the stimulation. The same stimulus may be excessive for one man but elicit little excitement in another. Furthermore, the lack of a consistent stimulus (partner variables, nature of sexual activity, presence or absence of foreplay, preference for single or multiple stimuli) precludes a rigorous experimental design.
- The lack of consistency and accuracy in measurements of time to ejaculation and other outcomes. The most common outcome parameter, time to ejaculation, is either recorded at the time or documented later by recall. These measurements lack accuracy but generally are useful when applied consistently within a single study. Application across multiple studies in a meta-analysis is problematic because any methodological differences will compromise the ability to make a valid comparison. Other common outcome measures concern patient and partner satisfaction. A variety of assessment tools are used, and there is no assurance of comparability between studies

The Panel determined that a meta-analysis was inappropriate due to the disparate outcome measures and populations in the existing studies. The amount of variation between studies also made other less rigorous forms of outcome estimation inaccurate. The Panel's recommendations were developed either solely by consensus or by consensus combined with a review of the available, though limited, evidence. Unless otherwise noted, the statistics cited in this document are derived from the evidence tables.

After this guideline was written, it was reviewed and approved by each member of the Panel and submitted for peer review by 57 physicians. Based on the results of peer assessment, revisions were made and the guideline was forwarded to the Panel again, to the Practice Guidelines Committee, and the Board of Directors of the AUA, all of which rendered approval.

III. Evaluation of the Patient With Premature Ejaculation

Premature ejaculation is a self-reported diagnosis. A sexual history in which the patient uses language that explicitly communicates the circumstances of the condition is the fundamental basis of assessment with time to ejaculation as the most important feature. The opinion of a partner can provide a significant contribution to clinician understanding. A complete description is essential in distinguishing PE from ED, i.e., the inability to attain or maintain an erection, because these conditions frequently coexist. Moreover, some men are unaware that loss of erection after ejaculation is normal; thus, they may erroneously complain of ED when the actual problem is PE.

Recommendation 1:

The diagnosis of PE is based on sexual history alone. A detailed sexual history should be obtained from all patients with ejaculatory complaints.

[Based on Panel consensus.]

When obtaining the patient's history, several important sexual and psychological characteristics should be assessed: frequency and duration of PE, relationship to specific partners, occurrence with all or some attempts, degree of stimulus resulting in PE, nature and frequency of sexual activity (foreplay, masturbation, intercourse, use of visual clues, etc.), impact of PE on sexual activity, types and quality of personal relationships and quality of life, aggravating or alleviating factors, and relationship to drug use or abuse. Laboratory or physiological testing is not required unless the history and a physical examination reveal indications beyond uncomplicated PE.

Recommendation 2:

In patients with concomitant PE and ED, the ED should be treated first.

[Based on Panel consensus.]

Another priority of assessment should be determining whether ED is a concurrent problem. Many patients with ED develop secondary PE, perhaps due to either the need for intense stimulation to attain and maintain an erection or due to the anxiety associated with difficulty in attaining and maintaining an erection. Premature ejaculation may improve in patients when concomitant ED is effectively treated.

IV. Treatment of Premature Ejaculation

Recommendation 3:

The risks and benefits of all treatment options should be discussed with the patient prior to any intervention. Patient and partner satisfaction is the primary target outcome for the treatment of PE.

[Based on Panel consensus.]

As outlined above, the treatments for PE range from psychological and behavioral therapies to pharmacologic therapies. While pharmacologic therapies are the focus of this guideline, other types of interventions may be considered. The patient plays a central role in determining the need for treatment. The patient and possibly his partner can be reassured that PE is a common and treatable disorder. Information on the risks and benefits of all therapeutic options should be presented to the patient (and partner) so that an educated treatment choice may be made by the patient in consultation with the physician. Premature ejaculation is not a life-threatening condition; therefore, safety should be a primary consideration. Some treatments, such as neurectomy and penile prosthesis implantation, have risks that far outweigh their benefits. In addition, none of the medical therapies currently employed in the management of PE have been approved by the U.S. Food and Drug Administration (FDA) for this specific indication. Thus, doses and dosing regimens frequently deviate from that employed for FDA-approved indications, and this difference should be considered in the risk-versus-benefit assessment of pharmacologic therapy.

Efficacy of Proposed Treatments

The preponderance of evidence together with Panel consensus strongly suggest that patients can benefit from the use of several oral or topical medications. At the dosages used in the management of PE, these treatments have been shown to have safety profiles that generally are appropriate to support their use.

Recommendation 4:

Premature ejaculation can be treated effectively with several serotonin reuptake inhibitors (SRIs) or with topical anesthetics. The optimal treatment choice should be based on both physician judgment and patient preference.

[Based on Panel consensus and review of data.]

Oral Medication — Antidepressants

Several antidepressants known to cause anorgasmia and delayed ejaculation have been evaluated in the management of PE. These antidepressants include SRIs, the majority of which are selective (SSRIs) — fluoxetine, paroxetine, and sertraline — and the tricyclic antidepressant clomipramine (Table 1). The SRIs have been successfully utilized in the management of PE. As a group, in clinical trials, the SRIs have provided significant benefit over placebo. Studies have suggested that nefazodone, citalopram, and fluvoxamine are ineffective for the treatment of PE and may be more suitable than other SSRIs for treatment of depression in men *not* wanting ejaculatory impairment.

Table 1. Medical therapy options for the treatment of premature ejaculation*

Oral Therapies	Trade Names [†]	Recommended Dose ‡§
Nonselective serotonin reuptak	e inhibitor	
Clomipramine	$Anafranil^{\circledR}$	25 to 50 mg/day
-		or
		25 mg 4 to 24 h pre-intercourse
Selective serotonin reuptake in		
Fluoxetine	Prozac [®] , Sarafem [®]	5 to 20 mg/day
Paroxetine	$Paxil^{ ext{ iny R}}$	10, 20, 40 mg/day
		or
		20 mg 3 to 4 h pre-intercourse
Sertraline	$Zoloft^{@}$	25 to 200 mg/day
		or
		50 mg 4 to 8 h pre-intercourse
Topical Therapies		
Lidocaine/prilocaine cream	EMLA® Cream	Lidocaine 2.5%/prilocaine 2.5%
		20 to 30 minutes pre-intercourse

^{*}This list does not reflect order of choice or efficacy.

Dosing

Various doses and dosing regimens of the SRIs have been evaluated in efficacy and safety studies of PE. Some studies have employed continuous daily dosing while others use a situational dosing regimen whereby the medication is only taken prior to sexual activity. Different situational dosing regimens also have been assessed, varying timing of the dose prior to sexual activity to the time of peak plasma concentrations of the prescribed agent. The limited data on situational dosing suggest that this regimen may be of use to some men because of the theoretical advantage that less of the drug will be used. In general, though, these SRIs have been designed for continuous usage, and their benefits in the treatment of depression are better established after a period of consistent drug administration. Conversely, continuous administration may foster a problem with patient compliance.

[†]Trade names listed may not be all-inclusive.

[‡] Peak plasma concentrations occur 2 to 8 hours (h) postdose and half-lives range from 1 to 3 days.

[§]Titrate doses from low to high based on response.

Whether continuous or situational dosing is more effective in the management of PE is unclear. The optimal interval for situational dosing before intercourse has not been established and the onset of action of these SRIs for this indication is unknown. However, all Panel members utilize a situational dosing regimen in their practices, and some initiate therapy with daily dosing (loading period). The choice of regimen often is based upon the frequency of sexual activity by the patient.

Duration of Therapy

Therapy for PE most likely will be needed on a continuing basis. There is no clear consensus as to whether SRIs will effect an eventual cure of PE, allowing for discontinuation of the medication, or whether SRIs will be required for life. The Panel members' experience is that PE usually returns upon discontinuing therapy.

Dosing of Specific Serotonin Reuptake Inhibitors

Doses of fluoxetine ranging from 5 to 20 mg/day (see Table 1) are reported to be more effective in delaying ejaculation and enhancing patient/partner satisfaction than placebo. A regimen in which the dose is increased after 1 week (to 40 mg/day or to 60 mg/day) also has been used with success ^{11, 12}. In addition, there is evidence that a clinically beneficial effect may be observed at daily doses as low as 5 mg¹³.

Both daily administration of paroxetine at 10, 20 and 40 mg/day and episodic administration at 20 mg 3 to 4 hours prior to intercourse (see Table 1) have been shown to increase ejaculatory latency^{14, 15, 16}. Due to the limited number of patients evaluated in these trials, the benefit of

increasing the dose to 40 mg/day has not been established. The majority of evidence shows effectiveness with 20 mg daily dosing, thus supporting a general suggestion that this dose of paroxetine provides the greatest benefit in remediating PE.

Sertraline, either given in daily doses of 25, 50, 100 or 200 mg or situationally in doses of 50 mg at 5 p.m. (4 to 8 hours before intercourse) (see Table 1), has been shown to increase ejaculatory latency¹⁷. Higher doses may increase efficacy, but logic suggests that higher doses may be associated with increased frequency of ED and decreased libido. Studies to date, though, have been too small to substantiate this conclusion about dose-related side effects.

Clomipramine, a tricyclic antidepressant with SRI effects, has improved ejaculatory latency and other measures of PE when prescribed at doses of 25 and 50 mg/day or 25 mg 4 to 24 hours prior to intercourse (see Table 1). Adverse event rates and the beneficial effects of clomipramine appear to be dose-related¹⁸.

Adverse Effects

Although the adverse effects of the SRIs have been well described in the management of clinical depression, the following facts should be considered when weighing the risks of prescribing these agents for the patient with PE:

First, men being treated for PE often are different from those being treated for
depression, and the adverse effects of these medications have not been well assessed
in settings other than depression. However, from evidence gathered to date, it
appears that the adverse event profiles of the SRIs reported in the treatment of PE are

similar to those reported in patients being treated for depression. The type and rate of occurrence of side effects appear to be acceptable to most patients and typically include nausea, dry mouth, drowsiness, and reduced libido (see Appendices 1 and 2). Isolated cases of more serious complications, such as mania¹⁹ and withdrawal symptoms, and potential drug interactions also have been associated with the use of SRIs. Pharmacodynamic drug interactions resulting in a "serotonergic syndrome" characterized in mild cases by headache, nausea, sweating, and dizziness and in severe cases by hyperthermia, rigidity, delirium, and coma have been reported rarely with concomitant use of monoamine oxidase inhibitors, lithium, sumatriptan and tryptophan. Pharmacokinetic interactions resulting in alterations in drug blood levels have been reported with the concomitant administration of agents that, like the SRIs, also are metabolized by the cytochrome P450 isoenzyme system or are bound to plasma proteins. Clinically significant pharmacokinetic interactions may rarely occur with the use of anticonvulsants, benzodiazepines, cimetidine, tricyclic antidepressants, antipsychotic agents, tolbutamide, antiarrhythmics, and warfarin especially in the elderly patient.

- Second, doses that are effective in the treatment of PE usually are lower than those recommended in the treatment of depression, suggesting that the frequency and severity of adverse events also could be less.
- Third, because two drug administration regimens, continuous daily dosing and situational dosing, are employed in the treatment of PE, adverse event profiles may differ among patients depending on the regimen prescribed.

The experience with SRIs, as reflected in the evidence tables, and the familiarity of Panel members to date with these medications in this clinical setting suggest that the level of adverse effects is acceptable for the benefit derived in the patient with PE.

Topical Anesthetic Agents

Topical anesthetic agents may be applied to the penis prior to intercourse to delay ejaculation. After topical application, these agents have been used either with or without a condom. The condom may be removed prior to sexual intercourse and the penis washed clean of any residual active compound. Lidocaine/prilocaine cream (2.5 g) applied for 20 to 30 minutes prior to intercourse (see Table 1) has been shown to increase latency time. No significant side effects have been noted. Prolonged application of topical anesthetic (30 to 45 minutes) has been reported to result in loss of erection due to numbness of the penis in a significant percentage of men²⁰. The reduction of penile sensation may limit the acceptability of this method of treatment. Diffusion of residual topical anesthetic on the penis into the vaginal wall also may result in numbness in the partner²¹. Topical anesthetics are contraindicated in patients who are either allergic themselves or have partners who are allergic to any component of the product.

Other Pharmacologic Therapies

Other pharmacologic therapies have been described in the treatment of PE in patients without ED. Intracorporal injection of a vasoactive agent, such as alprostadil, and the administration of sildenafil citrate, therapies effective in the management of ED, have been found to increase latency in patients with PE in a few small studies^{22, 23}. A recent study of 80 men without concomitant ED found that the administration of a combination of sildenafil citrate and

paroxetine on a situational basis enhanced the efficacy of paroxetine alone, although there was an increase in the frequency of the side effects of headache and flushing²⁴. Underlying these interventions is the hypothesis that pharmacologic maintenance of a rigid erection reduces the patient's need to rush to orgasm.

Because ejaculation involves the sympathetic nervous system, adrenergic blockade has been proposed as a treatment for delaying or inhibiting ejaculation. One clinical trial did show modest efficacy with alfuzosin and terazosin²⁵. Phenoxybenzamine and propranolol also have been studied, but the Panel did not believe the evidence was sufficient to support a recommendation for their use^{26, 27, 28}.

V. Future Research

Deficiencies and inconsistencies in the design of and lack of reporting standards for clinical studies on PE have hindered attempts to identify best practices. Future research efforts using well-planned and well-executed randomized, controlled trials are needed to:

- Determine ejaculation latency time in the general population;
- Develop a consensus on the definition of PE;
- Develop standardized, validated instruments to measure outcomes (i.e., patient/partner satisfaction and bother, ejaculatory latency);
- Determine more precisely the efficacy and risks of drug therapies;
- Determine ideal dosing regimens for SRIs (i.e., daily versus situational dosing regimens and whether loading is necessary prior to situational dosing);
- Determine the optimal treatment duration and how or whether to discontinue therapy;

- Determine the long-term acceptability of therapeutic agents to patients;
- Determine the efficacy of combining pharmacologic and behavioral approaches to therapy; and
- Identify the age-specific prevalence of PE.

Other authors^{29, 30} have made recommendations for reporting results in this field that should be considered by investigators studying PE.

VI. Conclusions

A common male sexual disorder, PE traditionally has been treated with psychotherapy or behavioral therapy. This guideline is the first to address the pharmacologic treatment of PE. Although not approved by the FDA for this indication, oral antidepressants and topical anesthetic agents have been shown to delay ejaculation in men with PE and have minimal side effects when used for the treatment of PE. Treatment with oral antidepressants should be started at the lowest possible dose that is compatible with a reasonable chance of success. The choice of additional therapy is based on the patient and partner reports of efficacy, side effects, and acceptance of the therapy as well as on a regular review of alternative approaches. Support and education of the patient and, when possible, the partner are an integral part of PE therapy.

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Appendix 1

Evidence Tables: Pharmacologic Treatment of Premature Ejaculation

- 1-A Fluoxetine
- 1-B Paroxetine
- 1-C Sertraline
- 1-D Clomipramine
- 1-E Topical Anesthetics
- 1-F Adrenergic Blockers
- 1-G Miscellaneous Treatments

Appendix 2

Summary Tables: Adverse Event Rates by Pharmacologic Treatment

Appendix 3

Summary Tables: Effects of Pharmacologic Treatment on Latency

Appendix 4

Articles Selected for Review: Sorted by Author

Appendix 5

Articles Selected for Review: Sorted by ProCite Reference Number

Appendix 1-A. Evidence Tables: Fluoxetine treatment studies

Reference*	Study Population [†]	Treatment Regimen	Outcomes	Comments
Atan, 2000	n=26: age 27 (range, 21-36)	Fluoxetine 20 mg/day for 1	Cured: 8 (30.8%)	There is no indication of
(795257)	With RE and without ED, substance abuse,	week followed by 40 mg/day	Improved: 11 (42.3%)	randomization in the article.
Controlled Trial	infections, diabetes, thyroid disease,	for 7 weeks	Failure: 7 (26.9%)	
Turkey	hypotension or loss of libido. Patients had			
N=43	normal psychiatric consultations.		Side effects:	
			Nausea 3	
			Headache 1	
			Insomnia 2	
			Total pts with side effects: 6	
	n=17: age 31 (range,19-48)	Fluoxetine 20 mg/day and	Cured: 9 (52.9%)	
	With PE and without ED, substance abuse,	local application of lidocaine	Improved: 5 (29.4%)	
	infections, diabetes, thyroid disease, hypo	ointment to the glans 20 min.	Failure: 3 (17.6%)	
	tension or loss of libido. Patients had normal	prior to intercourse.	, ,	
	psychiatric consultations.		Side effects:	
			Nausea 1	
			Headache 4	
			Insomnia 0	
			Total pts with side effects: 5	
Haensel, 1998	n=9: age 41.9±4.6 (range, 26-64)	Treatments included placebo	Latency increase with	All patients received both
(900017)	Patients with premature ejaculation	or fluoxetine (5 mg/day for	fluoxetine 190% (CI: 80-450%;	therapies. Groups were
Crossover	BL: 73±22 sec	two weeks followed by	p=.13) but combined groups 1	significantly different (p<.01)
RCT		10mg/day for two weeks). All	and 3 reached sig. (p=.007)	on age, Zung depression
Netherlands		patients had both treatments	6/7 pts increased latency with	scale, erectile function, and
N=40		in a random order. Each	fluoxetine 1/7 decreased	pretreatment latency.
		treatment lasted 4 weeks	latency	Treatment dose is much
		with a 4-week washout in	-	less than other fluoxetine
	n=7: age 54.6±3.7(range, 40-70)	between.	4/6 pts. decreased latency	studies. Article also has
	Patients with erectile dysfunction		with fluoxetine, 1/6 increased	data on response erotic
	BL: 360±91 sec		and 1/6 inability to ejaculate	stimuli with and without
			in a substantial s	vibratory stimulus.

^{*}Author, year/(Procite number)/Study Design/Location/N=Total patients †age=mean±standard deviation in years BL=baseline latency; RCT=randomized controlled trial

Appendix 1-A. Evidence Tables: Fluoxetine treatment studies

Reference*	Study Population [†]	Treatment Regimen	Outcomes	Comments
Haensel, cont.	n=9: age 51.2±2.7(range, 41-65)		Significant increase in latency	Side effects of fluoxetine:
	Patients with premature ejaculation and		with fluoxetine over placebo	Dry mouth 3
	erectile dysfunction		(p=.03)	Incr. Libido 2
	BL: 89±22 sec		6/8 increased latency with	Loose stools 2
			fluoxetine	Slight palpitations 1
			1/8 decreased	Penile pain 1
			1/8 unchanged	Dizziness 1
	n=15: age 41.3±2.0		7/15 increased latency with	Altered sleep 2
	Normal controls		fluoxetine, 8/15 decreased	Sweating 1
	BL: 535±116 sec		latency	_
				Side effects of placebo:
				Increased libido 1
				Decreased libido 1
				Burning on micturation 1
				Change in stools 1
Kara, 1996	All patients married age 15-50, ½ of whom	Fluoxetine 20 mg/day for 1	Initial latency: 25±12.6 sec	Article gives patient ages,
(12110)	had latencies (average of 3) < 2 min	week	4 wk latency: 180±99.5 sec	spouse ages, and length of
RCT		Fluoxetine 40 mg/day		marriage by groups.
Turkey	n=7 (Initially n=9, but 2 discontinued due to	thereafter	Side effects:	Change in latency status
N=40	side effects)		Headache 1	significant for fluoxetine but
			(discontinued)	not for controls.
			Nausea 2	
			Insomnia 1	
			(discontinued)	
	All patients married age 15-50, ½ of whom	Placebo 1 tablet/day for 1	Initial latency: 30±8.6 sec	
	had latencies (average of 3) < 2 min.	week	4 wk latency: 60±46.9	
		2 tablets/day thereafter	-	
	n =7 (Initially n=8, but 1 dropped from			
	efficacy analysis for not following directions)			

^{*}Author, year/(Procite number)/Study Design/Location/N=Total patients †age=mean±standard deviation in years BL=baseline latency; RCT=randomized controlled trial

Appendix 1-A. Evidence Tables: Fluoxetine treatment studies

Reference*	Study Population [†]	Treatment Regimen	Outcomes	Comments
Reference* Kim, 1998 (12047) Crossover RCT Korea N=53	n=37: age 44 (range, 30-60) 53 heterosexual patients enrolled; 37 completed the study. Reasons for withdrawal: loss to follow-up (5), no efficacy (fluoxetine 3, sertraline 1, placebo 1) and no efficacy plus side effects (clomipramine 1). One patient was excluded from analysis because of delayed (>30 min) ejaculation with sertraline or clomipramine.	Placebo 1 capsule/day first week and 2/day thereafter	OutcomesLatency: 46 ± 41 secPatient Satisfaction: Satisfied 0% Moderate 0% Dissatisfied 100% Partner Satisfaction: Satisfied 0% Moderate 22.2% Dissatisfied 77.8% Latency: 2.27 ± 3.78 minPatient Satisfaction: Satisfied 19.4% Moderate 27.8% Dissatisfied 52.8% Partner Satisfaction: Satisfied 52.8% Partner Satisfaction: Satisfied 36.1% Dissatisfied 52.8% Side effects: Drowsiness Drowsiness Dry mouth Reduced Potency Nausea Vomiting Other 3	Article gives patient ages, spouse ages, and length of marriage by groups. Change in latency status significant for fluoxetine but not for controls. All patients received each therapy and placebo for 4 weeks with a 4-week washout between therapies. Order of administration was randomized.

^{*}Author, year/(Procite number)/Study Design/Location/N=Total patients †age=mean±standard deviation in years BL=baseline latency; RCT=randomized controlled trial

Appendix 1-A. Evidence Tables: Fluoxetine treatment studies

Reference*	Study Population [†]	Treatment Regimen	Outcomes	Comments
Reference* Kim, cont.	Study Population [†]	Treatment Regimen Clomipramine 25 mg/day for first week, 50 mg/day thereafter	Dutcomes Latency: 5.75±6.68 min Patient Satisfaction: Satisfied 25.0% Moderate 38.9% Dissatisfied 36.1% Partner Satisfaction: Satisfied 19.4% Moderate 38.9% Dissatisfied 41.7% Side effects: Drowsiness 6 Dry mouth 2 Reduced Potency 3 Nausea 4	Comments
			Vomiting 0 Other 3 Total pts with side effects: 13	

^{*}Author, year/(Procite number)/Study Design/Location/N=Total patients †age=mean±standard deviation in years BL=baseline latency; RCT=randomized controlled trial

Appendix 1-A. Evidence Tables: Fluoxetine treatment studies

Reference*	Study Population [†]	Treatment Regimen	Outcomes	Comments
Reference* Kim, cont.	Study Population [†]	Treatment Regimen Fluoxetine 20 mg/day for first week, 40 mg/day thereafter	Dutcomes Latency: 2.30±2.08 min Patient Satisfaction: Satisfied 25.0% Moderate 38.9% Dissatisfied 36.1% Partner Satisfaction: Satisfied 19.4% Moderate 38.9% Dissatisfied 41.7% Side effects: Drowsiness 6 Dry mouth 2 Reduced Potency 3 Nausea 4 Vomiting 0 Other 3	Comments
			Total pts with side effects: 13	

^{*}Author, year/(Procite number)/Study Design/Location/N=Total patients †age=mean±standard deviation in years BL=baseline latency; RCT=randomized controlled trial

Appendix 1-A. Evidence Tables: Fluoxetine treatment studies

Sertraline 50 mg/day for first week, 100 mg/day thereafter Patient Satisfaction: Satisfied 41.7% Moderate 36.1% Dissatisfied 22.2%

^{*}Author, year/(Procite number)/Study Design/Location/N=Total patients †age=mean±standard deviation in years BL=baseline latency; RCT=randomized controlled trial

Appendix 1-A. Evidence Tables: Fluoxetine treatment studies

Reference*	Study Population [†]	Treatment Regimen	Outcomes	Comments
Lee, cont.	Otady i Opalation	2 wk washout followed by fluoxetine 20mg/day for 1-2 weeks titrated to 60mg/day based on tolerance. 8 week total trial	Median latency: Post-treatment 9.64±7.0 min YSFI II Scale: Sexual des. 55±21* Erect. Qual. 35±30 Anx. For RE 52±24* Ejac. Satis. 65±18 Partn.Ejac.Sat. 54±17* Overall sex sat. 64±19 Partn. Ovr. Sat. 53±13* *P<.05 Side effects: Nausea and other GI 4 Extremity tingling 1 Dizziness 1 All side effects disappeared in 2-3 weeks after treatment	Comments
Murat Basar, 1999 (12003) RCT Turkey N=57	n=26: age 27±1.1 (range, 21-36) Exclusion criteria: loss of libido; erection failure; alcohol or substance abuse; mental retardation; thyroid disease orthostatic hypotension; previous use of drugs for PE; recent myocardial infarction, uncontrolled diabetes, urogenic infections	Fluoxetine 20 mg/day for 1 week, 40 mg/day later	ceased. Cured 8 (30.8%) Improved 11 (42.3%) Failure 7 (29.6%) Side effects: Nausea Nausea 3 Headache 1 Insomnia 2	Cured and improved not defined. Authors concluded no significant difference between the two treatments.

^{*}Author, year/(Procite number)/Study Design/Location/N=Total patients †age=mean±standard deviation in years BL=baseline latency; RCT=randomized controlled trial

Appendix 1-A. Evidence Tables: Fluoxetine treatment studies

Reference*	Study Population [†]	Treatment Regimen	Outcomes	Comments
Murat Basar, cont.	n=31: age 34±1.7 (range, 21-45) Same exclusion criteria as above.	Sertraline 50 mg/day	Cured 12 (38.78%) Improved 10 (32.3%) Failure 9 (29.0%) Side effects: Nausea 2 Mouth dryness 6	
Raju, 1997 (500001) Letter-CS India N=44	N=44; age 31±6; without clinically detectable organic etiology, and ejaculation prior to intromission or within 10-20 strokes, usually less than 1 min		Total GRISS score Before 10.7±4 After 3.2±2 Side effects included glossitis, vague headache, and lack of concentration	The scale used is not clear since the article describes a 0-5 point scale, but reports larger values. No numbers given for complications
Waldinger, 1998 (12044a) RCT Netherlands N=51	After prescreening exclusions, 60 patients enrolled; 51 completed the study; 6 withdrew due to adverse events and 3 for lack of efficacy			This article describes two RCTs, the first of which is a 5-arm study. 6-week data are recorded here; graphs of intermediate results are
	n=10: age 38±7, partner age 36±7 BL: 21±12 sec	Fluoxetine 20 mg/day	Latency at 6 weeks: 211±251 sec Absolute change: 189±244 sec	presented in the paper.
	n=10: age 44±10, partner age 43±10 BL: 15±17 sec	Fluvoxamine 100 mg/day	Latency at 6 weeks: 55±70 sec Absolute change: 42±57 sec	
	n=11: age 41±8, partner age 39±8 BL: 16±10 sec	Paroxetine 20 mg/day	Latency at 6 weeks: 476±1146 sec Absolute change: 458±1142 sec	

^{*}Author, year/(Procite number)/Study Design/Location/N=Total patients †age=mean±standard deviation in years BL=baseline latency; RCT=randomized controlled trial

Appendix 1-A. Evidence Tables: Fluoxetine treatment studies

Reference*	Study Population [†]	Treatment Regimen	Outcomes	Comments
Waldinger, cont.	n=11: age 40±9, partner age 38±10 BL: 21±12 sec	Sertraline 50 mg/day	Latency at 6 weeks: 117±87 sec Absolute change: 96±84 sec	
	n=9: age 45±4, partner age 41±5 BL: 19±15 sec	Placebo	Latency at 6 weeks: 29±25 sec Absolute change: 10±18 sec	
Yilmaz, 1999 (12032) RCT	n=20: age 37.3 (range, 24-58)	Fluoxetine 20 mg/day for 1 month	Latency: Pre 1.2±1.0 min Post 6.6±7.7 min	40 of 48 patients enrolled. Latency was average reported by patient and not
Turkey N=40	n=20: age 36.5 (range, 22-56)	Placebo	Latency: Pre 1.1±1.3 min Post 4.8±1.0 min	verified. Study also reports data on penile sensory thresholds.

^{*}Author, year/(Procite number)/Study Design/Location/N=Total patients †age=mean±standard deviation in years BL=baseline latency; RCT=randomized controlled trial

Appendix 1-B. Evidence Tables: Paroxetine treatment studies

Reference*	Study Population [†]	Treatment Regimen	Outcomes	Comments
Abdel-Hamid, 2001 (700014) Crossover RCT Egypt N=51	n=31: age 34.09±4.29 (range, 27-42) Heterosexual, married at least 1 year, with primary PE and willing to have intercourse twice a week. Patients did not have history of psychiatric illness, current physical illness, previous surgery or drug known to affect sexual function, current substance abuse, or	Clomipramine 25 mg, 3-5 hrs before planned coitus and not more than twice/week	Results at 4 weeks: Median latency 4 (1-8) min Median anxiety score 11 (4-22) min Median sexual satisfaction score 11 (0-25) min	Study also included a normal control group of 20 patients who were not treated but supplied anxiety scores of 3.7±2.7 (range 1-9) vs. 12.7±5.8 (range 5-25) for the PE subjects.
	ED. Patients each received each of 5 treatments for 4 weeks separated by 2-week washout periods. The treatments were ordered randomly in a double blind manner. BL (med.): 1(0.5-1.5) min Baseline anxiety (med.): 12 (5-25)		2 patients dropped out for lack of efficacy and 1 for side effects and lack of efficacy. Side effects: Dry mouth 3 Anorexia 0 Nausea 1 Headache 0 Flushing 0 Drowsiness 1 Sleepiness 2 Nasal congestion 0	Study contains blinding problems because the Masters and Johnson pause squeeze technique cannot be blinded.
			Yawning 0 Total patients with side effects: 7/28	

^{*}Author, year/(Procite number)/Study Design/Location/N=Total patients †age=mean±standard deviation in years BL=baseline latency; RCT=randomized controlled trial

Appendix 1-B. Evidence Tables: Paroxetine treatment studies

Reference*	Study Population [†]	Treatment Regimen	Outcomes	Comments
Abdel-Hamid,		Paroxetine 20 mg, 3-5 hrs	Results at 4 weeks:	
cont.		before planned coitus and	Median latency	
		not more than twice/week	4(2-10) min	
			Median anxiety score	
			9(5-23) min	
			Median sexual satisfaction	
			score 12(0-29) min	
			2 patients dropped out for lack	
			of efficacy.	
			or omegay.	
			Side effects:	
			Dry mouth 2	
			Anorexia 0	
			Nausea 1	
			Headache 0	
			Flushing 0	
			Drowsiness 0	
			Sleepiness 0 Nasal congestion 0	
			Nasal congestion 0 Yawning 2	
			Total patients with side effects:	
			5/29	

^{*}Author, year/(Procite number)/Study Design/Location/N=Total patients †age=mean±standard deviation in years BL=baseline latency; RCT=randomized controlled trial

Appendix 1-B. Evidence Tables: Paroxetine treatment studies

Reference*	Study Population [†]	Treatment Regimen	Outcomes	Comments
Abdel-Hamid, cont.		Sertraline 50 mg, 3-5 hrs before planned coitus and not more than twice/week	Results at 4 weeks: Median latency	
			3/29	

^{*}Author, year/(Procite number)/Study Design/Location/N=Total patients †age=mean±standard deviation in years BL=baseline latency; RCT=randomized controlled trial

Appendix 1-B. Evidence Tables: Paroxetine treatment studies

Reference*	Study Population [†]	Treatment Regimen	Outcomes	Comments
Abdel-Hamid, cont.		Sildenafil 50 mg, 3-5 hrs before planned coitus and not more than twice/week	Results at 4 weeks: Median latency	
		Masters and Johnson pause squeeze technique	Total patients with side effects: 5/28 Results at 4 weeks: Median latency 3(1-7) min Median anxiety score 12(5-21) min Median sexual satisfaction score 6(0-22) min 2 patients dropped out for lack of efficacy.	

^{*}Author, year/(Procite number)/Study Design/Location/N=Total patients †age=mean±standard deviation in years BL=baseline latency; RCT=randomized controlled trial

Appendix 1-B. Evidence Tables: Paroxetine treatment studies

Reference*	Study Population [†]	Treatment Regimen	Outcomes	Comments
Ludovico, 1996 (12118) CS Italy N=32	n=32: mean age 28 BL: < 1 min (14 ejaculated before penetration)	Paroxetine 20 mg/day for two months.	Latency: 15-20 min Side effects: Severe sensory confusion 1 (withdrew) Sleepiness 14 Mild sensory confusion 21 Side effects disappeared after 15-20 days. PE recurred in 28 patients 2-3 weeks after cessation of therapy.	This is a prospective clinical series. No detailed data given.
McMahon, 1999 (12005) CS Australia N=94	n=94 Heterosexual patients in stable relationships with no other sexual disorders. BL: < 1 min Group A: n=61: age 40 (range, 22-61) Mean latency 0.4 min, 37 with lifelong PE, 6 severe (never had intravaginal ejaculation) Group B: n=33: age 37 (range, 18-56) Mean latency 0.4 min, 18 with lifelong PE, 4 severe	Group A, Phase 1: Paroxetine 20 mg/day for 4 weeks.	Latency 4.5 min Side effects: Anejaculation 5 (1 withdrawal) Drowsiness & anorexia 1 Minor Gl upset 2 Reduced libido 3 Inhibited orgasm 3 Ejaculation restored for anejaculation patients with lower dose.	Also data on frequency of intercourse.

^{*}Author, year/(Procite number)/Study Design/Location/N=Total patients †age=mean±standard deviation in years BL=baseline latency; RCT=randomized controlled trial

Appendix 1-B. Evidence Tables: Paroxetine treatment studies

Reference*	Study Population [†]	Treatment Regimen	Outcomes	Comments
McMahon,		Group A, Phase 2:	Latency 3.9 min	
cont.		Paroxetine 20 mg single		
		dose 3-4 hrs before	No side effects	
		intercourse (Phase II for		
		Group A patients with		
		success in Phase I above)		
		for 4 weeks		
		Group B: Paroxetine 20 mg	Latency 1.5 min	
		single dose 3-4 hrs before		
		intercourse for 4 weeks	No side effects	
McMahon,	n=26: mean age 39.5	Paroxetine 20 mg, 3-4 hrs	Pretreatment latency	First of two studies in one
1999	19 patients with primary PE, 3 never	before intercourse for 4	0.3min	article. Study is single
(12020a)	ejaculated vaginally.	weeks. 3-week washout	Paroxetine latency at 4 weeks	blind only. Also data on
Crossover RCT	Average latency = 0.3 min	followed by placebo 3-4 hrs	3.2 min	intercourse frequency.
Australia	The two groups each had 13 patients.	before intercourse for 4	Placebo latency at 4 weeks	
N=26		weeks.	0.45 min	
			No side effects with either pill.	
		Placebo, 3-4 hrs before	Pretreatment latency	
		intercourse for 4 weeks. 3-	0.3 min	
		week washout followed by	Placebo latency at 4 weeks	
		paroxetine 20 mg 3-4 hrs	0.6 min	
		before intercourse for 4	Paroxetine latency at 4 weeks	
		weeks.	3.5 min	
			No side effects with either pill.	

^{*}Author, year/(Procite number)/Study Design/Location/N=Total patients †age=mean±standard deviation in years BL=baseline latency; RCT=randomized controlled trial

Appendix 1-B. Evidence Tables: Paroxetine treatment studies

Reference*	Study Population [†]	Treatment Regimen	Outcomes	Comments
McMahon,	n=42: mean age 40.5	Paroxetine 10 mg/day for 3	BL: 0.5 min	Second of two studies in
1999	32 patients with primary PE, 10 never	weeks, followed by		one article. Study is single
(12020b)	ejaculated vaginally.	paroxetine 20 mg 3-4 hrs	Results at the end of each	blind only.
Crossover RCT	Average latency 0.5 min	before intercourse for 4	section:	
Australia	The two groups each had 21 patients.	weeks, followed by 3-week	Paroxetine daily latency	Side effects were not listed
N=42		washout, then placebo daily	4.3 min	by group and were as
		for 3 weeks followed by	Paroxetine PRN latency	follows:
		placebo 3-4 hrs before	5.8 min	Paroxetine Daily:
		intercourse for 4 weeks.	Placebo daily latency	Anorexia 1
			0.9 min	Anejaculation 3
			Placebo PRN latency	GI upset 3
			0.6 min	Reduced libido 2
		Placebo daily for 3 weeks,	BL: 0.5 min	
		then placebo 3-4 hrs before		Placebo Daily:
		intercourse for 4 weeks. 3-	Results at the end of each	Erectile dysfunction 2
		week washout followed by	section:	
		paroxetine 10 mg/day for 3	Paroxetine daily latency	Placebo PRN:
		weeks followed by	3.3 min	Headache 1
		paroxetine 20 mg 3-4 hrs	Paroxetine PRN latency	
		before intercourse for 4	6.1 min	
		weeks	Placebo daily latency	
			0.8 min	
			Placebo PRN latency	
			1.1 min	

^{*}Author, year/(Procite number)/Study Design/Location/N=Total patients †age=mean±standard deviation in years BL=baseline latency; RCT=randomized controlled trial

Appendix 1-B. Evidence Tables: Paroxetine treatment studies

Reference*	Study Population [†]	Treatment Regimen	Outcomes	Comments
Salonia, 2002	N=40; mean age 34 (range, 19-46); primary	10 mg/day paroxetine for 21	Latency:	Analysis of IIEF data
(500005)	PE 29, secondary PE 11, never ejaculated	days, followed by 20 mg 3-4	Baseline .33±.04	showed significant
CT	vaginally 8	hrs before intercourse for 6	3 mo 3.7±10	differences at 6 months
Italy		months	6 mo 4.2±.03	only in intercourse
N=80				satisfaction and overall
			Side effects:	satisfaction.
			Anejaculation 1/40	
			GI upset/nausea 5/40	
			Headache 4/40	
			Decr. Libido 2/40	
			Flushing 0/40	
	N=40; mean age 36 (range 21-47); primary	10 mg/day paroxetine for 21	Latency:	
	PE 33, secondary PE 7, never ejaculated	days, followed by 20 mg 3-4	Baseline .35±.05	
	vaginally 12	hrs plus 50 mg sildenafil	3 mo 4.5±07	
		before intercourse for 6	6 mo 5.3±.02	
		months		
			Side effects:	
			Anejaculation 1/40	
			GI upset/nausea 6/40	
			Headache 8/40	
			Decr. Libido 1/40	
			Flushing 6/40	
Waldinger,	n=15: age 38±11	Paroxetine 20 mg/day for 6	Geometric mean latency by	Randomized blinded
2001	Married or in a relationship for 15±10 years	weeks	week (sec):	controlled trial of a
(795220)	BL: 22±15 sec		0 17.58	superselective SSRI vs.
RCT			1 34.17	paroxetine. Also data on
Netherlands			2 57.27	intercourse frequency,
N=30			3 116.00	which rose with paroxetine
			4 141.16	and fell with citalopram.
			5 170.50	
			6 152.28	

^{*}Author, year/(Procite number)/Study Design/Location/N=Total patients †age=mean±standard deviation in years BL=baseline latency; RCT=randomized controlled trial

Appendix 1-B. Evidence Tables: Paroxetine treatment studies

Reference*	Study Population [†]	Treatment Regimen	Outcomes	Comments
Waldinger, cont.	n=15: age 39±8 Married or in a relationship for 24±10 years BL: 24±10 sec	Citalopram 20 mg/day for 6 weeks	Geometric mean latency by week (sec): 0	
Waldinger, 2001 (795222) RCT Netherlands N=48	n=12: age 38±10 Married for 15±12 years with primary PE	Placebo, morning and evening	Geometric mean latency by week (sec): 0 15.1 1 20.8 2 19.3 3 19.1 4 23.6 5 18.8 6 18.1	All patients had a 1-month baseline and were prohibited from using condoms or topical anesthetics through the study. Study was randomized and blinded. Patients used stopwatches to measure latency.
	n=12: age 40±7 Married for 17±7 years with primary PE	Paroxetine 20 mg/day, morning and evening	Geometric mean latency by week (sec): 0 17.1 1 36.7 2 71.1 3 119.2 4 88.3 5 146.0 6 107.9	,

^{*}Author, year/(Procite number)/Study Design/Location/N=Total patients †age=mean±standard deviation in years BL=baseline latency; RCT=randomized controlled trial

Appendix 1-B. Evidence Tables: Paroxetine treatment studies

Reference*	Study Population [†]	Treatment Regimen	Outcomes	Comments
Waldinger, cont.	n=12: age 41±9 Married for 15±9 years with primary PE	Sertraline 50 mg/day, morning and evening	Geometric mean latency by week (sec): 0 13.9 1 25.2 2 38.6 3 33.8 4 42.7 5 58.1 6 50.3	
	n=12: age 35±6 Married or in a relationship for 12±8 years with primary PE	Nefazodone 400 mg/day, morning and evening	Geometric mean latency by week (sec): 0 16.8 1 19.1 2 14.2 3 25.6 4 28.4 5 15.6 6 17.9	
Waldinger, 1998 (12044a) RCT	After prescreening exclusions, 60 patients enrolled; 51 completed the study; 6 withdrew due to adverse events and 3 for lack of efficacy			This article describes two RCTs, the first of which is a 5-arm study. 6-week data are recorded here; graphs
Netherlands N=51	n=10: age 38±7, partner age 36±7 BL: 21±12 sec	Fluoxetine 20 mg/day	Latency at 6 weeks: 211±251 sec Absolute change: 189±244 sec	of intermediate results are presented in the paper.
	n=10: age 44±10, partner age 43±10 BL: 15±17 sec	Fluvoxamine 100 mg/day	Latency at 6 weeks: 55±70 sec Absolute change: 42±57 sec	
	n=11: age 41±8, partner age 39±8 BL: 16±10 sec	Paroxetine 20 mg/day	Latency at 6 weeks: 476±1146 sec Absolute change: 458±1142 sec	

^{*}Author, year/(Procite number)/Study Design/Location/N=Total patients †age=mean±standard deviation in years BL=baseline latency; RCT=randomized controlled trial

Appendix 1-B. Evidence Tables: Paroxetine treatment studies

Reference*	Study Population [†]	Treatment Regimen	Outcomes	Comments
Waldinger, cont.	n=11: age 40±9, partner age 38±10 BL: 21±12 sec	Sertraline 50 mg/day	Latency at 6 weeks: 117±87 sec Absolute change: 96±84 sec	
	n=9: age 45±4, partner age 41±5 BL: 19±15 sec	Placebo	Latency at 6 weeks: 29±25 sec Absolute change: 10±18 sec	
Waldinger, 1998 (12044b) RCT Netherlands N=32	n=24 with latency ≤ 1min: age 46±6 (range, 31-45) BL: 18±13 sec (range, 1-46)	12 randomized to paroxetine 20 mg/day and 12 to placebo 1 paroxetine and 3 placebo patients dropped out	Paroxetine treated latency increased 580% (CI 314-1025%). No statistically significant increase in placebo group.	Study notes paroxetine had no clinically significant effects on libido or erectile function and no statistically significant differences between paroxetine and
	n=8 with latency > 1 min: age 47±3 (range 43-53) BL: 82±27 sec (range, 57-130)	5 patients randomized to paroxetine 20 mg/day and 3 to placebo	Paroxetine treated latency increased 596% (CI 225-1388%).	placebo on other side effects (unspecified).
		1 placebo patient dropped out	No significant increase in placebo group.	

^{*}Author, year/(Procite number)/Study Design/Location/N=Total patients †age=mean±standard deviation in years BL=baseline latency; RCT=randomized controlled trial

Appendix 1-B. Evidence Tables: Paroxetine treatment studies

Reference*	Study Population [†]	Treatment Regimen	Outcomes	Comments
Waldinger, 1997 (12088) RCT Netherlands N=34	n=17: age 44 (range, 23-57) All patients in stable heterosexual relationship with primary PE and no ED, substance or alcohol abuse, mental disorders, physical illnesses, use of medications, or history of sexual abuse of the patient or partner. SCL-90 depression and fear scores similar to normal population.	Paroxetine 20 mg/day for 8 weeks. 2 capsules/day (1 placebo).	BL: 13 3 wk latency: 300 8 wk latency: 300 2 patients discontinued for anejaculation and frequent yawning. 1 patient discontinued 2 nd pill (placebo) for yawning, perspiration, and fatigue. At 8 weeks, 6 patients still had yawning, perspiration, dry mouth, fatigue and/or nausea. One patient had slight reduction in erectile function at 8 weeks.	Study also contains partner data and estimates of latency, which matched the patients' estimates. Latency estimates are medians not mean. Patients who dropped were not included in the latency results, but authors said including them did not change results.
	n=17: age 43 (range 32-55) All patients in stable heterosexual relationship with primary PE and no ED, substance or alcohol abuse, mental disorders, physical illnesses, use of medications, or history of sexual abuse of the patient or partner. SCL-90 depression and fear scores similar to normal population.	Paroxetine 40 mg/day for 8 weeks. 2 capsules/day both paroxetine.	BL: 10 3 wk latency: 240 8 wk latency: 540 4 patients discontinued 2 nd capsule for anejaculation, yawning, fatigue and perspiration. One also experienced reduced libido and dry mouth. Reducing dose reduced symptoms. At 8 weeks 7 patients still had yawning, perspiration, dry mouth, fatigue and/or nausea.	

^{*}Author, year/(Procite number)/Study Design/Location/N=Total patients †age=mean±standard deviation in years BL=baseline latency; RCT=randomized controlled trial

Appendix 1-B. Evidence Tables: Paroxetine treatment studies

Reference*	Study Population [†]	Treatment Regimen	Outcomes	Comments
Waldinger, 1994 (12172) RCT Netherlands N=17	n=8: age 41 (range, 27-48) 7 with primary PE and 1 with secondary. 2 patients dropped out. One in the first week and one in the third because of latencies > 30 min Reducing dose to 20 mg yielded latency of 7 min, but patient was excluded from analysis.	Paroxetine 20 mg/day first week and 40 mg/day for 5 more weeks.	Latency – patient assessment: Initial 30 (3-4) sec 3 wks 7.5 (3-20) min 6 wks 10 (5-20) min Latency – partner assessment: Initial 10 (3-30) sec 3 wks 8.5 (2-17.5) min 6 wks 10.0 (5-17.5) min	Data on number of thrusts are also available in the study. Study stated there were no statistically significant differences in side effects at any time point.
	n=9: age 38 (range, 30-47) 7 with primary PE and 2 with secondary. One patient dropped out in the first week.	Placebo, 1 capsule/day for first week and 2 per day for 5 more weeks.	Latency – patient assessment: Initial 15 (5-90) sec 3 wks 20 (5-120) sec 6 wks 15 (5-120) sec Latency – partner assessment: Initial 30 (5-90) sec 3 wks 23 (5-60) sec 6 wks 33 (10-90) sec	

^{*}Author, year/(Procite number)/Study Design/Location/N=Total patients †age=mean±standard deviation in years BL=baseline latency; RCT=randomized controlled trial

Appendix 1-C. Evidence Tables: Sertraline treatment studies

Reference*	Study Population [†]	Treatment Regimen	Outcomes	Comments
Abdel-Hamid, 2001 (700014) Crossover RCT	n=31: age 34.09±4.29 (range, 27-42) Heterosexual, married at least 1 year, with primary PE and willing to have intercourse twice a week. Patients did not have history	Clomipramine 25 mg, 3-5 hrs before planned coitus and not more than twice/week	Results at 4 weeks: Median latency 4 (1-8) min Median anxiety score	Study also included a normal control group of 20 patients who were not treated but supplied anxiety
Egypt N=51	of psychiatric illness, current physical illness, previous surgery or drug known to affect sexual function, current substance abuse, or ED.		11 (4-22) min Median sexual satisfaction score 11 (0-25) min	scores of 3.7±2.7(range 1-9) vs. 12.7±5.8 (range 5-25) for the PE subjects.
	Patients each received each of 5 treatments for 4 weeks separated by 2-week washout periods. The treatments were ordered randomly in a double blind manner.		2 patients dropped out for lack of efficacy and 1 for side effects and lack of efficacy.	Study contains blinding problems because the Masters and Johnson pause squeeze technique
	BL (med.): 1 (0.5-1.5) min Baseline anxiety (med.): 12 (5-25) min		Side effects: Dry mouth 3 Anorexia 0 Nausea 1	cannot be blinded.
			Headache 0 Flushing 0 Drowsiness 1 Sleepiness 2	
			Nasal congestion 0 Yawning 0 Total patients with side effects: 7/28	

^{*}Author, year/(Procite number)/Study Design/Location/N=Total patients †age=mean±standard deviation in years BL=baseline latency; RCT=randomized controlled trial

Appendix 1-C. Evidence Tables: Sertraline treatment studies

Reference*	Study Population [†]	Treatment Regimen	Outcomes	Comments
Abdel-Hamid, cont.	Study Population	Paroxetine 20 mg, 3-5 hrs before planned coitus and not more than twice/week	Results at 4 weeks: Median latency	Comments
			5/29	

^{*}Author, year/(Procite number)/Study Design/Location/N=Total patients †age=mean±standard deviation in years BL=baseline latency; RCT=randomized controlled trial

Appendix 1-C. Evidence Tables: Sertraline treatment studies

Reference*	Study Population [†]	Treatment Regimen	Outcomes	Comments
Reference* Abdel-Hamid, cont.	Study Population ^T	Sertraline 50 mg, 3-5 hrs before planned coitus and not more than twice/week	Results at 4 weeks: Median latency 3 (1-10) min Median anxiety score 11 (5-22) min Median sexual satisfaction score 10 (0-31) min 2 patients dropped out for lack of efficacy. Side effects: Dry mouth Anorexia Nausea 1	Comments
			Headache 0 Flushing 0 Drowsiness 1 Sleepiness 0 Nasal congestion 0 Yawning 0 Total patients with side effects: 3/29	

^{*}Author, year/(Procite number)/Study Design/Location/N=Total patients †age=mean±standard deviation in years BL=baseline latency; RCT=randomized controlled trial

Appendix 1-C. Evidence Tables: Sertraline treatment studies

Reference*	Study Population [†]	Treatment Regimen	Outcomes	Comments
Abdel-Hamid, cont.		Sildenafil 50 mg, 3-5 hrs before planned coitus and not more than twice/week	Results at 4 weeks: Median latency	
		Masters and Johnson pause squeeze technique	5/28 Results at 4 weeks: Median latency 3 (1-7) Median anxiety score 12 (5-21) Median sexual satisfaction score 6 (0-22) 2 patients dropped out for lack of efficacy.	

^{*}Author, year/(Procite number)/Study Design/Location/N=Total patients †age=mean±standard deviation in years BL=baseline latency; RCT=randomized controlled trial

Appendix 1-C. Evidence Tables: Sertraline treatment studies

Reference*	Study Population [†]	Treatment Regimen	Outcomes	Comments
Balbay, 1997 (12063) CS N=22	n=22: age 26-53 With undefined PE. 4 patients lost to follow-up.	Oral sertraline 50 mg/day for 2 weeks.	Efficacy: Completely satisfied 14/16 Not completely satisfied 2/16 Side effects: Patients reported 5/18 2 patients discontinued due to	Study also has data on patient condition based on whether the patient was satisfied. Minimal data are available – side effects not defined, efficacy measure soft and patient condition not well defined.
Basar, 1999 (12003) RCT Turkey N=57	n=26: age 27±1.1 (range, 21-36) Exclusion criteria: loss of libido; erection failure; alcohol or substance abuse; mental retardation; thyroid disease orthostatic hypotension; previous use of drugs for PE; recent myocardial infarction, uncontrolled diabetes, urogenic infections	Fluoxetine 20 mg/day for 1 week 40 mg/day later	unspecified side effects Cured 8 (30.8%) Improved 11 (42.3%) Failure 7 (29.6%) Side effects: Nausea 3 Headache 1 Insomnia 2	Cured and improved not defined. Authors concluded no significant difference between the two treatments.
	n=31: age 34±1.7 (range, 21-45) Same exclusion criteria as above.	Sertraline 50 mg/day	Cured 12 (38.78%) Improved 10 (32.3%) Failure 9 (29.0%) Side effects: Nausea 2 Mouth dryness 6	

^{*}Author, year/(Procite number)/Study Design/Location/N=Total patients †age=mean±standard deviation in years BL=baseline latency; RCT=randomized controlled trial

Appendix 1-C. Evidence Tables: Sertraline treatment studies

Reference*	Study Population [†]	Treatment Regimen	Outcomes	Comments
Biri, 1998	n=37: age 31.1 (range, 21-54)	Sertraline 50 mg/day for 4	Latency:	
(12033)	Patients with latencies < 1 min during	weeks,	Initial 40.93±12.6	
RCT	previous 6 months.	22 patients	Treated 325.4±261.7	
Turkey	No impotence.			
N=37			Side effects:	
			Headache 6/22	
			Sleepiness 6/22	
			Diarrhea 3/22	
			Dry mouth 2/22	
			PE recurred in 19 patients 4	
			weeks after cessation of	
			treatment.	
		Placebo for 4 weeks,	Latency:	
		15 patients	Initial 43.53±20.2	
			Treated 114.4±93.7	
			Cide effects.	
			Side effects:	
			Headache 3/15	
			Sleepiness 3/15	
			Diarrhea 1/15	
			Dry mouth 0/15	

^{*}Author, year/(Procite number)/Study Design/Location/N=Total patients †age=mean±standard deviation in years BL=baseline latency; RCT=randomized controlled trial

Appendix 1-C. Evidence Tables: Sertraline treatment studies

Reference*	Study Population [†]	Treatment Regimen	Outcomes		Comments
Chia, 2002	N=52; without neurological disease or	Sertraline 50 mg 4 hours	Latency (sec)		
(500002)	depression, with primary PE. 12 age 20-30,	prior to intercourse for 6	Baseline	46	
ĊS	24 age 30-40, 14 age 40-50, 2 age 50-60.	months	6 months	247.2	
Singapore N=87			Treatment failures	3/52	
			Sexual Satisfaction		
			(mean of 5 pt scale)		
			Baseline	.81	
			6 months	2.21	
			Side effects		
			Dizziness	3/52	
			Nausea	2/52	
			Dry mouth	1/52	
	N=35; without neurological disease or]	Latency (sec)		
	depression, with ED treated successfully		Baseline	34.6	
	with sildenafil. Pts with PE prior to ED		6 months	111.6	
	excluded. 3 age 20-30, 13 age 30-40, 11 age 40-50, 8 age 50-60.		Treatment failures	7/35	
	age 40-30, 6 age 30-60.		Sexual Satisfaction		
			(mean of 5 pt scale)		
			Baseline	.51	
			6 months	1.17	
			Side effects		
			Dizziness	4/35	
			Nausea	3/35	
			Dry mouth	1/35	

^{*}Author, year/(Procite number)/Study Design/Location/N=Total patients †age=mean±standard deviation in years BL=baseline latency; RCT=randomized controlled trial

Appendix 1-C. Evidence Tables: Sertraline treatment studies

Reference*	Study Population [†]	Treatment Regimen	Outcomes	Comments
Kim, 1999 (12013) CS Korea N=24	n=24: age 32.5 (range, 27-52) Patients with primary PE. Latency < 1 min at least 50% of time. Patients had one stable partner, were not receiving psychotropic medication, were not depressed, or did not have any other medical disease or symptoms. Reasons for withdrawal: 6 patients were excluded from efficacy analysis due to drop out (3 non-compliance, 2 unknown, 1 intercurrent illness).	Pretreatment	Latency 23±19 sec Patient satisfaction (0-5) 0.8±0.8 Partner satisfaction 1.1±0.7 Side effects not linked to dose schedule: Delayed ejaculation 1 Fatigue 2 Numbness in extremities 1 No withdrawals due to side effects.	
		Sertraline 50 mg/day for 2 weeks	Latency 5.9±4.2 min Patient satisfaction (0-5) 3.8±1.2 Partner satisfaction 3.2±1.6	
		Sertraline 50 mg at 5 pm on days when intercourse was anticipated (4-8 hrs before intercourse). Dose titrated up to 100 mg in 3 rd week if needed. Total of 4 or 6 weeks (not clear).	At 2 weeks: Latency 5.1±3.8 min Patient satisfaction (0-5) 3.4±1.0 Partner satisfaction 3.1±1.4	
			At 4 weeks: Latency 4.5±2.7 min Pt. satisfaction (0-5) 3.2±0.7 Partner satisfaction 3.3±1.2	

^{*}Author, year/(Procite number)/Study Design/Location/N=Total patients †age=mean±standard deviation in years BL=baseline latency; RCT=randomized controlled trial

Appendix 1-C. Evidence Tables: Sertraline treatment studies

Reference*	Study Population [†]	Treatment Regimen	Outcomes	Comments
Reference* Kim, 1998 (12047) Crossover RCT Korea N=53	n=37: age 44 (range, 30-60). 53 heterosexual patients enrolled; 37 completed the study. Reasons for withdrawal: loss to follow-up (5), no efficacy (fluoxetine 3, sertraline 1, placebo 1) and no efficacy plus side effects (clomipramine 1). One patient was excluded from analysis because of delayed (> 30 min) ejaculation with sertraline or clomipramine.	Placebo 1 capsule/day first week and 2/day thereafter.	OutcomesLatency:46±41 secPatient Satisfaction: Satisfied0% Moderate0% DissatisfiedPartner Satisfaction: Satisfied0% Moderate22.2% DissatisfiedDissatisfied77.8%Latency:2.27±3.78 minPatient Satisfaction: Satisfied19.4% Moderate27.8% DissatisfiedDissatisfied52.8%Partner Satisfaction: Satisfied11.1% Moderate36.1% DissatisfiedDissatisfied52.8%Side effects: Drowsiness2 Dry mouth0 Reduced PotencyReduced Potency3 Nausea1 VomitingOther3	All patients received each therapy and placebo for four weeks with a 4-week washout between therapies. Order of administration was randomized.
			Total pts with side effects: 7	

^{*}Author, year/(Procite number)/Study Design/Location/N=Total patients †age=mean±standard deviation in years BL=baseline latency; RCT=randomized controlled trial

Appendix 1-C. Evidence Tables: Sertraline treatment studies

Reference*	Study Population [†]	Treatment Regimen	Outcomes	Comments
Kim, cont.		Clomipramine 25 mg/day for first week, 50 mg/day thereafter	Latency: 5.75±6.68 min Patient Satisfaction: Satisfied 25.0% Moderate 38.9% Dissatisfied 36.1% Partner Satisfaction: Satisfied 19.4% Moderate 38.9% Dissatisfied 41.7% Side effects: Drowsiness 6 Dry mouth 2 Reduced Potency 3 Nausea 4 Vomiting 0 Other 3	
			Total pts with side effects: 13	

^{*}Author, year/(Procite number)/Study Design/Location/N=Total patients †age=mean±standard deviation in years BL=baseline latency; RCT=randomized controlled trial

Appendix 1-C. Evidence Tables: Sertraline treatment studies

Reference*	Study Population [†]	Treatment Regimen	Outcomes	Comments
Kim, cont.		Fluoxetine 20 mg/day for first week, 40 mg/day thereafter	Latency: 2.30±2.08 min Patient Satisfaction: Satisfied 25.0% Moderate 38.9% Dissatisfied 36.1% Partner Satisfaction: Satisfied 19.4% Moderate 38.9% Dissatisfied 41.7% Side effects: Drowsiness 6 Dry mouth 2 Reduced Potency 3 Nausea 4 Vomiting 0 Other 3	
			Total pts with side effects: 13	

^{*}Author, year/(Procite number)/Study Design/Location/N=Total patients †age=mean±standard deviation in years BL=baseline latency; RCT=randomized controlled trial

Appendix 1-C. Evidence Tables: Sertraline treatment studies

Reference*	Study Population [†]	Treatment Regimen	Outcomes	Comments
Kim, cont.		Sertraline 50 mg/day for first	Latency: 4.27±5.69 min	
		week, 100 mg/day thereafter		
			Patient Satisfaction:	
			Satisfied 41.7%	
			Moderate 36.1%	
			Dissatisfied 22.2%	
			Partner Satisfaction:	
			Satisfied 30.6%	
			Moderate 38.9%	
			Dissatisfied 30.6%	
			Side effects:	
			Drowsiness 7	
			Dry mouth 4	
			Reduced Potency 3	
			Nausea 2	
			Vomitting 0	
			Other 0	
			Total pts with side effects: 12	
McMahon,	n=19: age 40	Sertraline 50 mg/day for 4	Latency:	Single blind study. Overall
1998	Heterosexual in stable relationships without	weeks, washout for 4 weeks,	Initial 0.3 min	patient age range (19-70).
(12057)	other sexual disorders or other physical or	placebo for 4 weeks.	After sertraline 3.4 min	In patients who had not
Crossover RCT	psychological ailments.	Patients asked to not use	After washout 0.6 min	previously achieved intra-
Australia	BL: < 1 min	condoms or anesthetics.	After placebo 0.5 min	vaginal ejaculation, 5/8
N=37				achieved it for the first time
			Frequency of intercourse:	with sertraline. 29/35 who
			Initial 0.6/wk	completed the study
			After sertraline 3.3/wk	continued sertraline open
			After washout 1.6/wk	label after the study. Stage
			After placebo 1.0/wk	withdrawal of the drug

^{*}Author, year/(Procite number)/Study Design/Location/N=Total patients †age=mean±standard deviation in years BL=baseline latency; RCT=randomized controlled trial

Appendix 1-C. Evidence Tables: Sertraline treatment studies

Reference*	Study Population [†]	Treatment Regimen	Outcomes	Comments
McMahon,	n=18: age 42	Placebo for 4 weeks,	Latency:	every 4 weeks allowed
cont.	Heterosexual in stable relationships without	washout for 4 weeks,	Initial 0.3 min	20/29 to discontinue
	other sexual disorders or other physical or	sertraline 50 mg/day for 4	After placebo 0.5 min	treatment after a mean of
	psychological ailments.	weeks. Patients asked to not	After washout 0.5 min	7.3 months and maintain
	BL: < 1 min	use condoms or anesthetics.	After sertraline 3.0 min	ejaculatory control with mean latency of 4.1
			Frequency of intercourse: Initial 0.4/wk	according to authors.
			After placebo 0.8/wk	Sertraline side effects:
			After washout 0.5/wk	Anejaculation, withdrawal:
			After sertraline 3.1/wk	2
				Drowsiness, anorexia: 1
				Gl upset: 2
				No ED, reduced libido, or
				reduced orgasmic intensity.
				One patient had minor ED
				on placebo.
McMahon,	n=46: age 42 (range, 22-63)	Setraline 25 mg/day for 3	Latency:	Study also includes data on
1998 (12037)	All heterosexual in stable relationships with no other sexual disorders. 36 had primary	weeks.	7.6 (0-20) min	intercourse frequency.
CS Australia	PE and 10 had secondary. 6 men had severe PE, who had never achieved vaginal		Anejaculation: 0	
N=46	ejaculation. Each patient received each		Intravaginal ejaculation in 4/6	
11-40	dose for 3 weeks followed by a 3week		severe cases	
	washout period. The doses were given in		Severe dudes	
	increasing order.		Adverse events:	
	BL: 1.0 min (range, 0 to 5 min).		Anorexia 0	
			Anxiety 0	
			Dizziness 1	
			Drowsiness 0	
			Dyspepsia 0	
			Erectile dysfunction 0	
			Reduced libido 0	

^{*}Author, year/(Procite number)/Study Design/Location/N=Total patients †age=mean±standard deviation in years BL=baseline latency; RCT=randomized controlled trial

Appendix 1-C. Evidence Tables: Sertraline treatment studies

Reference*	Study Population [†]	Treatment Regimen	Outcomes	Comments
McMahon,		Setraline 50 mg/day for 3	Latency:	
cont.		weeks.	13.1 (7-anej) min	
			Anejaculation: 4	
			Anejaculation. 4	
			Intravaginal ejaculation in all 6	
			severe cases	
			Adverse events:	
			Anorexia 1	
			Anxiety 0	
			Dizziness 0	
			Drowsiness 1	
			Dyspepsia 1	
			Erectile dysfunction 0	
			Reduced libido 0	
		Setraline 100 mg/day for	Latency:	
		three weeks	16.4 (7-anej) min	
			Anejaculation: 10	
			Intravaginal ejaculation in all 6	
			severe cases	
			Adverse events:	
			Anorexia 2 Anxiety 2 Dizziness 0	
			Drowsiness 2 Dyspepsia 2	
			Erectile dysfunction 2	
			Reduced libido 2	

^{*}Author, year/(Procite number)/Study Design/Location/N=Total patients †age=mean±standard deviation in years BL=baseline latency; RCT=randomized controlled trial

Appendix 1-C. Evidence Tables: Sertraline treatment studies

Reference*	Study Population [†]	Treatment Regimen	Outcomes	Comments
Mendels, 1995	n=52: age 25-52	Sertraline 50 mg/day titrated	Efficacy (min):	The data reporting in this
(12139)	Heterosexual with only 1 partner in the last 6	from 50 to 200 mg/day	Latency (n=22)	study is unclear and this
RCT	months.	during weeks 1-3. Total 8	Baseline 0.98±1.15	extraction attempts to
N=52		weeks and 26 patients. Mean	Endpoint 5.43±5.62	summarize the results.
	Reasons for exclusion: Patients taking	final dosage 121 mg/day. 2-	Change 3.58±3.82	Latency, patient
	psychotropic medications, depressed (> 14	week washout titrating down	Ejaculation during foreplay	satisfaction, and
	on Hamilton scale), receiving therapy for	to 0 mg/day.	(n=24)	ejaculation during foreplay
	sexual dysfunction or with significant medical		Baseline 3	based on patients'
	disease or symptoms.		During treatment 1	assessment. Partner
	4 patients excluded from analysis: 1		Patient satisfaction (n=24) 0-4	satisfaction based on
	sertraline without baseline data, 1 sertraline		Baseline 1.42±1.10	partner assessment.
	dropped out in the first 3 weeks, 2 placebo		Endpoint 2.42±1.28	Article contains data on
	dropped out in first 3 weeks.		Change 1.00±1.47	many variables from both
	BL: < 1 min on at least 50% of tries in last 6		Partner satisfaction (n=19) 0-4	patient and partner
	months.		Baseline 1.53±0.90	assessment as well as
			Endpoint 2.58±1.22	global clinical impression
			Change 1.05±1.18	from health care provider.
			Side effects (n=26):	
			Any adverse event 17	
			Discontinued due to AE 0	
			Diarrhea 6	
			Anejaculation 5	
			Dry mouth 4	
			Fatigue 4	
			Headache 3	
			Dizziness 3	
			Insomnia 3	
			Nausea 3	
			Somnolence 2	
			Dyspepsia 2	
			Flatulence 2	

^{*}Author, year/(Procite number)/Study Design/Location/N=Total patients †age=mean±standard deviation in years BL=baseline latency; RCT=randomized controlled trial

Appendix 1-C. Evidence Tables: Sertraline treatment studies

Reference*	Study Population [†]	Treatment Regimen	Outcomes	Comments
Mendels, cont.		Placebo 50 mg/day titrated	Efficacy (min):	
		from 50 to 200 mg/day	Latency (n=22)	
		during weeks 1-3. Total 8	Baseline 1.10±1.35	
		weeks and 26 patients. Mean	Endpoint 1.85±3.68	
		final dosage 145 mg/day. 2-	Change 1.22±3.86	
		week washout titrating down	Ejaculation during foreplay	
		to 0 mg/day.	(n=24)	
			Baseline 4	
			During treatment 8	
			Patient satisfaction (n=24)	
			Baseline 1.83±1.20	
			Endpoint 1.88±1.39	
			Change 0.04±1.60	
			Partner satisfaction (n=20)	
			Baseline 1.65±1.14	
			Endpoint 2.10±1.37	
			Change 0.45±1.57	
			Side effects (n=26):	
			Any adverse evenť 16	
			Discont. due to AE 2	
			Diarrhea 1	
			Anejaculation 0	
			Dry mouth 2	
			Fatigue 0	
			Headache 3	
			Dizziness 1	
			Insomnia 1	
			Nausea 1	
			Somnolence 3	
			Dyspepsia 1	
			Flatulence 1	

^{*}Author, year/(Procite number)/Study Design/Location/N=Total patients †age=mean±standard deviation in years BL=baseline latency; RCT=randomized controlled trial

Appendix 1-C. Evidence Tables: Sertraline treatment studies

Reference*	Study Population [†]	Treatment Regimen	Outcomes	Comments
Waldinger, 2001 (795222) RCT Netherlands N=48	n=12: age 38±10 Married for 15±12 years with primary PE.	Placebo, morning and evening	Geometric mean latency by week (sec): 0 15.1 1 20.8 2 19.3 3 19.1 4 23.6 5 18.8 6 18.1	All patients had a 1-month baseline and were prohibited from using condoms or topical anesthetics through the study. Study was randomized and blinded. Patients used stopwatches to measure latency.
	n=12: age 40±7 Married for 17±7 years with primary PE.	Paroxetine 20 mg/day, morning and evening	Geometric mean latency by week (sec): 0 17.1 1 36.7 2 71.1 3 119.2 4 88.3 5 146.0 6 107.9	
	n=12: age 41±9 Married for 15±9 years with primary PE.	Sertraline 50 mg/day, morning and evening	Geometric mean latency by week (sec): 0 13.9 1 25.2 2 38.6 3 33.8 4 42.7 5 58.1 6 50.3	

^{*}Author, year/(Procite number)/Study Design/Location/N=Total patients †age=mean±standard deviation in years BL=baseline latency; RCT=randomized controlled trial

Appendix 1-C. Evidence Tables: Sertraline treatment studies

Reference*	Study Population [†]	Treatment Regimen	Outcomes	Comments
Waldinger, cont.	n=12: age 35±6 Married or in a relationship for 12±8 years with primary PE.	Nefazodone 400 mg/day, morning and evening	Geometric mean latency by week (sec) 0 16.8 1 19.1 2 14.2 3 25.6 4 28.4 5 15.6 6 17.9	
Waldinger, 1998 (12044a) RCT	After prescreening exclusions, 60 patients enrolled; 51 completed the study; 6 withdrew due to adverse events and 3 for lack of efficacy			This article describes two RCTs, the first of which is a 5-arm study. 6-week data are recorded here; graphs
Netherlands N=51	n=10: age 38±7, partner age 36±7 BL: 21±12 sec	Fluoxetine 20 mg/day	Latency at 6 weeks: 211±251 sec Absolute change: 189±244 sec	of intermediate results are presented in the paper.
	n=10: age 44±10, partner age 43±10 BL: 15±17 sec	Fluvoxamine 100 mg/day	Latency at 6 weeks: 55±70 sec Absolute change: 42±57 sec	
	n=11: age 41±8, partner age 39±8 BL: 16±10 sec	Paroxetine 20 mg/day	Latency at 6 weeks: 476±1146 sec Absolute change: 458±1142 sec	
	n=11: age 40±9, partner age 38±10 BL: 21±12 sec	Sertraline 50 mg/day	Latency at 6 weeks: 117±87 sec Absolute change: 96±84 sec	
	n=9: age 45±4, partner age 41±5 BL: 19±15 sec	Placebo	Latency at 6 weeks: 29±25 sec Absolute change: 10±18 sec	

^{*}Author, year/(Procite number)/Study Design/Location/N=Total patients †age=mean±standard deviation in years BL=baseline latency; RCT=randomized controlled trial

Appendix 1-C. Evidence Tables: Sertraline treatment studies

Reference*	Study Population [†]	Treatment Regimen	Outcomes	Comments
Wise, 1994 (12173) Letter/CR N=1	n=1: age 43 Patient with primary PE and no intercourse for 6 months due to wife's frustration with PE.	Sertraline 50 mg/day	Latency 6-10 min Patient and wife satisfied.	This letter is a case report about a single patient who refused sexual therapy (partially because wife refused).

^{*}Author, year/(Procite number)/Study Design/Location/N=Total patients †age=mean±standard deviation in years BL=baseline latency; RCT=randomized controlled trial

Appendix 1-D. Evidence Tables: Clomipramine treatment studies

Reference*	Study Population [†]	Treatment Regimen	Outcomes	Comments
Abdel-Hamid, 2001 (700014) Crossover RCT Egypt N=51	n=31: age 34.09±4.29 (range, 27-42) Heterosexual, married at least 1 year, with primary PE and willing to have intercourse twice a week. Patients did not have history of psychiatric illness, current physical illness, previous surgery or drug known to affect sexual function, current substance abuse, or ED. Patients each received each of 5 treatments for 4 weeks separated by 2 week wash-out periods. The treatments were ordered randomly in a double blind manner. BL (med.): 1 (0.5-1.5) min Baseline anxiety (med.): 12 (5-25)	Clomipramine 25 mg, 3-5 hrs before planned coitus and not more than twice/week	Results at 4 weeks: Median latency 4 (1-8) min Median anxiety score 11 (4-22) min Median sexual satisfaction score 11 (0-25) min 2 patients dropped out for lack of efficacy and 1 for side effects and lack of efficacy. Side effects: Dry mouth 3 Anorexia 0 Nausea 1 Headache 0 Flushing 0 Drowsiness 1 Sleepiness 2 Nasal congestion 0 Yawning 0 Total pts with side effects: 7/28	Study also included a normal control group of 20 patients who were not treated but supplied anxiety scores of 3.7±2.7(range 1-9) vs. 12.7±5.8 (range 5-25) for the PE subjects. Study contains blinding problems because the Masters and Johnson pause squeeze technique cannot be blinded.

^{*}Author, year/(Procite number)/Study Design/Location/N=Total patients †age=mean±standard deviation in years BL=baseline latency; RCT=randomized controlled trial

Appendix 1-D. Evidence Tables: Clomipramine treatment studies

Reference*	Study Population [†]	Treatment Regimen	Outcomes	Comments
Reference* Abdel-Hamid, cont.	Study Population [†]	Treatment Regimen Paroxetine 20 mg, 3-5 hrs before planned coitus and not more than twice/week	Results at 4 weeks: Median latency	Comments
			Sleepiness 0 Nasal congestion 0	
			Yawning 2 Total pts with side effects: 5/29	

^{*}Author, year/(Procite number)/Study Design/Location/N=Total patients †age=mean±standard deviation in years BL=baseline latency; RCT=randomized controlled trial

Appendix 1-D. Evidence Tables: Clomipramine treatment studies

Reference*	Study Population [†]	Treatment Regimen	Outcomes	Comments
Abdel-Hamid, cont.		Sertraline 50 mg, 3-5 hrs before planned coitus and not more than twice/week	Results at 4 weeks: Median latency	
			Total pts with side effects: 3/29	

^{*}Author, year/(Procite number)/Study Design/Location/N=Total patients †age=mean±standard deviation in years BL=baseline latency; RCT=randomized controlled trial

Appendix 1-D. Evidence Tables: Clomipramine treatment studies

Study Population [†]	Treatment Regimen	Outcomes	Comments
Study Population	Sildenafil 50 mg, 3-5 hrs before planned coitus and not more than twice/week	Results at 4 weeks: Median latency	Comments
	Masters and Johnson pause squeeze technique	Total pts with side effects: 5/28 Results at 4 weeks: Median latency 3 (1-7)	
		Median anxiety score 12 (5-21) Median sexual satisfaction score 6 (0-22)	
	Study Population [†]	Sildenafil 50 mg, 3-5 hrs before planned coitus and not more than twice/week Masters and Johnson pause	Sildenafil 50 mg, 3-5 hrs before planned coitus and not more than twice/week Sildenafil 50 mg, 3-5 hrs before planned coitus and not more than twice/week Results at 4 weeks: Median latency 15 (5-30) Median anxiety score 8 (4-15) Median sexual satisfaction score 30 (17-34) 2 patients dropped out for side effects. Side effects: Dry mouth 0 Anorexia 0 Nausea 0 Headache 2 Flushing 2 Drowsiness 0 Sleepiness 0 Nasal congestion 1 Yawning 0 Total pts with side effects: 5/28 Results at 4 weeks: Median latency 3 (1-7) Median anxiety score 12 (5-21) Median sexual satisfaction

^{*}Author, year/(Procite number)/Study Design/Location/N=Total patients †age=mean±standard deviation in years BL=baseline latency; RCT=randomized controlled trial

Appendix 1-D. Evidence Tables: Clomipramine treatment studies

Reference*	Study Population [†]	Treatment Regimen	Outcomes	Comments
Althof, 1995	n=15: age 38±8.94 (range, 23-56)	Each patient had a run-in	Latency:	Satisfaction scores are only
(12146)	All patients in stable relationships (married	period long enough for 3	Run-in	on a graph. Study also lists
Crossover RCT	or cohabitating) recruited through newspaper	attempts at coitus. 3	81 sec	data on a symptom
Cleveland	ads. All had primary PE with latency < 4 min	treatment periods were each	25mg/day clomipramine	checklist 90-R results.
N=15	and in good physical health without history of	set long enough to anticipate	202 sec	
	mental illness, not taking prescription meds,	5 attempts. The treatment	50mg/day clomipramine	
	ED less than 10% of time, and no substance	periods were separated by 1-	419 sec	
	or alcohol dependency in last 2 years.	week washout. At the end	Placebo (from graph)	
	BL: < 4 min	was a run-out equivalent to	137 sec	
		the run-in. 3 treatments	Run-out (from graph)	
		were used in random order:	80 sec	
		Clomipramine 25 mg/day		
		Clomipramine 50 mg/day	Side effects (days with effect):	
		Placebo 1/day	Placebo	
			Nausea 4	
			Claminramina 25 mg/day	
			Clomipramine 25 mg/day Dry mouth 7	
			Feeling different 8	
			Constipation 1	
			Consupation	
			Clomipramine 50 mg/day	
			Dry mouth 33	
			Feeling different 21	
			Constipation 18	
			Dizziness 10	
			Nausea 8	
			Sleep disturbance 6	
			Fatigue 4	
			Hot flashes 4	
			Headache 3	
			Ears ringing 3	
			Moist mouth 1	
			Decreased concentration 1	

^{*}Author, year/(Procite number)/Study Design/Location/N=Total patients †age=mean±standard deviation in years BL=baseline latency; RCT=randomized controlled trial

Appendix 1-D. Evidence Tables: Clomipramine treatment studies

Reference*	Study Population [†]	Treatment Regimen	Outcomes	Comments
Assalian, 1988 (105301) CR (5) Canada N=5	n=5: age 39.6 (range, 26-52) Patients were reported as individual case reports. 4 had primary PE and 1 had secondary.	All were started on clomipramine 25 mg/day. 1 patient partially complied by only taking the tablet when sexual activity was anticipated. Another patient took the dose only on Friday, Saturday, and Sunday when sex was possible. 1 patient reduced to 10 mg/day due to sleepiness.	All patients reported significant delay and heightened control over ejaculation. 2 patients had side effects including daytime sedation, dry mouth, and constipation.	No real data are supplied. Results and side effect data are not clear in this study.
Eaton, 1973 (105303) CS England N=13	n=13: age 35 (range, 19-58) 11 in stable relationships, 2 with multiple partners. 11 with primary PE (but 1 had a period of normal response in the past). 7 had some ED and 5 had some loss of libido. Most had some amount of anxiety and/or depression.	Clomipramine 25 mg/day titrated up to 75 mg/day in 25 mg steps every 2 weeks	12/13 responded "positively" in 2 weeks to 2 months. Side effects included dyspepsia, dry mouth and perspiration.	No numbers for side effects or definition of PE or positive response given.
Girgis, 1982 (12409) Crossover RCT Egypt N=50	n=50: age 34.5±8.2 (range, 19-57) Married with history of PE from 1 months to 17 years. Patients in good health with normal libido and erection, no diabetes, pyuria or prostatitis.	Clomipramine 10 mg bid (12.00 and 18.00 hrs) for 6 weeks followed by placebo bid for 6 weeks	n=22 (Initially 25 patients but 3 dropouts.) Number of satisfactory sexual performances: Clomipramine 161/317 Placebo 163/313 No side effects observed.	Dropouts were for no apparent reason. There was no washout.

^{*}Author, year/(Procite number)/Study Design/Location/N=Total patients †age=mean±standard deviation in years BL=baseline latency; RCT=randomized controlled trial

Appendix 1-D. Evidence Tables: Clomipramine treatment studies

Reference*	Study Population [†]	Treatment Regimen	Outcomes	Comments
Girgis, cont.		Pacebo bid (12.00 and 18.00 hrs) for 6 weeks followed by clomipramine 10 mg bid for 6 weeks.	n=17 (Initially 25 patients but 8 dropouts.) Number of satisfactory sexual performances: Clomipramine 106/216 Placebo 73/228 No side effects observed.	
Goodman, 1980 (13710) Crossover RCT England N=20	n=10: age 28 (range, 18-43) 4 with primary PE and 3 with secondary PE. 2 with occasional ED. All patients offered or trained in squeeze technique without success. 3 withdrawals from study – net 7 patients.	Clomipramine 10 mg/day increasing up to 40 mg/day for 1 month. Second month clomipramine at doses up to 150 mg/day. At 4 weeks 2 8 weeks 3/7 16 weeks 1 further treatr better using	At 4 weeks 2/7 were better. At 8 weeks 3/7 were better. At 16 weeks 1 was cured (no further treatment needed), 2 better using clomipramine occasionally and the rest not	This study is difficult to decode. There are no numeric outcomes, only subjective terms like better or no better. Side effects are listed in a divided table that makes it unclear which drug or group they apply to.
	n=10: age 30 (range, 22-41) 7 with primary PE and 2 with secondary PE. 3 with occasional ED. All patients offered or trained in squeeze technique without success. 1 withdrawal from study – net 9 patients.	Placebo 10 mg/day increasing up to 40 mg/day for 1 month. Second month clomipramine at doses up to 150 mg/day.	At 4 weeks 4/9 were better. At 8 weeks 5/9 were better. At 16 weeks 6 were better (2 using clomipramine occasionally, 4 using it regularly), 2 were better but stopped using clomipramine due to side effects and 1 was not better.	

^{*}Author, year/(Procite number)/Study Design/Location/N=Total patients †age=mean±standard deviation in years BL=baseline latency; RCT=randomized controlled trial

Appendix 1-D. Evidence Tables: Clomipramine treatment studies

Reference*	Study Population [†]	Treatment Regimen	Outcomes	Comments
Haensel, 1996	n=8: age 42.2 (range, 31-62)	Patients in all three groups	Latency: (n=6)	There is no mention of this
(10696)	Patients with SMD-IV criteria for primary	were given two treatments in	Pretest 1.4±.3	being a double-blind trial.
Crossover RCT	premature ejaculation, heterosexual, with	crossover. The treatments	Clomipramine 2.8±.6	The lack of washout and
Netherlands	dysfunction for at least 6 months, willingness	were 25 mg of clomipramine	Placebo 1.7±.3	the fact that more patients
N=22	to attempt coitus or masturbation at least	12-24 hrs before anticipated		began with clomipramine
	once a week, no concomitant psychiatric	sexual activity (coitus or	Approximately 2 min placebo,	may have impacted results.
	disease, previous surgery or drug use that	masturbation) and placebo.	8 min clomipramine:	A variety of other scores
	would affect sexual function.	Patients received each	Number of thrusts (n=6)	and frequencies were also
		treatment for 3 weeks and	Pretest 1.9±0.4	listed in the article.
		were crossed over to the	Clomipramine 4.0±0.0	Latency and number of
		alternate treatment for 3	Placebo 3.3±0.5	thrusts were scored on
		weeks. Patients were	Orgasm sooner than desired	unusual scales:
		randomized as to order with	(n=8)	
		13 beginning with	Pretest 1.0±0.0	Latency(min)
		clomipramine and 9 with	Clomipramine 4.0±0.8	Score
		placebo.	Placebo 1.1±0.1	<1 1
	n=6: age 41.2 (range, 26-52)		Latency: (n=5)	1-3 2
	Patients with SMD-IV criteria for secondary		Pretest 1.0±0.0	4-6 3
	premature ejaculation who also had erectile		Clomipramine 1.2±0.2	7-10 4
	dysfunction. These patients were		Placebo 1.2±0.2	11-15 5
	heterosexual, with dysfunction for at least 6			>15 6
	months, willingness to attempt coitus or		Approximately 2 min with	
	masturbation at least once a week, no		either treatment:	Number of thrusts
	concomitant psychiatric disease, previous		Number of thrusts (n=5)	Score
	surgery or drug use that would affect sexual		Pretest 1.5±0.5	<5 1
	function.		Clomipramine 1.8±0.4	6-10 2 11-20 3
			Placebo 1.5±0.4	
			Orgasm sooner than desired	>20 4
			(n=6)	
			Pretest 1.2±0.2	Orgasm sooner than
			Clomipramine 1.8±0.5	desired was scored on a 1
			Placebo 1.8±0.8	– 7 scale

^{*}Author, year/(Procite number)/Study Design/Location/N=Total patients †age=mean±standard deviation in years BL=baseline latency; RCT=randomized controlled trial

Appendix 1-D. Evidence Tables: Clomipramine treatment studies

Reference*	Study Population [†]	Treatment Regimen	Outcomes	Comments
Haensel, cont.	n=8: age 40.5 (range, 26-52) recruited		Latency: (n=5)	1 – always, 7 – never
	through word of mouth healthy normal controls		Pretest 3.7±0.3 Clomipramine 3.6±0.2	Side effects, not broken
			Placebo 3.8±0.2	down by group, included:
			Approx. 9 min placebo and 11	Fatigue or low energy 8
			min clomipramine:	Dizziness 3
			Number of thrusts (n=6) Pretest 3.7±0.3	Nausea, headache, yawning 1
			Clomipramine 3.7±0.3	
			Placebo 4.0±0.0 Orgasm sooner than desired	9/14 patients discriminated drug from placebo
			(n=7) Pretest 5.3±0.6	
			Clomipramine 5.6±0.6	
			Placebo 5.7±0.5	
Kim, 1998 (12047)	n=37: age 44 (range; 30-60). 53 heterosexual patients enrolled; 37	Baseline – no treatment	Latency: 46±41 sec	All patients received each therapy and placebo for 4
Crossover RCT	completed the study.		Patient Satisfaction:	weeks with a 4-week
Korea	Reasons for withdrawal: loss to follow-up (5),		Satisfied 0%	washout between
N=53	no efficacy (fluoxetine 3, sertraline 1,		Moderate 0%	therapies. Order of
	placebo 1) and no efficacy plus side effects (clomipramine 1). One patient was excluded		Dissatisfied 100%	administration was randomized.
	from analysis because of delayed (>30 min)		Partner Satisfaction:	
	ejaculation with sertraline or clomipramine.		Satisfied 0%	
			Moderate 22.2%	
			Dissatisfied 77.8%	

^{*}Author, year/(Procite number)/Study Design/Location/N=Total patients †age=mean±standard deviation in years BL=baseline latency; RCT=randomized controlled trial

Appendix 1-D. Evidence Tables: Clomipramine treatment studies

Reference*	Study Population [†]	Treatment Regimen	Outcomes	Comments
Kim, cont.		Placebo 1 capsule/day first week and 2/day thereafter	Latency: 2.27±3.78 min Patient Satisfaction: Satisfied 19.4% Moderate 27.8% Dissatisfied 52.8% Partner Satisfaction: Satisfied 11.1% Moderate 36.1% Dissatisfied 52.8% Side effects: Drowsiness 2 Dry mouth 0 Reduced Potency 3 Nausea 1 Vomiting 0 Other 3	
			Total pts with side effects: 7	

^{*}Author, year/(Procite number)/Study Design/Location/N=Total patients †age=mean±standard deviation in years BL=baseline latency; RCT=randomized controlled trial

Appendix 1-D. Evidence Tables: Clomipramine treatment studies

Reference*	Study Population [†]	Treatment Regimen	Outcomes	Comments
Kim, cont.		Clomipramine 25 mg/day for first week, 50 mg/day thereafter	Latency: 5.75±6.68 min Patient Satisfaction: Satisfied 25.0% Moderate 38.9% Dissatisfied 36.1% Partner Satisfaction: Satisfied 19.4% Moderate 38.9% Dissatisfied 41.7% Side effects: Drowsiness 6 Dry mouth 2 Reduced Potency 3 Nausea 4 Vomiting 0 Other 3 Total pts with side effects: 13	
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^{*}Author, year/(Procite number)/Study Design/Location/N=Total patients †age=mean±standard deviation in years BL=baseline latency; RCT=randomized controlled trial

Appendix 1-D. Evidence Tables: Clomipramine treatment studies

Reference*	Study Population [†]	Treatment Regimen	Outcomes	Comments
Kim, cont.		Fluoxetine 20 mg/day for first week, 40 mg/day thereafter	Latency: 2.30±2.08 min Patient Satisfaction: Satisfied 25.0% Moderate 38.9% Dissatisfied 36.1% Partner Satisfaction: Satisfied 19.4% Moderate 38.9% Dissatisfied 41.7% Side effects: Drowsiness 6 Dry mouth 2 Reduced Potency 3 Nausea 4 Vomiting 0 Other 3	
			Total pts with side effects: 13	

^{*}Author, year/(Procite number)/Study Design/Location/N=Total patients †age=mean±standard deviation in years BL=baseline latency; RCT=randomized controlled trial

Appendix 1-D. Evidence Tables: Clomipramine treatment studies

Reference*	Study Population [†]	Treatment Regimen	Outcomes	Comments
Kim, cont.		Sertraline 50 mg/day for first	Latency: 4.27±5.69 min	
		week, 100 mg/day thereafter	5 " 10 " 1 "	
			Patient Satisfaction:	
			Satisfied 41.7%	
			Moderate 36.1%	
			Dissatisfied 22.2%	
			Partner Satisfaction:	
			Satisfied 30.6%	
			Moderate 38.9%	
			Dissatisfied 30.6%	
			Side effects:	
			Drowsiness 7	
			Dry mouth 4	
			Reduced Potency 3	
			Nausea 2	
			Vomitting 0	
			Other 0	
			Total pts with side effects: 12	
Montorsi, 1995	N=17 with available partner, without ED,	Placebo for 8 weeks followed	Placebo response	Dropouts due to side
(500007)	penile defects with PE at least 50% of time	by clomipramine 50 mg/day	Complete 1/17	effects: tremor in 5, nausea
RCT, Italy	·	given at bedtime for 8 weeks	Partial 2/17	in 2 – not clear which
N=40			Failure 14/17	group. Overall
			Overall complication rate 10%	clomipramine complication
				rate (both groups) is 40%.
			Clomipramine response	The 3 month data is with all
			Complete 5/17	patients taking
			Partial 6/17	clomipramine.
			Failure 6/17	

^{*}Author, year/(Procite number)/Study Design/Location/N=Total patients †age=mean±standard deviation in years BL=baseline latency; RCT=randomized controlled trial

Appendix 1-D. Evidence Tables: Clomipramine treatment studies

Reference*	Study Population [†]	Treatment Regimen	Outcomes	Comments
Montorsi, cont.	N=16 with available partner, without ED, penile defects with PE at least 50% of time	Clomipramine 50 mg/day given at bedtime for 8 weeks (presumably continued for at least 3 months)	Clomipramine response Complete 4/16 Partial 4/16 Failure 8/16	Complete 8/33 Partial 9/33 Failure 16/33 Complete response is ability to control ejaculation at least 90% of the time. Partial response is controlled at least 70% of the time.
Rowland, 2001 (795018) CS US N=4	n=4: age 53.5 (range, 44-72) Out of 13 who failed with 25 mg. Clomipramine PRN who were in stable relationships. Other 9 had either moved out or were no longer in stable relationships with a willing partner. PE persisted for 7.5 (3-10) years with BL: ≤1 min	Patients received increasing daily doses of clomipramine of 10, 20, and 30 mg/day for 3 weeks for each dose. Data were also reported from a previous study with these patients where they received 25 mg PRN or placebo.	Latency in sec (mean): Baseline 25.5 10 mg 50.65 (median from graph) 20 mg 91.5 30 mg 221.0 Ejaculatory control (0-10 increasing scale): Baseline 4.0 10mg 4.5 20mg 5.9 30mg 6.2	This is a very small study that shows that daily dosing may salvage some patients who fail PRN dosing. Patients were failures from article 12022.

^{*}Author, year/(Procite number)/Study Design/Location/N=Total patients †age=mean±standard deviation in years BL=baseline latency; RCT=randomized controlled trial

Appendix 1-D. Evidence Tables: Clomipramine treatment studies

Reference*	Study Population [†]	Treatment Regimen	Outcomes	Comments
Segraves, 1993 (12219) RCT Czech. Rep. N=29	n=10: age 44.7±7.5 Recruited by newspaper ad. Requirements include age 21-65, and no history of psychiatric disorder, current substance abuse, significant illness, or use of psychotropic drugs. 8 were married and 7 had primary PE.	25 mg clomipramine to take 6 hrs prior to coitus. After 2 attempts patients could double dose in the absence of side effects or unsatisfactory ejaculatory delay. Trial continued for a total 10 attempts at coitus.	At 25 mg: 7/10 had latency ≥ 2 min Mean at 25 mg, 6.1 min At 50 mg: 8/10 had latency ≥ 2 min 6/7 men increased latency by 2 min or more by spousal estimate. Rating of libido, erections, ejaculation timing and quality, and overall satisfaction significantly increased. Side effects: Mild nausea 3 Drowsiness 3 Diarrhea 1 Transient ED 2	9 patients (out of 29) dropped out and were not included in the analysis, but it is not clear which groups they were in. Only data for completers was supplied. Numbers lacking for some outcomes.

^{*}Author, year/(Procite number)/Study Design/Location/N=Total patients †age=mean±standard deviation in years BL=baseline latency; RCT=randomized controlled trial

Appendix 1-D. Evidence Tables: Clomipramine treatment studies

Reference*	Study Population [†]	Treatment Regimen	Outcomes	Comments
Segraves, cont.	n=10: age 47.2±10.2 Recruited by newspaper ad. Requirements include age 21-65, and no history of psychiatric disorder, current substance abuse, significant illness, or use of psychotropic drugs. 6 were married and 8 had primary PE.	25 mg placebo to take 6 hrs prior to coitus. After two attempts patients could double dose in the absence of side effects or unsatisfactory ejaculatory delay. Trial continued for a total 10 attempts at coitus.	At 25mg: 0 men had latency ≥ 2 min Mean at 25mg, 51 sec At 50 mg: 2 men had latency ≥ 2 min 2/5 men increased latency by 2 min or more by spousal estimate. Rating of libido, erections, ejaculation timing and quality, and overall satisfaction were unchanged. Side effects: Mild paymen	
Strassberg, 1999 (12022) Crossover RCT Salt Lake City N=34	n=23: age 46.3±11.738 PE patients recruited from newspaper ads of which 28 completed the study. 5 of these were dropped because the studies indicated placebo latencies greater than 3 min Patients had no history of psychiatric disorder, no current substance abuse, no current relevant medications, no previous surgery or drug use known to impact sexual function, and in a stable heterosexual relationship.	Patients each received 2 treatments for 2 weeks with no apparent washout. Treatment 1: 25 mg (2 capsules) of clomipramine taken 4 hrs before coitus. Treatment 2: 2 capsules of placebo taken similarly. Patients were randomized to sequence.	Mild nausea 1 Drowsiness 1 Constipation 2 Latency: (n=22) Placebo 52±45 sec Clomipramine 229±286 sec Control over ejaculation: (n=16) (1-10 scale) Placebo 2.70±1.84 Clomipramine 5.08±2.45	This article contains many complex statistical analyses, showing that clomipramine increased latency and control for the patients with PE. The study included lab studies with arousing videos and vibrators, but only the at home diary data is included under outcomes here.

^{*}Author, year/(Procite number)/Study Design/Location/N=Total patients †age=mean±standard deviation in years BL=baseline latency; RCT=randomized controlled trial

Appendix 1-D. Evidence Tables: Clomipramine treatment studies

Reference*	Study Population [†]	Treatment Regimen	Outcomes	Comments
Strassberg,	n=13: age 45.6±7.3		Latency:	
cont.	15 normal control patients recruited from		(n=4)	
	newspaper ads of which 13 completed		Placebo 491±245 sec	
	study. 2 of these were dropped because the		Clomipramine 665±307 sec	
	studies indicated placebo latencies less than			
	150 sec		Control over ejaculation:	
	Patients had no history of psychiatric		(n=4)	
	disorder, no current substance abuse, no		(1-10 scale)	
	current relevant medications, no previous		Placebo 8.25±1.5	
	surgery or drug use known to impact sexual		Clomipramine 8.5±1.0	
	function, and in a stable heterosexual			
	relationship.			

^{*}Author, year/(Procite number)/Study Design/Location/N=Total patients †age=mean±standard deviation in years BL=baseline latency; RCT=randomized controlled trial

Appendix 1-E. Evidence Tables: Topical anesthetic treatment studies

Reference*	Study Population [†]	Treatment Regimen	Outcomes		Comments
Atan, 2000	n=26: age 27 (range, 21-36)	Fluoxetine 20 mg/day for 1	Cured:	8 (30.8%)	There is no indication of
(795257)	With PE and without ED, substance abuse,	week followed by 40 mg/day	Improved:	11 (42.3%)	randomization in the article.
Controlled Trial Turkey	infections, diabetes, thyroid disease, hypotension or loss of libido. Patients had	for 7 weeks	Failure:	7 (26.9%)	
N=43	normal psychiatric consultations.		Side effects:		
			Nausea	3	
			Headache	1	
			Insomnia	2	
			Total patients w	ith side effects:	
				6	
	n=17: age 31 (range,19-48)	Fluoxetine 20 mg/day and	Cured:	9 (52.9%)	
	With PE and without ED, substance abuse,	local application of lidocaine	Improved:	5 (29.4%)	
	infections, diabetes, thyroid disease, hypo tension or loss of libido. Patients had normal	ointment to the glans 20 min prior to intercourse	Failure:	3 (17.6%)	
	psychiatric consultations.	•	Side effects:		
			Nausea	1	
			Headache	4	
			Insomnia	0	
			Total patients w	ith side effects:	
				5	

^{*}Author, year/(Procite number)/Study Design/Location/N=Total patients
†age=mean±standard deviation in years
BL=baseline latency; RCT=randomized controlled trial

Appendix 1-E. Evidence Tables: Topical anesthetic treatment studies

Reference*	Study Population [†]	Treatment Regimen	Outcomes	Comments
Atikeler, 2002 (500006) RCT Turkey	N=10, age 29.4 (range, 20-40) with latency less than 1 minute and married or with constant partner for more than 2 years and no organic causes.	Placebo cream used at least 5 times (range 5-8)	Mean latency 1.01 min	
N=40	N=10, age 29.4 (range, 20-40) with latency less than 1 minute and married or with constant partner for more than 2 years and no organic causes.	Prilocaine-lidocaine cream applied and left in place under a condom for 20 min and removed—used at least 5 times (range 5-9)	Mean latency 6.71 min in 8 patients. One patient still had 1 min latency and one patient increased to 3-5 min. Overall average latency 6.5 min	
	N=10, age 29.4 (range, 20-40) with latency less than 1 minute and married or with constant partner for more than 2 years and no organic causes.	Prilocaine-lidocaine cream applied and left in place under a condom for 30 min and removed—used at least 5 times (range 5-9)	Mean latency 8.71 min in 4 patients while 6 patients complained of erection loss due to numbness and delayed ejaculation	
	N=10, age 29.4 (range, 20-40) with latency less than 1 minute and married or with constant partner for more than 2 years and no organic causes.	Prilocaine-lidocaine cream applied and left in place under a condom for 45 min and removed—used at least 5 times (range 5-9)	All patients complained of erection loss due to numbness and delayed ejaculation	

^{*}Author, year/(Procite number)/Study Design/Location/N=Total patients
†age=mean±standard deviation in years
BL=baseline latency; RCT=randomized controlled trial

Appendix 1-E. Evidence Tables: Topical anesthetic treatment studies

Reference*	Study Population [†]	Treatment Regimen	Outcomes	Comments
Berkovitch,	n=11: age 36±7 (range, 26-44)	Lidocaine-prilocaine cream	Patients estimated latency and	
1995	All patients heterosexual and married or in	2.5 mg dose applied to glans	graded it as follows:	
(12140)	stable relationship without ED or risk factors	and shaft and covered with a	Excellent 5	
CS	for ED or extensive drug or alcohol use.	condom for 30 min After that	Better than usual 4	
Canada N=11		time the condom could be removed at the patient's	Unsatisfied 2	
		option.	Patients rating excellent used	
			the cream 8.4±1.7 times with latency 15-20 min	
			Patients rating better used the cream 5±0.41 time with latency 5-10 min	
			One unsatisfied patient complained of numbness despite 20 min latency and spousal satisfaction. The other unsatisfied patient had no meaningful improvement.	
Damrau, 1963	n=13: age 31.2 (range, 22-39)	Ethyl aminobenzoate cream	PE "corrected" in all cases.	Study also reports some
(105302)	11 with ejaculation prior to intromission, 1	(3%) to glans and prepuce,	Latency 1.6 (range, 0.5-5).	data on the use of the
CS	with immediate ejaculation and 1 with 1 min	wiped off after 5 min	Duration of anesthesia mean	cream on normal
New Jersey N=13	latency. Duration of condition 2.7 years (range, 0.5-5).		25.2 min (range, 10-50).	volunteers.

^{*}Author, year/(Procite number)/Study Design/Location/N=Total patients
†age=mean±standard deviation in years
BL=baseline latency; RCT=randomized controlled trial

Appendix 1-E. Evidence Tables: Topical anesthetic treatment studies

Reference*	Study Population [†]	Treatment Regimen	Outcomes	Comments
Slob, 2000	N=15; age 43±11(range, 21-64)	Lidocaine-prilocaine cream	Latency	Patients were own controls
(500003)	Estimated latency between 1-2 min with less	(1/2 tube) applied for 10 min	489±465	in an unblended crossover.
CS	than 8 thrusts with ejaculation prior to	contained with condom, then	Sense of control	
Netherlands	intromission 31% of the time. 11/15 patients	removed. Patients given 5	4.7±3.1	
N=15	experienced ejaculation prior to intromission	tubes and 12 condoms.	3 pts reported inability to reach	
	at some times. Full erections 92% of the		orgasm on some occasions	
	time.		with the cream	
		Coitus without cream or	Latency	
		intervention. Patients asked	115±142	
		to report on 2-3 encounters.	Sense of control	
			2.5±2.2	
			No patients reported inability	
			to achieve orgasm	

^{*}Author, year/(Procite number)/Study Design/Location/N=Total patients †age=mean±standard deviation in years BL=baseline latency; RCT=randomized controlled trial

Appendix 1-F. Evidence Tables: Adrenergic blockade treatment studies

Reference*	Study Population [†]	Treatment Regimen	Outcomes	Comments
Beretta, 1986 (12350) CS Italy N=15	n=15: age 34 (range, 20-55) All patients with primary PE who refused psychological treatment (at least initially).	Phenoxybenzamine 10 mg/day	Outcomes given by patient. 7 patients ejaculated ante portam before treatment but only 4 did so after treatment. The other 3 had an average latency of 15 min after treatment. Average pretreatment latency of 8 patients who ejaculated intravaginally pretreatment was 1.38 min pretreatment and 5.63 min post-treatment. Of these 8 patients, 3 had no change in latency. Patients report of sexual response: Improved 8 No variation 5 Worsened 2 Partner satisfaction: Improved 8 No variation 5 Worsened 2 Two patients reported dry ejaculation	Also data on semen volume.

^{*}Author, year/(Procite number)/Study Design/Location/N=Total patients †age=mean±standard deviation in years BL=baseline latency; RCT=randomized controlled trial

Appendix 1-F. Evidence Tables: Adrenergic blockade treatment studies

Reference*	Study Population [†]	Treatment Regimen	Outcomes	Comments
Cavallini, 1995 (12160) Crossover RCT Italy N=101	n=101: age 34.3 (range, 21-63)	Patients received alphuzosine (2x3 mg/day, terazosine (1 mg/day up to 5 mg once a day), vitamin C 2x500 mg/day (placebo). The patients were in 3 groups which received treatment in 1 of the following 3 orders: 1. Alphuzosine, terazosine, placebo 2. Terazosine, placebo, alphuzosine 3. Placebo, alphuzosine, terazosine, terazosine	Results deemed positive if patients and spouses agreed they were satisfied. Alphuzosine: 1st administered 14/30 2nd administered 16/31 3rd administered 13/30 Terazosine: 1st administered 15/30 2nd administered 15/30 2nd administered 16/30 3rd administered 16/30 3rd administered 17/31 Placebo: 1st administered 4/31 2nd administered 4/31 2nd administered 8/30 Side effects: Alphuzosin: 5/90 (3 hypotension, 2 withdrawals, 1 hypotension plus epigastralgia) Terazosine: 3/90 (2 hypotension leading to withdrawal, 1 headache) Placebo: 1/90 (weak epigastralgia)	No washout periods between treatments and terazosine followed by alphuzosine directly twice, but the reverse never happened. Only 3 of 6 possible orders were used. 6 patients were excluded from analysis because patients and spouses disagreed about results. It is not clear which groups they came from. Dosages did not match (terazosine 1/day, others bid) so blinding was compromised. Side effect numbers do not add up and refer to withdrawals at various stages in the study. It is not clear how this was handled in the results statistics.

^{*}Author, year/(Procite number)/Study Design/Location/N=Total patients †age=mean±standard deviation in years BL=baseline latency; RCT=randomized controlled trial

Appendix 1-F. Evidence Tables: Adrenergic blockade treatment studies

Reference*	Study Population [†]	Treatment Regimen	Outcomes		Comments
Cooper, 1984	n=12: age 28 (range, 18-58)	Patients each had a 4 week	Latency, mean (min)):	Also data about erectile
(12387)	Patients with PE for mean 4.2 years (range,	run-in followed by a 4 week	Run-in	1.5	quality, blood pressure,
Crossover RCT	1-12.5).	treatment (either propranolol	Placebo	1.7	pulse rate, Hamilton rating
Canada	Patients also had chronic anxiety for at least	120 mg/day or placebo), 4	Wash-out	1.5	scale. No real impact in
N=12	6 months scoring > 9 on the somatic scale of	week washout, 4 week	Propranolol	1.7	any case. Not all subjects
	the Hamilton Anxiety Rating Scale.	treatment with the alternate	Run-out	1.6	were able to meet the
		treatment, and a 4 week run-			projected goal of
		out. Patients were randomly	Overall satisfaction,	mean	intercourse 2x per week.
		assigned to treatment	(0-100 scale):		
		sequence.	Run-in	51.7	
			Placebo	51.7	
			Wash-out	50.3	
			Propranolol	52.5	
			Run-out	52.2	

^{*}Author, year/(Procite number)/Study Design/Location/N=Total patients †age=mean±standard deviation in years BL=baseline latency; RCT=randomized controlled trial

Appendix 1-F. Evidence Tables: Adrenergic blockade treatment studies

Reference*	Study Population [†]	Treatment Regimen	Outcomes		Comments
Shilon, 1984	n=9: age 36.3±7.7 (range, 28-50)	Patients were treated with	Premature Ejaculation		Erection quality:
(12371)	All patients with undefined premature	phenoxybenzamine.	Baseline:	9/9	2 patients with fair erection
CS	ejaculation.		10 mg :		at baseline progressed to
Israel		Patients received increasing	No improvement	5/9	good erections on any drug
N=9		doses of 10, 20, and 30	Slight improvement	2/9	dose. One patient with
		mg/day, but the study is not	Good	2/9	good erections at baseline
		clear whether there was a	20 mg:		had fair erections on 10
		washout period, or even the	No improvement	1/7	mg/day and 20 mg/day and
		sequence of doses.	Slight improvement	1/7	good erections at 30
		Presumably the doses were	Good	4/7	mg/day.
		escalated based on results.	Excellent	1/7	
		Duration of treatment ranged			Most other patients had
		from 7 to 150 days. All	Wife's response:		significant reductions in
		patients received 10 mg. 7	Baseline:		semen volume.
		also received 20 mg and 3 of	Unsatisfied	7/9	
		those received 30 mg.	Satisfied	2/9	
			10 mg.:		
			Unsatisfied	2/9	
			No improvement	5/9	
			Slight improvement	1/9	
			Good	1/9	
			20 mg.:		
			No improvement	2/7	
			Satisfied	4/7	
			Good	1/7	
			30 mg.:	0.40	
			Satisfied	2/2	
			Zero semen volume		
			10 mg	4/9	
			20 mg	6/7	
			30 mg	2/2	

^{*}Author, year/(Procite number)/Study Design/Location/N=Total patients †age=mean±standard deviation in years BL=baseline latency; RCT=randomized controlled trial

Appendix 1-G. Evidence Tables: Miscellaneous treatment studies

Reference*	Study Population [†]	Treatment Regimen	Outcomes	Comments
Brown, 2000 (105252) CR	n=1: age 31 Patient with acute prostatitis who also had unreported PE for three years.	Ciprofloxacin 500 mg bid for 30 days (for prostatitis)	After treatment: Latency 6-15	Individual patient report of an incidental finding after ciprofloxacin therapy.
New York N=1	BL: 1-2 minutes.		Increase persisted after treatment discontinued.	
Fein, 1990 (900015) CS Florida N=16	n=8 Complaining of life-long PE, ejaculating prior to or immediately on vaginal penetration. Reasons for withdrawal: 8 patients declined to participate in the study due to an unwillingness to have penile injections.	Penile injection with 0.2 ml of papaverine 30 mg/ml and phentolamine 1mg/ml	Success was defined as having an erection of sufficient length for satisfactory intercourse regardless of ejaculation. All patients were successful with doses ranging from 0.1 to 0.4 ml. Three patients said they were cured and having successful intercourse with no PE. Other 5 continuing injections. No side effects reported in 14 months of treatment.	All 16 patients trained in Masters and Johnson squeeze technique, gluteal muscle contraction technique, mental divergence and 15 min masturbation exercise with no success.

^{*}Author, year/(Procite number)/Study Design/Location/N=Total patients †age=mean±standard deviation in years BL=baseline latency; RCT=randomized controlled trial

Clomipramine

Reference	Description/Comments	No. of AE's	No. of Patients	Rate
Anorexia 700014	25 mg 3-5 hrs before coitus, not > 2 x per week	0	31	0.0%
Diarrhea 12219	25 mg 6 hrs prior to coitus	1	10	10.0%
Dizziness	OF and 40 04 has a size to convert activity	2	00	42.00/
10696 Drowsiness	25 mg 12-24 hrs prior to sexual activity	3	22	13.6%
12047 12219 700014	25 mg/day for 1 week, 50 mg/day later 25 mg 6 hrs prior to coitus 25 mg 3-5 hrs before coitus, not > 2 x per week	8 3 1	53 10 31	15.1% 30.0% 3.2%
Dry mouth 10696 12047 700014	25 mg 12-24 hrs prior to sexual activity 25 mg/day for 1 week, 50 mg/day later 25 mg 3-5 hrs before coitus, not > 2 x per week	4 12 3	22 53 31	18.2% 22.6% 9.7%
Erectile dysfunction - transient 12219	25 mg 6 hrs prior to coitus	2	10	20.0%
Fatigue or low energy 10696	25 mg 12-24 hrs prior to sexual activity	8	22	36.4%
Flushing 700014	25 mg 3-5 hrs before coitus, not > 2 x per week	0	31	0.0%
Headache 700014	25 mg 3-5 hrs before coitus, not > 2 x per week	0	31	0.0%
Nasal congestion 700014	25 mg 3-5 hrs before coitus, not > 2 x per week	0	31	0.0%
Nausea 12047 500007 700014	25 mg/day for 1 week, 50 mg/day later 50 mg/day given at bedtime for 8 weeks 25 mg 3-5 hrs before coitus, not > 2 x per week	3 2 1	53 33 31	5.7% 6.1% 3.2%
Nausea - mild 12219	25 mg 6 hrs prior to coitus	3	10	30.0%
Nausea, headache, yawning 10696	25 mg 12-24 hrs prior to sexual activity	1	22	4.5%
Other 12047	25 mg/day for 1 week, 50 mg/day later	2	53	3.8%
Potency - reduced 12047	25 mg/day for 1 week, 50 mg/day later	5	53	9.4%
Sleepiness 700014	25 mg 3-5 hrs before coitus, not > 2 x per week	2	31	6.5%
Tremor 500007	50 mg/day given at bedtime for 8 weeks	5	33	15.1%
Vomiting 12047	25 mg/day for 1 week, 50 mg/day later	2	53	3.8%
Yawning 700014	25 mg 3-5 hrs before coitus, not > 2 x per week	0	31	0.0%

Fluoxetine

Reference	Description/Comments	No. of AE's	No. of Patients	Rate
Altered Sleep				
900017	5 mg/day for 2 weeks, 10 mg/day for 2 weeks	2	40	5.0%
Dizziness				
12112	20 mg/day for 1-2 weeks, titrated to 60 mg/day	1	11	9.1%
900017	5 mg/day for 2 weeks, 10 mg/day for 2 weeks	1	40	2.5%
Drowsiness				
12047	20 mg/day for 1 week, 40 mg/day later	6	53	11.3%
Dry mouth				
12047	20 mg/day for 1 week, 40 mg/day later	2	53	3.8%
900017	5 mg/day for 2 weeks, 10 mg/day for 2 weeks	3	40	7.5%
Extremity tingling				
12112	20 mg/day for 1-2 weeks, titrated to 60 mg/day	1	11	9.1%
Headache				
12003 12110	20 mg/day for 1 week, 40 mg/day later 20 mg/day for 1 week, 40 mg/day later	1 1	26 9	3.8% 11.1%
795257	20mg/day for 1 week, 40 mg/day later 20mg/day for 1 week then 40mg/day	1	26	3.8%
Insomnia				
12003	20 mg/day for 1 week, 40 mg/day later	2	26	7.7%
12110	20 mg/day for 1 week, 40 mg/day later	1	9	11.1%
795257	20mg/day for 1 week then 40mg/day	2	26	7.7%
Libido - increased				
900017	5 mg/day for 2 weeks, 10 mg/day for 2 weeks	2	40	5.0%
Loose stools				
900017	5 mg/day for 2 weeks, 10 mg/day for 2 weeks	2	40	5.0%
Nausea				
12003	20 mg/day for 1 week, 40 mg/day later	3	26	11.5%
12047 12110	20 mg/day for 1 week, 40 mg/day later 20 mg/day for 1 week, 40 mg/day later	4 2	53 9	7.5% 22.2%
795257	20mg/day for 1 week then 40mg/day	3	26	11.5%
Nausea and other GI				
12112	20 mg/day for 1-2 weeks, titrated to 60 mg/day	4	11	36.4%
Other				
12047	20 mg/day for 1 week, 40 mg/day later	3	53	5.7%
Palpitations - slight				
900017	5 mg/day for 2 weeks, 10 mg/day for 2 weeks	1	40	2.5%
Penile pain				
900017	5 mg/day for 2 weeks, 10 mg/day for 2 weeks	1	40	2.5%
Potency - reduced	3,			
12047	20 mg/day for 1 week, 40 mg/day later	3	53	5.7%
Sweating	. 3	· ·		2 ,3
900017	5 mg/day for 2 weeks, 10 mg/day for 2 weeks	1	40	2.5%
	o my day for 2 woods, to my day for 2 woods	•	70	2.070
Vomiting 12047	20 mg/day for 1 week, 40 mg/day later	0	53	0.0%
12041	20 mgraay loi 1 week, 40 mgraay latel	U	JJ	0.0 /0

Fluoxetine/Lidocaine

Reference	Description/Comments	No. of AE's	No. of Patients	Rate
Headache 795257	Fluoxetine 20mg/day + topical lidocaine cream	4	17	23.5%
Insomnia 795257	Fluoxetine 20mg/day + topical lidocaine cream	0	17	0.0%
Nausea 795257	Fluoxetine 20mg/day + topical lidocaine cream	1	17	5.9%

Paroxetine

Reference	Description/Comments	No. of AE's	No. of Patients	Rate
Anejaculation				
12005	20 mg/day for 4 weeks	5	61	8.2%
12020	10 mg/day for 3 weeks; followed by 20mg 3-4 hrs prior to intercourse	3	42	7.1%
500005	10 mg/day for 21 days, followed by 20 mg 3-4 hrs before intercourse for 6 months	1	40	2.5%
500005	10 mg/day for 21 days, followed by 20 mg 3-4 hrs plus 50 mg sildenafil before intercourse for 6 months	1	40	2.5%
Anejaculation and frequent yawnin	g			
12088	20 mg/day for 8 weeks	2	17	11.8%
Anejaculation, yawning, fatigue, pe	erspiration			
12088	40 mg/day for 8 Weeks	4	17	23.5%
Anorexia				
12020	10 mg/day for 3 weeks; followed by 20mg 3-4 hrs prior to intercourse	1	42	2.4%
700014	20 mg 3-5 hrs before coitus, not > 2 x per week	0	31	0.0%
Any side effect				
12005	20 mg single dose 3-4 hrs prior to intercourse	0	37	0.0%
12020	20 mg single dose 3-4 hours before coitus for 4 weeks	0	13	0.0%
Decreased Libido				
500005	10 mg/day for 21 days, followed by 20 mg 3-4 hrs before intercourse	2	40	5.0%
500005	for 6 months 10 mg/day for 21 days, followed by 20 mg 3-4 hrs plus 50 mg sildenafil	1	40	2.5%
	before intercourse for 6 months		40	2.570
Drowsiness				
700014	20 mg 3-5 hrs before coitus, not > 2 x per week	0	31	0.0%
Drowsiness & anorexia				
12005	20 mg/day for 4 weeks	1	61	1.6%
Dry mouth				
12088	40 mg/day for 8 Weeks	1	17	5.9%
700014	20 mg 3-5 hrs before coitus, not > 2 x per week	2	31	6.5%
Erectile function - slight reduction				
12088	20 mg/day for 8 weeks	1	17	5.9%
Flushing				
500005	10 mg/day for 21 days, followed by 20 mg 3-4 hrs before intercourse	0	40	0.0%
500005	for 6 months 10 mg/day for 21 days, followed by 20 mg 3-4 hrs plus 50 mg sildenafil			
00000	before intercourse for 6 months	6	40	15.0%
700014	20 mg 3-5 hrs before coitus, not > 2 x per week	0	31	0.0%
GI upset				
12020	10 mg/day for 3 weeks; followed by 20mg 3-4 hrs prior to intercourse	3	42	7.1%
500005	10 mg/day for 21 days, followed by 20 mg 3-4 hrs before intercourse for 6 months	5	40	12.5%
500005	10 mg/day for 21 days, followed by 20 mg 3-4 hrs plus 50 mg sildenafil	6	40	15.0%
	before intercourse for 6 months	-	-	
GI upset - minor				
12005	20 mg/day for 4 weeks	2	61	3.3%
Headache				
700014	20 mg 3-5 hrs before coitus, not > 2 x per week	0	31	0.0%
500005	10 mg/day for 21 days, followed by 20 mg 3-4 hrs before intercourse for 6 months	4	40	10.0%
500005	10 mg/day for 21 days, followed by 20 mg 3-4 hrs plus 50 mg sildenafil	8	40	20.0%
	before intercourse for 6 months		-	

Paroxetine

Reference	Description/Comments	No. of AE's	No. of Patients	Rate
Libido - reduced 12005 12020 12088	20 mg/day for 4 weeks 10 mg/day for 3 weeks; followed by 20mg 3-4 hrs prior to intercourse 40 mg/day for 8 Weeks	3 2 1	61 42 17	4.9% 4.8% 5.9%
Nasal congestion				
700014	20 mg 3-5 hrs before coitus, not > 2 x per week	0	31	0.0%
Nausea 700014	20 mg 3-5 hrs before coitus, not > 2 x per week	1	31	3.2%
Orgasm - inhibited 12005	20 mg/day for 4 weeks	3	61	4.9%
Sensory confusion - mild				
12118	20 mg/day for 2 months	21	32	65.6%
Sensory confusion - severe				
12118	20 mg/day for 2 months	1	32	3.1%
Sleepiness 12118 700014	20 mg/day for 2 months 20 mg 3-5 hrs before coitus, not > 2 x per week	14 0	32 31	43.8% 0.0%
Yawning				
700014	20 mg 3-5 hrs before coitus, not > 2 x per week	2	31	6.5%
Yawning, perspiration, dry mouth, t	fatigue, nausea			
12088 12088	20 mg/day for 8 weeks 40 mg/day for 8 Weeks	6 7	17 17	35.3% 41.2%

Placebo

Reference	Description/Comments	No. of AE's	No. of Patients	Rate
Anejaculation 12139	145 mg/day mean final dosage	0	26	0.0%
Any side effect 12020 12139	Placebo for 4 weeks after 3 week washout 145 mg/day mean final dosage	0 16	13 26	0.0% 61.5%
Burning on micturation 900017	5 mg/day for 2 weeks, 10 mg/day for 2 weeks	1	40	2.5%
Change in stools 900017	5 mg/day for 2 weeks, 10 mg/day for 2 weeks	1	40	2.5%
Constipation 12219	25 mg 6 hrs prior to coitus	2	10	20.0%
Diarrhea 12033	Placebo	1	15	6.7%
12139 Discontinued due to AE	145 mg/day mean final dosage	1	26	3.8%
12139 Dizziness	145 mg/day mean final dosage	2	26	7.7%
12139 Drowsiness	145 mg/day mean final dosage	1	26	3.8%
12047 12219	1 capsule day for 1 week, 2 per day later 25 mg 6 hrs prior to coitus	2 1	53 10	3.8% 10.0%
Dry mouth 12033 12047	Placebo 1 capsule day for 1 week, 2 per day later	0	15 53	0.0% 0.0%
12139 Dyspepsia	145 mg/day mean final dosage	2	26	7.7%
12139 Erectile dysfunction	145 mg/day mean final dosage	1	26	3.8%
12020 Erectile dysfunction - minor	Placebo daily	2	42	4.8%
12057 Fatigue	Placebo	1	18	5.6%
12139 Flatulence	145 mg/day mean final dosage	0	26	0.0%
12139 Headache	145 mg/day mean final dosage	1	26	3.8%
12020 12033 12139	Placebo PRN Placebo 145 mg/day mean final dosage	1 3 3	42 15 26	2.4% 20.0% 11.5%
Insomnia 12139	145 mg/day mean final dosage	1	26	3.8%
Libido - decreased 900017	5 mg/day for 2 weeks, 10 mg/day for 2 weeks	1	40	2.5%
Libido - increased 900017	5 mg/day for 2 weeks, 10 mg/day for 2 weeks	1	40	2.5%

Placebo

Reference	Description/Comments	No. of AE's	No. of Patients	Rate
Nausea 12047 12139	1 capsule day for 1 week, 2 per day later 145 mg/day mean final dosage	1 1	53 26	1.9% 3.8%
Nausea - mild 12219	25 mg 6 hrs prior to coitus	1	10	10.0%
Other	· ·	2		
12047 Potency - reduced	1 capsule day for 1 week, 2 per day later	3	53	5.7%
12047 Sleepiness	1 capsule day for 1 week, 2 per day later	3	53	5.7%
12033	Placebo	3	15	20.0%
Somnolence 12139	145 mg/day mean final dosage	3	26	11.5%
Vomiting 12047	1 capsule day for 1 week, 2 per day later	0	53	0.0%

Topical Anesthetics

Reference	Description/Comments	No. of AE's	No. of Patients	Rate
Delayed ejaculation				
500006	Prilocaine-lidocaine cream applied and left in place under a condom for 30 min and removed – used at least 5 times (range 5-9)	6	10	60.0%
500006	Prilocaine-lidocaine cream applied and left in place under a condom for 45 min and removed – used at least 5 times (range 5-9)	10	10	100.0%
Inability to reach orgasm				
500003	Lidocaine-prilocaine cream (1/2 tube) applied for 10 min, contained with condom, then removed. Patients given 5 tubes and 12 condoms.	3	15	20.0%
Numbness				
500006	Prilocaine-lidocaine cream applied and left in place under a condom for 30 min and removed – used at least 5 times (range 5-9)	6	10	60.0%
500006	Prilocaine-lidocaine cream applied and left in place under a condom for 45 min and removed – used at least 5 times (range 5-9)	10	10	100.0%

Sertraline

Reference	Description/Comments	No. of AE's	No. of Patients	Rate
Anejaculation				
12037	25 mg/day for 3 weeks	0	46	0.0%
12037	100 mg/day for 3 weeks	10	46	21.7%
12037	50 mg/day for 3 weeks	4	46	8.7%
12139	121 mg/day mean final dosage	5	26	19.2%
Anejaculation & withdrawal from st	udy			
12057	50 mg/day for 4 weeks	2	19	10.5%
Anorexia				
12037	50 mg/day for 3 weeks	1	46	2.2%
12037	100 mg/day for 3 weeks	2	46	4.3%
12037	25 mg/day for 3 weeks	0	46	0.0%
700014	50 mg 3-5 hrs before coitus, not > 2 x per week	1	31	3.2%
Anxiety				
12037	25 mg/day for 3 weeks	0	46	0.0%
12037	50 mg/day for 3 weeks	0	46	0.0%
12037	100 mg/day for 3 weeks	2	46	4.3%
Any side effect				
12139	121 mg/day mean final dosage	17	26	65.4%
Delayed ejaculation				
12013	50mg/day for 2 weeks / 50 mg at 5 pm when intercourse	1	24	4.2%
Diarrhea				
12033	50 mg/day for 4 weeks	3	22	13.6%
12139	121 mg/day mean final dosage	6	26	23.1%
Discontinued due to AE				
12139	121 mg/day mean final dosage	0	26	0.0%
Dizziness				
12037	25 mg/day for 3 weeks	1	46	2.2%
12037	50 mg/day for 3 weeks	0	46	0.0%
12037	100 mg/day for 3 weeks	0	46	0.0%
12139	121 mg/day mean final dosage	3	26	11.5%
500002	50 mg 4 hours prior to intercourse for 6 months, pts with primary PE	3	52	5.8%
500002	50 mg 4 hours prior to intercourse for 6 months, pts with ED treated successfully with sildenafil	4	35	11.4%
Parameters	Successionly with shoenam			
Drowsiness	400 maldov for 2 wools	2	46	4.20/
12037	100 mg/day for 3 weeks	2	46 46	4.3%
1203 <i>7</i> 12037	25 mg/day for 3 weeks 50 mg/day for 3 weeks	0 1	46 46	0.0% 2.2%
12047	50 mg/day for 1 week, 100 mg/day later	7	53	13.2%
700014	50 mg 3-5 hrs before coitus, not > 2 x per week	1	31	3.2%
Drowsiness & anorexia				
12057	50 mg/day for 4 weeks	1	19	5.3%
Dry mouth				
12003	50 mg/day	6	31	19.4%
12033	50 mg/day for 4 weeks	2	22	9.1%
12047	50 mg/day for 1 week, 100 mg/day later	4	53	7.5%
12139	121 mg/day mean final dosage	4	26	15.4%
500002	50 mg 4 hours prior to intercourse for 6 months, pts with primary PE	1	52	1.9%
500002	50 mg 4 hours prior to intercourse for 6 months, pts with ED treated	1	35	2.9%
700014	successfully with sildenafil 50 mg 3-5 hrs before coitus, not > 2 x per week	0	31	0.0%
700014	oo mg o-o mo belore collus, not > 2 x per week	U	٥١	0.076

Sertraline

Reference	Description/Comments	No. of AE's	No. of Patients	Rate
Dyspepsia				
12037	50 mg/day for 3 weeks	1	46	2.2%
12037	100 mg/day for 3 weeks	2	46	4.3%
12037	25 mg/day for 3 weeks	0	46	0.0%
12139	121 mg/day mean final dosage	2	26	7.7%
	3 ,			
Erectile dysfunction	05 markley (m. 0 mm d. n	0	40	0.00/
12037	25 mg/day for 3 weeks	0	46	0.0%
12037	50 mg/day for 3 weeks	0	46 46	0.0%
12037 12057	100 mg/day for 3 weeks 50 mg/day for 4 weeks	2 0	46 19	4.3% 0.0%
	30 mg/day for 4 weeks	O	19	0.076
Fatigue		_		
12013	50mg/day for 2 weeks / 50 mg at 5 pm when intercourse	2	24	8.3%
12139	121 mg/day mean final dosage	4	26	15.4%
Flatulence				
12139	121 mg/day mean final dosage	2	26	7.7%
Flushing				
700014	50 mg 3-5 hrs before coitus, not > 2 x per week	0	31	0.0%
GI upset				
12057	50 mg/day for 4 weeks	2	19	10.5%
Headache	os mgras, is. i mosno	_		. 0.0 / 0
	50 // (4 /		00	07.00/
12033	50 mg/day for 4 weeks	6	22	27.3%
12139 700014	121 mg/day mean final dosage	3 0	26 31	11.5%
	50 mg 3-5 hrs before coitus, not > 2 x per week	U	31	0.0%
Insomnia				
12139	121 mg/day mean final dosage	3	26	11.5%
Libido - reduced				
12037	25 mg/day for 3 weeks	0	46	0.0%
12037	100 mg/day for 3 weeks	2	46	4.3%
12037	50 mg/day for 3 weeks	0	46	0.0%
12057	50 mg/day for 4 weeks	0	19	0.0%
Nasal congestion				
700014	50 mg 3-5 hrs before coitus, not > 2 x per week	0	31	0.0%
Nausea				
12003	50 mg/day	2	31	6.5%
12047	50 mg/day for 1 week, 100 mg/day later	2	53	3.8%
12139	121 mg/day mean final dosage	3	26	11.5%
500002	50 mg 4 hours prior to intercourse for 6 months, pts with primary PE	2	52	3.8%
500002	50 mg 4 hours prior to intercourse for 6 months, pts with ED treated	2	25	0.60/
	successfully with sildenafil	3	35	8.6%
700014	50 mg 3-5 hrs before coitus, not > 2 x per week	1	31	3.2%
Numbness in extremities				
12013	50mg/day for 2 weeks / 50 mg at 5 pm when intercourse	1	24	4.2%
Orgasmic intensity - reduced				
12057	50 mg/day for 4 weeks	0	19	0.0%
Other	3.17			
	FO maildou for 1 wools 100 maildou later	0	5 0	0.00/
12047	50 mg/day for 1 week, 100 mg/day later	0	53	0.0%
Potency - reduced				
12047	50 mg/day for 1 week, 100 mg/day later	3	53	5.7%
Sleepiness				
12033	50 mg/day for 4 weeks	6	22	27.3%
700014	50 mg 3-5 hrs before coitus, not > 2 x per week	0	31	0.0%

Sertraline

Reference	Description/Comments	No. of AE's	No. of Patients	Rate
Somnolence 12139	121 mg/day mean final dosage	2	26	7.7%
Vomiting 12047	50 mg/day for 1 week, 100 mg/day later	0	53	0.0%
Yawning 700014	50 mg 3-5 hrs before coitus, not > 2 x per week	0	31	0.0%

Sildenafil

Reference	Description/Comments	No. of AE's	No. of Patients	Rate
Anejaculation				
500005	10 mg/day paroxetine for 21 days, followed by 20 mg 3-4 hrs plus 50 mg sildenafil before intercourse for 6 months	1	40	2.5%
Anorexia				
700014	50 mg 3-5 hrs before coitus, not > 2 x per week	0	31	0.0%
Decreased libido				
500005	10 mg/day paroxetine for 21 days, followed by 20 mg 3-4 hrs plus 50 mg sildenafil before intercourse for 6 months	1	40	2.5%
Drowsiness				
700014	50 mg 3-5 hrs before coitus, not > 2 x per week	0	31	0.0%
Dry mouth				
700014	50 mg 3-5 hrs before coitus, not > 2 x per week	0	31	0.0%
Flushing				
500005	10 mg/day paroxetine for 21 days, followed by 20 mg 3-4 hrs plus 50 mg sildenafil before intercourse for 6 months	6	40	20.0%
700014	50 mg 3-5 hrs before coitus, not > 2 x per week	2	31	6.5%
GI upset/nausea				
500005	10 mg/day paroxetine for 21 days, followed by 20 mg 3-4 hrs plus 50 mg sildenafil before intercourse for 6 months	6	40	15.0%
Headache				
500005	10 mg/day paroxetine for 21 days, followed by 20 mg 3-4 hrs plus 50 mg sildenafil before intercourse for 6 months	8	40	15.0%
700014	50 mg 3-5 hrs before coitus, not > 2 x per week	2	31	6.5%
Nasal congestion				
700014	50 mg 3-5 hrs before coitus, not > 2 x per week	1	31	3.2%
Nausea				
700014	50 mg 3-5 hrs before coitus, not > 2 x per week	0	31	0.0%
Sleepiness				
700014	50 mg 3-5 hrs before coitus, not > 2 x per week	0	31	0.0%
Yawning				
700014	50 mg 3-5 hrs before coitus, not > 2 x per week	0	31	0.0%

Appendix 3-A. Summary Tables: Citalopram treatment studies

	Treatment regimen				Latency in Second				
Treatment Groups	Week	Dosage	Treatment Duration	Number of Patients	Mean	SD	Min	Max	
		. H., Olivier, B. SSRIs and			ed, fixed-dose stu	dy with paro	xetine and		
citalopram. J Cli	in Psychopharma	col 2001 Dec; 21: 556-60.	(ProCite no. 79522	20)					
1	Baseline		6 weeks	15	21*				
	1	10 mg/day			34				
	2	20 mg/day			43				
	3	20 mg/day			40				
	4	20 mg/day			33				
	5	20 mg/day			44				
	6	20 mg/day			43				
	* @	Seometric means							

Appendix 3-B. Summary Tables: Clomipramine treatment studies

		Treatment regimen	t regimen Latency in Seconds					
Treatment Groups	Week	Dosage	Treatment Duration	Number of Patients	Mean	SD	Min	Max
		, E. A., El Gilany, A. H. Asses npot Res. 2001 Feb; 13: 41-5.			erapy and the p	ause-squeez	e technique	e in
1	Baseline			31	60		30	90
	Overall/ entire time period	25 mg 3-5 hrs before coitus, not > 2 x per week Crossover design - 2-week	4 weeks + 2 week washout	28	240		60	480
		orty, E. W., Risen, C. B., Sterr 1995 Sep; 56: 402-7. (ProCit	ı, E. B., Kurit, D. M	I. A double-blind cros	sover trial of clo	mipramine fo	or rapid ejac	ulation in
1	Baseline			15	81			
	Overall/ entire time period	1-4 week run-out period 3 intercourse attempts	1-4 weeks	15	80			
		Crossover design, time point	s vary to allow for	normal pattern of inte	ercourse attempt	ts		
	Overall/ entire time period	25 mg/day for 2-7 weeks 5 intercourse attempts	2-7 weeks	15	202			
	Overall/ entire time period	50 mg/day for 2-7 weeks 5 intercourse attempts	2-7 weeks	15	419			
		and safety of fluoxetine, sert Feb; 159: 425-7. (ProCite no.		amine in patients with	n premature ejac	ulation: a do	uble-blind, p	olacebo
1	Baseline			53	46	41		
	Overall/ entire time period	25 mg/day for 1 week, 50 mg/day later	4 weeks	37	345	401		
		Crossover design - 4-week v	vashout period					
, ,	, ,	imboli, F., Rigatti, P., Pizzini, a Urol Ital. 1995; 1: 5-6. (ProC	, ,	ipramine for prematu	re ejaculation: a	randomized	, double blir	nd,
·	Overall/ entire time period	50 mg/day given at bedtime for 8 weeks (following 8 weeks placebo treatment)	8 weeks	17				
2	Baseline	Only efficacy outcome is 5 (2	29.4%) complete re	esponse, 6 (35.3%) p	artial response,	6 (35.3%) fa	ilure	
	Overall/ entire time period	50 mg/day given at bedtime for 8 weeks (presumably continued for at least 3 months)	8 weeks	16				
		Only efficacy outcome is 4 (2	25.0%) complete re	esponse, 4 (25.0%) p	artial response,	8 (50.0%) fa	ilure	

Appendix 3-B. Summary Tables: Clomipramine treatment studies

Арренаіх	o B. Oui	nmary rables: Clor	•	i catillolit Sta				
		Treatment regimen			Late	ency in Se	econds	
Treatment Groups	Week	Dosage	Treatment Duration	Number of Patients	Mean	SD	Min	Max
		Brazao, C. A., Koos Slob, A. E e. BJU Int. 2001 Mar; 87: 357-6			ine in men with p	remature eja	culation who	en 25
1	Baseline		9 weeks	4	26			
	3	1 mg/day for 3 weeks then 20 mg/day for 3 weeks*		4	51			
	6	1 mg/day for 3 weeks then 20 mg/day for 3 weeks		4	92			
	9	1 mg/day for 3 weeks then 20 mg/day for 3 weeks		4	221			
		These are patients who failed * This value is a median from egraves, K., Maguire, E. Clom 98-200. (ProCite no. 12219)	a graph		ent of premature e	ejaculation: a	a pilot study.	J Sex
1	Overall/ entire time period	25 mg 6 hrs prior to coitus, could double dose to 50 mg after 2 attempts at coitus and no side effects	10 attempts at coitus	10	366			
		a Brazao, C. A., Rowland, D. L 101. (ProCite no. 12022)	, Tan, P., Slob, A.	K. Clomipramine in	the treatment of	rapid (prema	ature) ejacul	ation. J
1	Overall/ entire time	25 mg 4 hrs precoitus	2 weeks	22	229	286		
	period	PE patients – crossover desi	gn, no washout – 2	weeks drug, then 2	2 weeks placebo a	and vice-vers	sa	
2	Overall/ entire time period	25 mg 4 hrs precoitus	2 weeks	4	665	307		
	•	Normal controls - crossover of	design, no washou	t period - 2 weeks d	rug, then 2 weeks	placebo an	d vice-versa	l

Appendix 3-C. Summary Tables: Fluoxetine treatment studies

		Treatment regimen			Lat	ency in S	econds	
Treatment Groups	Week	Dosage	Treatment Duration	Number of Patients	Mean	SD	Min	Max
		anli, L. Comparison of the effi sp Urol. 2000 Nov; 53: 856-8.			plus local lidocai	ne ointment i	n the treatm	ent of
premature ejac	diation. Alch L	,	(110Cite 110. 7752	37)				
1	7	20 mg/day for 1 week, 40 mg/day later	7 weeks	26	'26 09/) failed			
Haensel SM k	(lem Tmal Ho	Only efficacy outcome is 8 (3 p, WJC and Slob, AK Fluoxet				acology 100	Q·1Q·1₋6 (DroCito
no. 900017)	dem, mai, no	p, woo and olob, Art i doxet	ille and i remature	Ejaculation. 3 Olimic	ан зуспорнанн	acology. 199	0, 10. 1-0. (riocite
1	Baseline (pts with PE		4 weeks	9	73	22		
	ົ only) Overall/	5 mg/day for 2 weeks, 10						
	entire time period	mg/day for 2 weeks		9				
	Baseline	(Latency increase 180% with	Confidence Interv	ral 80%-450%)				
2	(pts with		4 weeks	7	360	91		
	ED only) Overall/	5 mg/day for 2 weeks, 10						
	entire time period	mg/day for 2 weeks		7				
	·	(1/6 increase latency, 4/6 de	crease)					
3	Baseline (pts with		4 weeks	9	89	22		
3	ED and PE)		4 Weeks	9	09	22		
	Overall/ entire time	5 mg/day for 2 weeks, 10 mg/day for 2 weeks		9				
	period	(6/8 increase latency, 1/8 dec	crease. 1/8 no cha	nge)				
	Baseline	(-, -, -, -, -, -, -, -, -, -, -, -, -, -		•				
4	(Normal controls)		4 weeks	15	535	116		
	Overall/ entire time	5 mg/day for 2 weeks, 10 mg/day for 2 weeks		15				
	period	(7/15 increase latency, 8/15	,					
Kara, H., Avdin	. S Yucel. M	All groups had Fluoxetine an Agargun, M. Y., Odabas, O.,			•	premature e	iaculation: a	double-
		v. J Urol. 1996 Nov; 156: 1631					,	
1	Baseline		4 weeks	9	25	13		
	4	20 mg/day for 1 week, 40 mg/day later		7	180	100		
		and safety of fluoxetine, sertr Feb;159: 425-7. (ProCite no. 7		mine in patients wit	h premature ejac	ulation: a do	uble-blind, _l	olacebo
2	Baseline	100/107. 120 7. (1100He 110.	4 weeks	53	46	41		
2	Overall/	20 mg/day for 1 week,	+ WCCK3	33	40	71		
	entire time period	40 mg/day later		37	138	125		
	penou	Crossover design - 4-week	washout period					
Lee, H. S., Son 1996 Oct; 16: 3		C. H., Choi, H. K. An open clir e no. 12112)	nical trial of fluoxet	ine in the treatment	of premature ejac	culation. J Cl	in Psychoph	armacol.
1	Baseline		8 weeks	14	55	3		
		20 mg/day for 1-2						
	8	weeks, titrated to 60		11	578	420		
		mg/day later 2 week washout before active	ve drug					
		2 WOOK WASHOUL DETOILE ACTIV	ro arag					

Appendix 3-C. Summary Tables: Fluoxetine treatment studies

		Treatment regimer	า	<u> </u>	Late	ency in S	econds	
Treatment Groups	Week	Dosage	Treatment Duration	Number of Patients	Mean	SD	Min	Max
Raju, G. A. R., 1997; 29: 204-5	•	t., Ramesh T., Schobha, J. C 500001).	. Evaluation of fluox	cetine in premature e	jaculation. Indiar	n Journal of	Pharmacolo	gy.
1	Baseline		4 weeks	44	10.7	4		
	Overall/ entire time period	20 mg/day in the morning for 4 weeks		44	3.2	2		
12044)		fluoxetine, fluvoxamine, paro					1. (ProCite r	10.
1	Baseline		6 weeks	10	21	12		
	6	20 mg/day		10	211	251		
		n, H., Arman, F., Ekmekcioglu 1999 Jan; 161: 107-11. (Prod	·	fluoxetine on several	neurophysiologi	cal variables	in patients	with
1	Baseline		4 weeks	20	72	60		
	4	20 mg/day for 1 week, 40 mg/day later		20	396	462		

Appendix 3-D. Summary Tables: Fluoxetine/Lidocaine treatment studies

		Treatment regimen			Latency in Seconds			
Treatment Groups	Week	Dosage	Treatment Duration	Number of Patients	Mean	SD	Min	Max
	. , .	anli, L. Comparison of the eff sp Urol. 2000 Nov; 53: 856-8	•		plus local lidocain	e ointment i	n the treatm	ent of
2	7	Fluoxetine 20 mg/day + local lidocaine cream Only efficacy outcome is 9 (7 weeks 52.9%) cured, 5 (2	17 29.4%) improved an	d 3 (17.6%) failed.			

Appendix 3-E. Summary Tables: Fluvoxamine treatment studies

		Treatment regimen			Latency in Seconds			
Treatment Groups	Week	Dosage	Treatment Duration	Number of Patients	Mean	SD	Min	Max
•		M. W., Zwinderman, A. H., O uoxetine, fluvoxamine, parox	•	•	•			-
2	Baseline		6 weeks	10	15	17		
	6	100 mg/day		10	55	70		

Appendix 3-F. Summary Tables: Nefazodone treatment studies

_		Treatment regimen		_	Late	ency in S	econds	
Treatment Groups	Week	Dosage	Treatment Duration	Number of Patients	Mean	SD	Min	Max
•	•	n, A. H., Olivier, B. Antidepres d nefazodone. J Clin Psychop	•			ebo- controll	ed, fixed-do	se study
4	Baseline		6 weeks	12	17*			
	1	400 mg/day morning and evening			19			
	2	400 mg/day morning and evening			14			
	3	400 mg/day morning and evening			26			
	4	400 mg/day morning and evening			28			
	5	400 mg/day morning and evening			16			
	6	400 mg/day morning and evening			18			
		* Geometric means						

Pharmacologic Treatment on Latency

Appendix 3-G. Summary Tables: Paroxetine treatment studies

		Treatment regime	n		Late	ency in Se	econds	
Treatment Groups	Week	Dosage	Treatment Duration	Number of Patients	Mean	SD	Min	Max
	, 00	r, E. A., El Gilany, A. H. Asse npot Res. 2001 Feb; 13: 41-5			herapy and the pa	ause-squeez	e technique	in
2	Baseline			31	60		30	90
	Overall/ entire time period	20 mg 3-5 hrs before coitus, not > 2 x per week Crossover design - 2-week	4 weeks + 2 week washout washout period	29	240		120	600
		., Pagliarulo, G., Cirillo-Maruo 2. (ProCite no. 12118)	cco, E., Marano, A.,	Pagliarulo, A. Paro	xetine in the treat	ment of pren	nature ejacu	lation.
1	Baseline (latency < 1 min.)		2 months	32				
	8	20 mg/day		32			900	1200
McMahon, C. G 246. (ProCite n		Freatment of premature ejacu	llation with paroxetir	ne hydrochloride. In	t J Impot Res. 199	99 Oct; 11: 2	41-245; disc	cussion
1	Baseline 4	20 mg/day	4 weeks	61 61	24 270			
	8	20 mg single dose 3-4 hrs precoitus	4 weeks	53*	234			
2	Baseline	* Only included responders	to first 4-week trial	33	24			
	4	20 mg single dose 3-4 hrs precoitus	4 weeks	33	90			
		Freatment of premature ejacu 99 Jun; 161: 1826-30. (ProCit		ne hydrochloride as	needed: 2 single-	blind placeb	o controlled	
1a	Baseline			13	18			
		Crossover design - drug, 3-	week washout, then	placebo				
	4	20 mg single dose 3-4 hrs precoitus	4 weeks	13	192			
2a	11	20 mg single dose 3-4 hrs precoitus	4 weeks	13	210			
Study b								
1b	Baseline			21	30			
	3	10 mg/day	3 weeks	21	258			
	7	20 mg single dose 3-4 hrs precoitus	4 weeks	21	348			
2b	13	3-week washout, then 20 mg/day for 3 weeks	6 weeks	21	198			
	17	20 mg single dose 3-4 hrs precoitus	4 weeks	21	366			

Appendix 3-G. Summary Tables: Paroxetine treatment studies

		Treatment regimen		_	Late	ency in S	econds	
Treatment Groups	Week	Dosage	Treatment Duration	Number of Patients	Mean	SD	Min	Max
		oo, R., Scattoni, V., Briganti, A xetine plus sildenafil in patient						
1	Baseline			40	.33	.04		
	12	10 mg/day for 21 days, followed by 20 mg 3-4 hrs before intercourse 10 mg/day for 21 days,	3 months	40	3.7	.10		
	24	followed by 20 mg 3-4 hrs before intercourse	6 months	40	4.2	.03		
2	Baseline	10 mg/day paroxetine for 21 days, followed by 20		40	.35	.05		
	12	mg 3-4 hrs plus 50 mg sildenafil before intercourse	3 months	40	4.5	.07		
	24	10 mg/day paroxetine for 21 days, followed by 20 mg 3-4 hrs plus 50 mg sildenafil before intercourse	6 months	40	5.3	.02		
		n, A. H., Olivier, B. SSRIs and macol 2001 Dec; 21: 556-60.			, fixed-dose stu	dy with paro	xetine and	
1	Baseline		6 weeks	15	18*			
	1	20 mg/day			34			
	2	20 mg/day			57			
	3	20 mg/day			116			
		20 mg/ady			141			
		20 ma/day						
	4	20 mg/day						
	4 5	20 mg/day			170			
	4							
	4 5 6 0., Zwindermal	20 mg/day 20 mg/day * Geometric means n, A. H., Olivier, B. Antidepres d nefazodone. J Clin Psychop 20 mg/day morning and evening 20 mg/day morning and			170 152 andomized, plac	ebo- control	led, fixed-do	se study
with paroxetine,	4 5 6 0., Zwindermal sertraline, an Baseline 1	20 mg/day 20 mg/day 20 mg/day * Geometric means n, A. H., Olivier, B. Antidepres d nefazodone. J Clin Psychop 20 mg/day morning and evening	harmacol. 2001 Ju	ın; 21: 293-7. (ProCite	170 152 andomized, place e no. 795222) 17* 37	ebo- control	led, fixed-do	se study
with paroxetine,	4 5 6 0., Zwindermal sertraline, an Baseline 1	20 mg/day 20 mg/day 20 mg/day * Geometric means n, A. H., Olivier, B. Antidepres d nefazodone. J Clin Psychop 20 mg/day morning and evening	harmacol. 2001 Ju	ın; 21: 293-7. (ProCite	170 152 andomized, place e no. 795222) 17* 37	ebo- control	led, fixed-do	se study
with paroxetine,	4 5 6 0., Zwindermal sertraline, an Baseline 1 2	20 mg/day 20 mg/day 20 mg/day * Geometric means n, A. H., Olivier, B. Antidepres d nefazodone. J Clin Psychop 20 mg/day morning and evening 20 mg/day morning and	harmacol. 2001 Ju	ın; 21: 293-7. (ProCite	170 152 andomized, place e no. 795222) 17* 37 71	ebo- control	led, fixed-do	se study

Appendix 3-G. Summary Tables: Paroxetine treatment studies

		Treatment regimen	า		Lat	ency in Se	econds	
Treatment Groups	Week	Dosage	Treatment Duration	Number of Patients	Mean	SD	Min	Max
		M. W., Zwinderman, A. H., C fluoxetine, fluvoxamine, paro						
3a	Baseline 6	20 mg/day	6 weeks	11 11	16 476	10 1146		
Study b								
1b	Baseline (BL < 60 sec)		6 weeks	12	18	13		
	6	20 mg/day		11	92			
3b	Baseline (BL > 60 sec)	(latency increase 580%)	6 weeks	5	82	27		
	6	20 mg/day		5	602			
		(latency increase 596%) M. W., Zwinderman, A. H. Ej				vith primary p	remature	
•	•	ndomized, dose-response stu	•		,			
1	Baseline		8 weeks	17	13			
	3	20 mg/day		14	300			
	8	20 mg/day		14	300			
2	Baseline		8 weeks	17	10			
	3	20 mg/day		13	240			
	8	20 mg/day		13	540			
		M. W., Zwinderman, A. H. Pa atry. 1994 Sep; 151: 1377-9.			ation: a double-b	lind, random	ized, placeł	00-
1	Baseline	20 mg/day for 1 week,	6 weeks	8	30			
	3	40 mg/day for 5 weeks		8	450		180	1200
	6	20 mg/day for 1 week, 40 mg/day for 5 weeks		8	600		300	1200

Appendix 3-H. Summary Tables: Pause/Squeeze treatment studies

		Treatment regimen			Lat	ency in Se	econds	
Treatment Groups	Week	Dosage	Treatment Duration	Number of Patients	Mean	SD	Min	Max
		A., El Gilany, A. H. Asses Res. 2001 Feb; 13: 41-5.		•	therapy and the pa	ause-squeez	e technique	in
premature ejac	diation. Int 3 impot	Nes. 20011 eb, 15. 41-5.	(Frocite no. 7000)	14)				
5	Baseline			31	60		30	90
	Overall/ entire time period		4 weeks + 2 week washout	29	180		60	420
	Cro	ossover design – 2-week v udy unclear about washou		oup)				

Appendix 3-I. Summary Tables: Phenoxybenzamine treatment studies

		Treatment regime	n		Lat	ency in S	econds	Max		
Treatment Groups	Week	Dosage	Treatment Duration	Number of Patients	Mean	SD	Min	Max		
		ci, F., Zanollo, A. Effect o tatis. 1986 Jan-Feb; 17:			camine) in the ma	anagement o	f premature			
1	Baseline			8	83					
	Overall/ entire time period	10 mg/day		8	338					

Appendix 3-J. Summary Tables: Placebo treatment studies

		Treatment regime	1		Late	ency in S	econds	
Treatment Groups	Week	Dosage	Treatment Duration	Number of Patients	Mean	SD	Min	Max
		orty, E. W., Risen, C. B., Ster 1995 Sep; 56: 402-7. (ProCi		. A double-blind cross	sover trial of clor	nipramine fo	or rapid ejac	ulation in
1	Overall/ entire time period	1 capsule/day for 2-7 weeks – 5 intercourse attempts Crossover design, time poir	2-7 weeks	15 normal pattern of inte	137 ercourse attempt	s		
		aran, M., Kupeli, B., Bozkirli, 30: 611-5. (ProCite no. 1203	I. Sertraline in the tr				l placebo co	ntrolled
2	Baseline 4	Placebo control	4 weeks	15 15	44 114	20 94		
Cooper, A. J., N no. 12387)	/lagnus, R. V.	A clinical trial of the beta bloc	ker propranolol in p	remature ejaculation	. J Psychosom F	Res. 1984; 2	28: 331-6. (P	roCite
1	Overall/ entire time period	Placebo	4 weeks	12	102			
		, Agargun, M. Y., Odabas, O. v. J Urol. 1996 Nov; 156: 163			the treatment of	premature e	ejaculation: a	double-
1	Baseline	1 consula/dou for 1	4 weeks	8	30	9		
	4	1 capsule/day for 1 week, 2 capsule/day later		7	60	47		
		and safety of fluoxetine, sert Feb;159:425-7. (ProCite no.		mine in patients with	premature ejacı	ulation: a do	ouble-blind, _l	olacebo
4	Baseline Overall/	1 capsule/day for 1		53	46	41		
	entire time period	week, 2 capsule/day	4 weeks	37	136	227		
		Crossover design - 4-week	washout period					
		rimboli, F., Rigatti, P., Pizzini, a Urol Ital. 1995; 1: 5-6. (ProC		pramine for prematui	re ejaculation: a	randomized	l, double blir	ıd,
,	Overall/ entire time	8 weeks placebo treatment	8 weeks	17				
	Overall/ entire time period	treatment Only efficacy outcome is 1 (5.9%) complete res	sponse, 2 (11.8%) pa				
McMahon, C. G	Overall/ entire time period 6., Touma, K. T	treatment	5.9%) complete resulation with paroxeting	sponse, 2 (11.8%) pa				
McMahon, C. G	Overall/ entire time period 6., Touma, K. T	treatment Only efficacy outcome is 1 (Freatment of premature ejaculary Jun; 161: 1826-30. (ProCital Placebo for 4 weeks	5.9%) complete resulation with paroxeting	sponse, 2 (11.8%) pa				
McMahon, C. G crossover studi Study a	Overall/ entire time period G., Touma, K. T es. J Urol. 199	treatment Only efficacy outcome is 1 (Freatment of premature ejacu 9 Jun; 161: 1826-30. (ProCit Placebo for 4 weeks after 3 week washout	5.9%) complete res lation with paroxetin e no. 12020)	sponse, 2 (11.8%) pa ne hydrochloride as n	eeded: 2 single-			
McMahon, C. G crossover studi Study a 1a	Overall/ entire time period 5., Touma, K. T es. J Urol. 199	treatment Only efficacy outcome is 1 (Freatment of premature ejaculary Jun; 161: 1826-30. (ProCital Placebo for 4 weeks	5.9%) complete res lation with paroxetin e no. 12020)	sponse, 2 (11.8%) pa ne hydrochloride as n 13	eeded: 2 single- 27			
McMahon, C. G crossover studi Study a 1a	Overall/ entire time period 5., Touma, K. T es. J Urol. 199 11 Baseline	treatment Only efficacy outcome is 1 (Freatment of premature ejaculary Jun; 161: 1826-30. (ProCitive Placebo for 4 weeks after 3 week washout Placebo for 3 weeks, then 3 week washout	5.9%) complete res lation with paroxetin e no. 12020) 4 weeks	sponse, 2 (11.8%) pa ne hydrochloride as n 13 13	eeded: 2 single- 27 18			
McMahon, C. G crossover studi Study a 1a 2a	Overall/ entire time period 5., Touma, K. T es. J Urol. 199 11 Baseline	treatment Only efficacy outcome is 1 (Freatment of premature ejacu 9 Jun; 161: 1826-30. (ProCit Placebo for 4 weeks after 3 week washout Placebo for 3 weeks, then 3 week washout 3 week washout, then placebo daily for 3 weeks	5.9%) complete res lation with paroxetin e no. 12020) 4 weeks	sponse, 2 (11.8%) pa ne hydrochloride as n 13 13	eeded: 2 single- 27 18			
McMahon, C. G crossover studi Study a 1a 2a Study b	Overall/ entire time period 5., Touma, K. T es. J Urol. 199 11 Baseline 4	treatment Only efficacy outcome is 1 (Freatment of premature ejacu 9 Jun; 161: 1826-30. (ProCite Placebo for 4 weeks after 3 week washout Placebo for 3 weeks, then 3 week washout 3 week washout, then placebo daily for 3	5.9%) complete res lation with paroxetin e no. 12020) 4 weeks 4 weeks	sponse, 2 (11.8%) pa ne hydrochloride as n 13 13 13	eeded: 2 single- 27 18 36			
McMahon, C. G crossover studi Study a 1a 2a Study b	Overall/ entire time period 5., Touma, K. T es. J Urol. 199 11 Baseline 4	treatment Only efficacy outcome is 1 (Freatment of premature ejacu 9 Jun; 161: 1826-30. (ProCite Placebo for 4 weeks after 3 week washout Placebo for 3 weeks, then 3 week washout 3 week washout, then placebo daily for 3 weeks Placebo 3-4 hours before coitus for 4 weeks	5.9%) complete resulation with paroxetine no. 12020) 4 weeks 4 weeks	sponse, 2 (11.8%) pa ne hydrochloride as n 13 13 13	eeded: 2 single- 27 18 36			
McMahon, C. G crossover studi Study a 1a 2a Study b	Overall/ entire time period 3., Touma, K. T es. J Urol. 199 11 Baseline 4 13	treatment Only efficacy outcome is 1 (Freatment of premature ejacu 9 Jun; 161: 1826-30. (ProCite Placebo for 4 weeks after 3 week washout Placebo for 3 weeks, then 3 week washout 3 week washout, then placebo daily for 3 weeks Placebo 3-4 hours	5.9%) complete resulation with paroxetine no. 12020) 4 weeks 4 weeks	sponse, 2 (11.8%) pa ne hydrochloride as n 13 13 13 21	eeded: 2 single- 27 18 36 54 36			

Appendix 3-J. Summary Tables: Placebo treatment studies

		Treatment regime	n		Latency in Second			
Tuestan				Name				
Treatment Groups	Mook	Decem	Treatment	Number of Patients	Maan	en.	Min	Max
•	Week	Dosage f premature ejaculation with	Duration		Mean	SD orongover of	Min	Max
Jun; 159: 1935-			sertraline nydrochlor	ide: a single-blind p	iacedo controlled	crossover si	udy. J Oloi.	1996
1	12	Placebo for 4 weeks,	4 weeks	19	30			
2	Dogalina	Crossover design - 4-week	washout	40	10			
2	Baseline 4	End of placebo	4 weeks	18 18	18 30			
	•	Crossover design - 4-week		10	00			
	8	Washout for 4 weeks	4 weeks	18	30			
	mera, A., Sike	es, C. Sertraline treatment fo	r premature ejaculat	ion. J Clin Psychopl	narmacol. 1995 O	ct; 15: 341-6	. (ProCite n	٥.
12139)								
2	Baseline		8 weeks	26	66	81		
_	24000	145 mg/day mean final	o moone	_0		٠.		
	8	dosage, 50 mg/day		22	111	221		
		titrated to 200 mg/day during weeks 1-3						
Segraves, R. T.,	Saran, A., Se	graves, K., Maguire, E. Clom	ipramine versus plad	cebo in the treatme	nt of premature e	iaculation: a	pilot study.	J Sex
		98-200. (ProCite no.12219)			p	,	J	
	0 "'							
2	Overall/ entire time	25 mg 6 hrs prior to	10 attempts at	10	51			
_	period	coitus	coitus	10	01			
	•	(Could double dose to 50 m	ng after 2 attempts a	t coitus and no side	effects)			
		Brazao, C. A., Rowland, D.	L., Tan, P., Slob, A.	K. Clomipramine in	the treatment of	rapid (prema	ture) ejacul	ation. J
Sex Marital The	r 1999; 25:89-	-101. (ProCite no. 12022)						
	Overall/							
		05 +- 1						
1	entire time	25 mg taken 4 hr precoitus	2 weeks	22	52	45		
1	entire time period	precoitus						
1		precoitus PE patients – crossover de					sa	
2	period	precoitus PE patients – crossover de 25 mg taken 4 hr					sa	
	period Overall/	precoitus PE patients – crossover de 25 mg taken 4 hr precoitus	sign, no washout – 2 2 weeks	2 weeks drug, then 2	2 weeks placebo a	and vice-vers		
2	period Overall/ entire time period	precoitus PE patients – crossover de 25 mg taken 4 hr precoitus Normal controls - crossover	sign, no washout – 2 2 weeks r design, no washou	2 weeks drug, then 2 4 t period - 2 weeks d	2 weeks placebo a 491 rug, then 2 weeks	and vice-vers 245 s placebo and	d vice-versa	
2 Waldinger, M. D	period Overall/ entire time period O., Zwindermal	precoitus PE patients – crossover de 25 mg taken 4 hr precoitus Normal controls - crossover n, A. H., Olivier, B. Antidepre	sign, no washout – 2 2 weeks r design, no washou essants and ejaculati	2 weeks drug, then 2 4 t period - 2 weeks d ion: a double-blind,	2 weeks placebo a 491 rug, then 2 weeks randomized, plac	and vice-vers 245 s placebo and	d vice-versa	
2 Waldinger, M. D	period Overall/ entire time period O., Zwindermal	precoitus PE patients – crossover de 25 mg taken 4 hr precoitus Normal controls - crossover	sign, no washout – 2 2 weeks r design, no washou essants and ejaculati	2 weeks drug, then 2 4 t period - 2 weeks d ion: a double-blind,	2 weeks placebo a 491 rug, then 2 weeks randomized, plac	and vice-vers 245 s placebo and	d vice-versa	
2 Waldinger, M. D	period Overall/ entire time period O., Zwindermal	precoitus PE patients – crossover de 25 mg taken 4 hr precoitus Normal controls - crossover n, A. H., Olivier, B. Antidepre	sign, no washout – 2 2 weeks r design, no washou essants and ejaculati	2 weeks drug, then 2 4 t period - 2 weeks d ion: a double-blind,	2 weeks placebo a 491 rug, then 2 weeks randomized, plac	and vice-vers 245 s placebo and	d vice-versa	
2 Waldinger, M. D with paroxetine,	period Overall/ entire time period O., Zwinderman sertraline, an	precoitus PE patients – crossover de 25 mg taken 4 hr precoitus Normal controls - crossover n, A. H., Olivier, B. Antidepred nefazodone. J Clin Psycho	sign, no washout – 2 2 weeks r design, no washou essants and ejaculati opharmacol. 2001 Ju	2 weeks drug, then 2 4 t period - 2 weeks d ion: a double-blind, in; 21: 293-7. (ProCi	2 weeks placebo a 491 rug, then 2 weeks randomized, plac te no. 795222)	and vice-vers 245 s placebo and	d vice-versa	
2 Waldinger, M. D with paroxetine,	period Overall/ entire time period O., Zwindermal sertraline, an Baseline 1	precoitus PE patients – crossover de 25 mg taken 4 hr precoitus Normal controls - crossover n, A. H., Olivier, B. Antidepred nefazodone. J Clin Psychology Placebo morning and evening	sign, no washout – 2 2 weeks r design, no washou essants and ejaculati opharmacol. 2001 Ju	2 weeks drug, then 2 4 t period - 2 weeks d ion: a double-blind, in; 21: 293-7. (ProCi	2 weeks placebo a 491 rug, then 2 weeks randomized, plac te no. 795222) 15* 21	and vice-vers 245 s placebo and	d vice-versa	
2 Waldinger, M. D with paroxetine,	period Overall/ entire time period O., Zwindermal sertraline, an Baseline	precoitus PE patients – crossover de 25 mg taken 4 hr precoitus Normal controls - crossover n, A. H., Olivier, B. Antidepre d nefazodone. J Clin Psycho Placebo morning and evening Placebo morning and	sign, no washout – 2 2 weeks r design, no washou essants and ejaculati opharmacol. 2001 Ju	2 weeks drug, then 2 4 t period - 2 weeks d ion: a double-blind, in; 21: 293-7. (ProCi	2 weeks placebo a 491 rug, then 2 weeks randomized, plac te no. 795222)	and vice-vers 245 s placebo and	d vice-versa	
2 Waldinger, M. D with paroxetine,	period Overall/ entire time period O., Zwindermal sertraline, an Baseline 1	precoitus PE patients – crossover de 25 mg taken 4 hr precoitus Normal controls - crossover n, A. H., Olivier, B. Antidepre d nefazodone. J Clin Psycho Placebo morning and evening Placebo morning and evening Placebo morning and	sign, no washout – 2 2 weeks r design, no washou essants and ejaculati opharmacol. 2001 Ju	2 weeks drug, then 2 4 t period - 2 weeks d ion: a double-blind, in; 21: 293-7. (ProCi	2 weeks placebo a 491 rug, then 2 weeks randomized, plac te no. 795222) 15* 21	and vice-vers 245 s placebo and	d vice-versa	
2 Waldinger, M. D with paroxetine,	period Overall/ entire time period O., Zwinderman sertraline, an Baseline 1 2 3	precoitus PE patients – crossover de 25 mg taken 4 hr precoitus Normal controls - crossover n, A. H., Olivier, B. Antidepre d nefazodone. J Clin Psycho Placebo morning and evening Placebo morning and evening Placebo morning and evening Placebo morning and evening	sign, no washout – 2 2 weeks r design, no washou essants and ejaculati opharmacol. 2001 Ju	2 weeks drug, then 2 4 t period - 2 weeks d ion: a double-blind, in; 21: 293-7. (ProCi	2 weeks placebo a 491 rug, then 2 weeks randomized, plac te no. 795222) 15* 21 19	and vice-vers 245 s placebo and	d vice-versa	
2 Waldinger, M. D with paroxetine,	period Overall/ entire time period O., Zwindermal sertraline, an Baseline 1 2	precoitus PE patients – crossover de 25 mg taken 4 hr precoitus Normal controls - crossover n, A. H., Olivier, B. Antidepre d nefazodone. J Clin Psycho Placebo morning and evening Placebo morning and	sign, no washout – 2 2 weeks r design, no washou essants and ejaculati opharmacol. 2001 Ju	2 weeks drug, then 2 4 t period - 2 weeks d ion: a double-blind, in; 21: 293-7. (ProCi	2 weeks placebo a 491 rug, then 2 weeks randomized, place te no. 795222) 15* 21 19	and vice-vers 245 s placebo and	d vice-versa	
2 Waldinger, M. D with paroxetine,	period Overall/ entire time period O., Zwinderman sertraline, an Baseline 1 2 3	precoitus PE patients – crossover de 25 mg taken 4 hr precoitus Normal controls - crossover n, A. H., Olivier, B. Antidepre d nefazodone. J Clin Psycho Placebo morning and evening Placebo morning and evening Placebo morning and evening Placebo morning and evening	sign, no washout – 2 2 weeks r design, no washou essants and ejaculati opharmacol. 2001 Ju	2 weeks drug, then 2 4 t period - 2 weeks d ion: a double-blind, in; 21: 293-7. (ProCi	2 weeks placebo a 491 rug, then 2 weeks randomized, plac te no. 795222) 15* 21 19	and vice-vers 245 s placebo and	d vice-versa	
2 Waldinger, M. D with paroxetine,	period Overall/ entire time period O., Zwindermal sertraline, an Baseline 1 2 3 4 5	precoitus PE patients – crossover de 25 mg taken 4 hr precoitus Normal controls - crossover n, A. H., Olivier, B. Antidepred nefazodone. J Clin Psycho Placebo morning and evening	sign, no washout – 2 2 weeks r design, no washou essants and ejaculati opharmacol. 2001 Ju	2 weeks drug, then 2 4 t period - 2 weeks d ion: a double-blind, in; 21: 293-7. (ProCi	2 weeks placebo a 491 rug, then 2 weeks randomized, plac te no. 795222) 15* 21 19 19 24 19	and vice-vers 245 s placebo and	d vice-versa	
2 Waldinger, M. D with paroxetine,	period Overall/ entire time period O., Zwinderman sertraline, an Baseline 1 2 3 4	PE patients – crossover de 25 mg taken 4 hr precoitus Normal controls - crossover n, A. H., Olivier, B. Antidepred nefazodone. J Clin Psycho Placebo morning and evening Placebo morning and	sign, no washout – 2 2 weeks r design, no washou essants and ejaculati opharmacol. 2001 Ju	2 weeks drug, then 2 4 t period - 2 weeks d ion: a double-blind, in; 21: 293-7. (ProCi	2 weeks placebo a 491 rug, then 2 weeks randomized, placete no. 795222) 15* 21 19 19	and vice-vers 245 s placebo and	d vice-versa	

Appendix 3-J. Summary Tables: Placebo treatment studies

_		Treatment regimer	1	_	Late	ency in Se	econds	
Treatment Groups	Week	Dosage	Treatment Duration	Number of Patients	Mean	SD	Min	Max
		M. W., Zwinderman, A. H., O fluoxetine, fluvoxamine, paro						
Study a 5a	Baseline 6	Placebo control	6 weeks	9 9	19 29	15 25		
Study b 2b	Baseline 6	Placebo control	6 weeks	12 9	18			
4b	Baseline 6	(no sig. increase in latency) Placebo control (no sig. increase in latency)	6 weeks	3 2	82	27		
U ,	, ,	, M. W., Zwinderman, A. H. Pa jatry. 1994 Sep; 151: 1377-9.		'	ation: a double-b	ind, random	ized, placeb	0-
2	Baseline		6 weeks	9	15		5	90
	3	1 capsule/day for 1 week, 2 capsule/day for 5 weeks		8	20		5	120
	6	1 capsule/day for 1 week, 2 capsule/day for 5 weeks		8	15		5	120
		Latency assessed by patient	İ					
		n, H., Arman, F., Ekmekcioglu . 1999 Jan; 161: 107-11. (Prod		fluoxetine on several	neurophysiologi	cal variables	in patients	with
2	Baseline		4 weeks	20	66	78		
	4	Placebo control		20	288	60		

Appendix 3-K. Summary Tables: Propanolol treatment studies

		Treatment regim	en		Lat	econds		
Treatment Groups	Week	Dosage	Treatment Duration	Number of Patients	Mean	SD	Min	Max
Cooper, A. J., N no. 12387)	Magnus, R. V. A	A clinical trial of the beta b	locker propranolol in p	remature ejaculatio	n. J Psychosom I	Res. 1984; 2	8: 331-6. (P	roCite
1	Baseline			12				
	4	Run-in period	4 weeks		90			
	8	Washout period	4 weeks		90			
	Overall/	•						
	entire time period	120 mg/day			102			

Appendix 3-L. Summary Tables: Sertraline treatment studies

		Treatment regimen			Lat	ency in S	econds	
Treatment Groups	Week	Dosage	Treatment Duration	Number of Patients	Mean	SD	Min	Max
		, E. A., El Gilany, A. H. Asses pot Res. 2001 Feb; 13: 41-5.			therapy and the p	ause-squeez	e technique	: in
3	Baseline Overall/		4 weeks + 2 week washout	31	60		30	90
	entire time period	50 mg/3-5 hrs before coitus, not > 2 x per week		29	180		60	600
5	0: " = 0	Crossover design - 2-week w	· · · · · · · · · · · · · · · · · · ·					
		aran, M., Kupeli, B., Bozkirli, I 30: 611-5. (ProCite no. 12033		eatment of premat	ure ejaculation: a	double-blind	placebo co	ntrolled
1	Baseline 4	50 mg/day	4 weeks	22 22	41 325	13 262		
		emature ejaculation - a compa Cite no. 500002)	arison of treatment	outcome in patient	s with and withou	t erectile dysf	function. Int	J
	Baseline (pts with		6 months	52	46			
1	primary PE) Overall/	50 mg 4 hours prior to	o monuto	02	40			
	entire time period	intercourse for 6 months		52	247.2			
2	Baseline (pts with ED only)		6 months	35	34.6			
	Overall/ entire time period	50 mg 4 hours prior to intercourse for 6 months		35	111.6			
	ponoa	(Pts with ED treated success	fully with sildenafil.	Pts with PE prior t	o ED excluded)			
Kim, S. W., Pai 1999 Sep; 54: 5		term analysis of the effects of no. 12013)	f as needed use of	sertraline at 5 PM t	for the treatment	of premature	ejaculation.	Urology.
1	Baseline 2	50 mg/day	2 weeks	18	23 354	19 252		
	2	50 mg/day at 5 pm on days when intercourse	2 Weeks		334	232		
	4	planned, dose titrated to 100 mg in week 3 if needed	2 weeks		306	228		
	6	50 mg/day at 5 pm on days when intercourse planned	2 weeks		270	162		
		r and safety of fluoxetine, sert Feb; 159: 425-7. (ProCite no.		mine in patients wi	th premature ejac	culation: a do	uble-blind, p	olacebo
3	Baseline			53	46	41		
	Overall/ entire time period	50 mg/day for 1 week, 100 mg/day later	4 weeks	37	256	341		
McMahon, C. G	•	Crossover design - 4-week w premature ejaculation with se		ide. Int J Impot Re	s. 1998 Sep; 10:	181-4; discus	sion 185. (F	roCite
1	Baseline		15 weeks	46	60		0	300
	3 6	25 mg/day Washout			456		0	1200
	9	50 mg/day (4 pts w anejaculation)			786		420	
	12 15	Washout 100 mg/day (10 pts w anejaculation)			984		420	

Appendix 3-L. Summary Tables: Sertraline treatment studies

		Treatment regimen	1	_	Lat	ency in Se	econds	
Treatment Groups	Week	Dosage	Treatment Duration	Number of Patients	Mean	SD	Min	Max
McMahon, C. G Jun; 159: 1935-		f premature ejaculation with so . 12057)	ertraline hydrochlor	ide: a single-blind pl	acebo controlled	l crossover st	tudy. J Urol.	1998
1	Baseline 4 8	50 mg/day Washout	4 weeks	19 19 19	18 204 36			
2 Mendels J. Ca	12 mera A Sike	50 mg/day Crossover design - 4-week ves, C. Sertraline treatment for		18	180 armacol 1995 C	oct: 15: 341-6	(ProCite n	10
12139)	inora, 7t., Onco	o, o. deritaine treatment for	promature ejaculat	on. o omr r oyonopn	iaimaooi. 1000 C	701, 10. 041 0	. (i roone n	
1	Baseline		8 weeks	26	59	69		
	8	121 mg/day mean final dosage * 50 mg/day titrated to 200 m	ng/day during week	22 s 1-3	326	337		
		n, A. H., Olivier, B. Antidepres	sants and ejaculati	on: a double-blind, r	andomized, plac	ebo- controll	ed, fixed-do	se study
with paroxetine,	, sertraline, and	d nefazodone. J Clin Psychop	harmacol. 2001 Ju					Í
3	, sertraline, and Baseline		oharmacol. 2001 Ju 6 weeks					·
•	•	50 mg/day morning and evening		n; 21: 293-7. (ProĆi	te no. 795222)		•	·
•	Baseline	50 mg/day morning and evening 50 mg/day morning and evening		n; 21: 293-7. (ProĆi	te no. 795222) 14*			·
•	Baseline 1	50 mg/day morning and evening 50 mg/day morning and evening 50 mg/day morning and evening		n; 21: 293-7. (ProĆi	14* 25			·
	Baseline 1	50 mg/day morning and evening		n; 21: 293-7. (ProĆi	14* 25 39			·
•	Baseline 1 2 3	50 mg/day morning and evening		n; 21: 293-7. (ProĆi	14* 25 39 34			ŕ
	Baseline 1 2 3 4	50 mg/day morning and evening 50 mg/day morning and		n; 21: 293-7. (ProĆi	14* 25 39 34 43			,
3 Waldinger, M. C	Baseline 1 2 3 4 5 6	50 mg/day morning and evening	6 weeks	n; 21: 293-7. (ProĆi	14* 25 39 34 43 58 50 s on ejaculation:			

Appendix 3-M. Summary Tables: Sildenafil treatment studies

		Treatment regimen			Late	ency in S	econds	
Treatment Groups	Week	Dosage	Treatment Duration	Number of Patients	Mean	SD	Min	Max
·		r, E. A., El Gilany, A. H. Asses: npot Res. 2001 Feb; 13: 41-5.		•	nerapy and the pa	ause-squeez	e technique	in
4	Baseline		4 weeks + 2 week washout	31	60		30	90
	Overall/ entire time period	50 mg 3-5 hrs before coitus, not > 2 x per week		29	900		300	1800
	•	oo, R., Scattoni, V., Briganti, A ketine plus sildenafil in patient				•	, ,	•
2	Baseline	10 mg/day paroxetine for 21 days, followed by 20		40	.35	.05		
	12	mg 3-4 hrs plus 50 mg sildenafil before intercourse 10 mg/day paroxetine for 21 days, followed by 20	3 months	40	4.5	.07		
	24	mg 3-4 hrs plus 50 mg sildenafil before intercourse	6 months	40	5.3	.02		

Appendix 3-N. Summary Tables: Topical anesthetic treatment studies

		Treatment regimen			Late	ency in S	econds	
Treatment Groups	Week	Dosage	Treatment Duration	Number of Patients	Mean	SD	Min	Max
Atikeler, M. K., no. 500006)	Gecit, I., Senol	, F. A. Optimum usage of price	caine-lidocaine cre	eam in premature ej	aculation. Adnrolo	ogia. 2002; 3	34: 356-9. (F	ProCite
	Baseline		Used at least five times (range 5-9)	40	< 1 min			
1	Overall/ entire time period	Placebo cream		10	1.01 min			
2	Overall/ entire time period	Prilocaine-lidocaine cream applied and left in place under a condom for 20 min and removed		10	6.5 min			
3	Overall/ entire time period	Prilocaine-lidocaine cream applied and left in place under a condom for 30 min and removed		10	8.71 min *			
4	Overall/ entire time period	Prilocaine-lidocaine cream applied and left in place under a condom for 45 min and removed		10	**			
		ency 8.71 min in 4 patients, 6 ents complained of erection lo				and delayed	ejaculation	
		n der Werff ten Bosch, J. J. P 37: 244-7. (ProCite no. 50000		on by local penile an	esthsia in an unc	ontrolled clir	nical replicat	tion
1	Baseline			15	1-2 min			
	Overall/ entire time period	Lidocaine-prilocaine cream (1/2 tube) applied for 10 min contained with condom, then removed.	Patients given 5 tubes and 12 condoms.		489	465		
	Overall/ entire time period	3 pts reported inability to reac Coitus without cream or intervention.No patients reported inability	Patients asked to report on 2-3 encounters.	e occasions with cre	eam 115	142		

Appendix 4. Summary Tables:

Articles Selected for Review: Sorted by Author

- **700014** Abdel-Hamid, I. A., El Naggar, E. A., El Gilany, A. H. Assessment of as needed use of pharmacotherapy and the pause-squeeze technique in premature ejaculation. Int J Impot Res. 2001 Feb; 13: 41-5
- 12146 Althof, S. E., Levine, S. B., Corty, E. W., Risen, C. B., Stern, E. B., Kurit, D. M. A double-blind crossover trial of clomipramine for rapid ejaculation in 15 couples. J Clin Psychiatry. 1995 Sep; 56: 402-7
- 105301 Assalian, P. Clomipramine in the treatment of premature ejaculation. J Sex Res. 1988; 24: 213-5
- 795257 Atan, A., Basar, M. M., Aydoganli, L. Comparison of the efficacy of fluoxetine alone vs. fluoxetine plus local lidocaine ointment in the treatment of premature ejaculation. Arch Esp Urol. 2000 Nov; 53: 856-8
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- **12160** Cavallini, G. Alpha-1 blockade pharmacotherapy in primitive psychogenic premature ejaculation resistant to psychotherapy. Eur Urol. 1995; 28: 126-30
- 500002 Chia, S. J. Management of premature ejaculation a comparison of treatment outcome in patients with and without erectile dysfunction. Int J Androl. 2002; 25: 301-5
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- 105302 Damrau, F. Premature ejaculation: use of ethyl aminobenzoate to prolong coitus. J Urol. 1963 Jun; 89: 936-9
- 105303 Eaton, H. Clomipramine (ananfranil) in the treatment of premature ejaculation. J Int Med Res. 1973; 1: 432-4
- 900015 Fein, R.L. Intracavernous medication for treatment of premature ejaculation. Urology. 1990; 34: 301-3
- **12409** Girgis, S. M., El-Haggar, S., El-Hermouzy, S. A double-blind trial of clomipramine in premature ejaculation. Andrologia. 1982 Jul-Aug; 14: 364-8
- **13710** Goodman, R. E. An assessment of clomipramine (Anafranil) in the treatment of premature ejaculation. J Int Med Res. 1980; 8 Suppl 3: 53-9
- 10696 Haensel, S. M., Rowland, D. L., Kallan, K. T. Clomipramine and sexual function in men with premature ejaculation and controls. J Urol. 1996 Oct; 156: 1310-5
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Appendix 4. Summary Tables:

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Total number of articles from all journals: 51

Appendix 5. Summary Tables:

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Total number of articles from all journals: 51