Article

Serenoa repens (saw palmetto) for lower urinary tract symptoms (LUTS): The evidence for efficacy and safety of lipidosterolic extracts. Part III.

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Abstract: Parts I and II of this 3-part series indicated how a global review of both English-language and non-English language papers plus a focus on a lipidosterolic extract of Serenoa repens (LSESr) having a standardized fatty acid profile have together engendered new insights about the biological activity of LSESr vs. LUTS. In this last of a 3-part series, data from the world literature is presented that confirms that LSESr efficacy is the predominant finding in clinical trials. Despite two placebocontrolled clinical trials performed in the U.S. that failed to confirm a benefit of LSESr vs. placebo in LUTS, the global body of the peer-reviewed literature attests not only to efficacy but also to safety. Results will be presented of important trials that compare LSESr to alpha-blockers such as tamsulosin (Flomax®) as well as to 5α -reductase inhibitors such as finasteride (Proscar®) that demonstrate consistent findings of near equivalency between LSESr and these pharmacologic agents. Studies relating data indicative of an additive effect or synergy between LSESr and tamsulosin will be presented as well. The heightened effectiveness of LSESr in men with severe LUTS vs. moderate LUTS expands the importance of our scrutinization of the global literature concerning LSESr. Of great consequence are the contributions of non-English language peer-reviewed publications that have consistently provided evidence of LSESr efficacy in treating LUTS/BPH. These peer-reviewed articles have shown that the effect of LSESr is not that of a placebo. Finally, a comparison of the LSESr extraction products used in the treatment of LUTS, and a discussion of the milieu factors that affect the natural history of LUTS and influence the outcome of clinical trials complete this sedulous analysis of LSESr vs. LUTS.

Keywords: lower urinary tract symptoms; LUTS; benign prostatic hyperplasia; BPH; saw palmetto; Serenoa repens; phytotherapy; lipidosterolic extract of Serenoa repens (LSESr); hexanic extract of Serenoa repens (HESr)

1. LSESr and the Placebo Effect: Is There a Resolution?

How Can We Address the Negative Clinical Trials of LSESr vs. Placebo in Male LUTS?

At least 48 systematic reviews and meta-analyses on Serenoa repens have been analyzed as part of due diligence in scrutinizing 190 studies involved in this report. The Willetts study from Australia and the STEP and CAMUS studies from the U.S. are the three major reports presenting negative findings on LSESr efficacy vs. LUTS. The *S. repens* Treatment for Enlarged Prostates (STEP) [1] and Complementary and Alternative Medicine for Urological Symptoms (CAMUS) trials [2], were randomized, placebo-controlled trials. The findings of STEP and CAMUS contributed to the negative assessment of the efficacy of Serenoa repens later reported in the Cochrane 2012 meta-analysis [3]. Note that the expansions of the acronyms for both STEP and CAMUS are misleading. In the STEP study, the mean prostate size was 34.7 cc, so the title of the study dealing with treatment for enlarged prostates is incorrect. This study randomized a total of 216 men with moderate-to-severe BPH to treatment with placebo (104 patients) or to a supercritical carbon dioxide extraction of Serenoa repens (sCESr) (102 patients) using a dose of 160 mg bid [1].

The clinical endpoints of the STEP and CAMUS trials along with additional information are shown in Table 1.

Table 1. The Bent (STEP) and Barry (CAMUS) studies are the preeminent placebo-controlled trials with negative results.

Author	Year	Extraction	Pt.	Study	IP	SS	Q	οL	Qma	x	Fatty acids%	
(senior)	rear	process	#	(mo)	Δ*	%	Δ	%	Δ	%	ratty actus /o	
Bent	2006	CO ₂	102	12	-0.7	4.3			+ 0.42	3.7	92.1 TFA	
Barry	2011	Ethanol	151	18	-2.2	15.0					54.1 FFA	

^{*} values rounded off to one decimal point

The Barry study is a negative study; the placebo group had Δ in IPSS of -2.99 or 20% improvement.

IPSS, International Prostate Symptom Score; QoL, quality of life; Qmax, peak urinary flow (ml/s); Δ, mean change; mo., months; #, number; Pt., patients; %, percent; TFA, total fatty acids; FFA, free fatty acids.

In STEP, patients were stratified into two groups using the AUA-SI with moderate LUTS being a score of 8-19 vs. severe LUTS, with a score of 20-35. Both the treatment and placebo groups had a decrease in the AUA-SI of approximately -1.5 during the one-month run-in phase. At 12-months post-randomization there were no significant differences in AUA-SI or urinary flow rate. The AUA-SI decreased -0.68 points from a baseline of 15.7 in patients receiving sCESr vs. a decrease of 0.72 points from a baseline of 15.0 in the placebo group. Similar non-significant changes for sCESr and placebo in the BPHII (BPH Impact Index) were seen from baseline to study end (-0.33) vs. (-0.09), and for Qmax (+ 0.42 ml/s) vs. (- 1.01 ml/s), respectively (Table 1). STEP evaluated a sCESr product that had never been used in a prior study or assessed in a subsequent study. Perhaps STEP is valid, or perhaps the product used has a lower quality profile than that of other LSESr products that have demonstrated efficacy in the 60 peer-reviewed studies of LSESr vs. LUTS/BPH.

The CAMUS study, published in 2011, was not an investigation of the efficacy of complementary medicine, as per the title, but instead of a particular ethanolic extract of Serenoa repens (EESr) having the brand name Prostamol Urgenin Uno®. This was the identical EESr used 14 years earlier by Derakhshani et al. [4], but surprisingly not discussed by the CAMUS authors [2]. CAMUS randomized 306 men with LUTS to placebo (n=155) vs. Prostamol Urgenin Uno (n=151) over a study duration of 72 weeks (1.5 years). Eligibility criteria included an AUA-SI between 8 to 24, and a Qmax ≥ 4 ml/s [2]. The daily dose of the EESr was escalated every 24 weeks, from 320 mg to 640 mg and to 960 mg/d. Results showed a decrease in AUA-SI from 14.69 to 11.70 (20% improvement) for placebo vs. 14.42 to 12.22 (15% improvement) for the EESr product (not statistically significant) [2,5] (Table 1). In the Derakhshani 1997 study, there were 1461 patients from 357 practices in Germany that were assessed for IPSS at the end of three months. Therefore, almost ten times the number of patients were assessed in Derakhshani 1997 vs. CAMUS 2011, but with strikingly different results. In Derakhshani 1997, the mean decrease in IPSS after three months was -7.4 points, representing a 40.4% improvement. The QoL improved by 45.9% (n=1461) and the Qmax by +3.7 ml/s (30.8%)(n=1277) (Table 2). The change in IPSS of -7.4 in the Derakhshani study far exceeds the threshold of -3 points cited by many authors as defining a significant therapeutic response. There is no obvious explanation to reconcile these significant differences in outcomes. In the course of this global analysis of LSESr vs. LUTS, we found a total of 58 peer-reviewed articles that met our criteria for evaluability; these included the CAMUS and the Derakhshani studies. The results of the mean changes in IPSS, QoL and Qmax for the 55 positives of the 58 total reports indicate significant improvements in all parameters. This will be discussed in detail in the sections that follow.

Table 2. Endpoints in IPSS, QoL and Qmax for Barry and Derakhshani using Prostamol Urgenin Uno for intervention.

Author	Year	Pt.	Study	IPSS	QoL	Qmax	Fatty acids %
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(senior)		Extraction process	#	(mo.)	Δ*	%	Δ	%	Δ	%	
Barry	2011	Ethanol	151	18	-2.2	15.0					54.1 FFA
Derakhs hani	1997	Ethanol	1461	3	-7.4	40.4	-1.61	45.9	+3.7	30.8	54.1 FFA

A striking difference in endpoints in IPSS, QoL and Qmax in two studies [2,4] that used the identical ethanolic extraction product of Serenoa repens.

IPSS, International Prostate Symptom Score; QoL, quality of life; Qmax, peak urinary flow (ml/s); Δ, mean change; mo., months; #, number; Pt., patient; %, percent; TFA, total fatty acids; FFA, free fatty acids.

The bottom line is that STEP and CAMUS lessened enthusiasm for the use of Serenoa repens in the United States. Scrutiny of all publications related to both trials does not disclose any obvious shortcomings to account for the lack of efficacy of LSESr in the study participants.

2. Therapeutic Comparator Studies of LSESr vs. LUTS

2.1. HESr not inferior to tamsulosin or to finasteride

In contrast to findings in STEP and CAMUS, a consistent benefit has been observed in controlled trials of HESr vs. placebo or therapeutic comparators. The most frequently studied HESr has been Permixon®, and several trials have compared the therapeutic activity of Permixon with that of an alpha-blocker such as tamsulosin, or to a 5α -reductase inhibitor like finasteride (Table 3).

Table 3. Clinical evaluation of HESr vs. Tamsulosin (Flomax®) in four studies and HESr vs. Finasteride (Proscar®) in one study.

Senior Author	Study Duration	C L., J., A	Patients (#)a	II	PSS .	Q	oL_	Q	max
Year, Ref. (#)	(mo.)	Study Arm	ratients (#)"	Δ	%	Δ	%	ml/s	%
Debruyne 2002 [6]	12	HESr Tam	350 354	-4.4 -4.4	28% 29%	NR NR	NR NR	+1.9 +1.8	17% 16%
Hizli 2007 [7]	6	HESr Tam	20 20	-6.1 -4.6	34% 28%	-2.6 -2.1	62% 60%	+3.2 +3.7	34% 35%
Latil 2015 [8]	3	HESr Tam	83 86	-4.5 -6.5	25% 39%	-0.9 -1.3	23% 34%	+1.65 +2.13	15% 20%
Alcaraz 2020 [9]	6	HESr Tam Combo	262 263 184	-5.4 -5.7 -7.2	29% 30.5% 36.9%	-2.5 -2.6 -2.2	65.8% 66.7% 55%	+3.1‡ +2.9 +2.0	23% 24% 16%
Carraro 1996 [10]	6.5	HESr Fin	467 484	-5.8 -6.1	37% 39%	-1.38 -1.51	38% 41%	+2.7 +3.2	25% 30%

^a Number of patients at study end

One such investigation in LUTS therapy involving Permixon vs. the alpha-blocker tamsulosin (Flomax®) was the PERMAL 12-month study reported by Debruyne in 2002 [6]. This large study of over 700 men, with an IPSS eligibility criterion of > 10, disclosed a

^{*} values rounded off to one decimal point

[‡] Number of patients for Qmax were 49, 37 and 56 for HESr, tamsulosin and the combination, respectively.

Δ, mean change; -, negative change; #, number; %, percent change; +, positive change; AUA, American Urological Association; IPSS, International Prostate Symptom Score; LSESr, lipidosterolic extract of *Serenoa repens*; LUTS, lower urinary tract symptoms; ml/s, milliliters per second; NR, not reported; mo., month; QoL, quality of life; Qmax, peak urinary flow (ml/s); Ref., reference citation; Tam, tamsulosin.

decrease in the IPSS of -4.4 points in both the Permixon and tamsulosin groups, with almost identical percentage improvements of 28% and 29%, respectively (Table 3). The changes in the Qmax for Permixon vs. tamsulosin were + 1.8 vs. +1.9 ml/sec, respectively. After three months of treatment, 34% of the patients on Permixon had an improvement in Qmax of at least 3 ml/s that persisted at 12 months. Similarly, at three months, 35% of the tamsulosin group improved the Qmax to at least 3 ml/s, and at 12 months 37% of the tamsulosin group had this response. The results of Permixon vs. tamsulosin are almost identical.

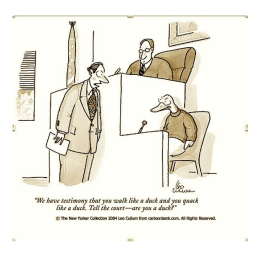
Five years after the PERMAL study, Hizli et al. [7] reported the results of a 6 months investigation of HESr vs. tamsulosin. They used Prostagood®, which for all practical purposes is identical to Permixon. This study involved three arms: HESr vs. tamsulosin vs. the combination of both agents, with 20 patients in each cohort. In this open-label study, HESr was given as 320 mg/d and tamsulosin 0.4 mg/d, with eligibility criteria of IPSS \geq 10, Qmax of < 15 ml/s, a gland volume of \geq 25 cc, and a PSA of \leq 4. At 6-months, the IPSS changes were -6.1, -4.6, and -4.9, for HESr, tamsulosin and the combination, respectively. The percentage improvements were 34%, 28% and 31%, respectively. Qmax changes were +3.2, +3.7 and +4.2 for percentage improvements of 34%, 35% and 42%, respectively. QoL results showed improvements of -2.6, -2.1 and -2.2, with respective percentage improvements of 62%, 60% and 63% (Table 3).

Another comparative study was done by Latil, et al. in 2015. They performed a randomized, double-blind study comparing HESr at 320 mg/d in 83 patients vs. tamsulosin 0.4 mg/d in 86 patients over three months [8]. The average IPSS score with Permixon decreased from 17.7 at baseline to 13.2 (-4.5) on Day 90. In comparison, the respective values for tamsulosin were 16.8 vs.10.3 (-6.5) (Table 3).

In 2020, Alcaraz et al. [9] reported their follow-up to the QUALIPROST study of 2016 [11] that compared HESr (Permixon) vs tamsulosin vs. a combination of the two agents. The combination arm results were statistically superior at p < 0.001. The QoL and Qmax results are shown in Table 3.

A comparison of HESr with the 5-alpha reductase inhibitor finasteride was done in 1996 by Carraro et al. [10]. This trial evaluated 951 men over 26 weeks to ascertain the efficacy of Permixon (n=467) vs. finasteride (n=484) using IPSS as the primary endpoint. The baseline IPSS was 15.7 ± 5.8 vs. 15.7 ± 5.7 , for Permixon vs. finasteride, respectively. At 26-weeks, the IPSS decreased by -5.8 and -6.1, representing improvements of 37% vs. 39%, respectively. The onset of action for both Permixon and finasteride was as early as six weeks after initiating treatment and was associated with a similar degree of improvement in IPSS, with both treatment approaches improving by 22% (p < 0.001). At 26 weeks, further improvement approached nearly 40% for both treatments. This change in IPSS over time in response to LSESr is important when looking at the results of studies of short duration, (i.e., 4 to 6 weeks). Regarding QoL, 70% of patients reported an improvement at 26-weeks with the QoL measurement (question 8 of the IPSS) dropping from -3.63 to -2.25 with Permixon, and from -3.66 to -2.15 with finasteride. These represent improvements of 38% and 41%, respectively. Qmax at baseline was 10.6 ml/s for Permixon and 10.8 ml/s for finasteride, and at 26-weeks was +2.7 ml/s and + 3.2 ml/s, respectively, reflecting 25% improvement for Permixon vs. 30% improvement for finasteride (Table 3). Comparing side effects, patients receiving finasteride experienced a statistically significant deterioration in sexual function vs. those receiving Permixon. This difference was noted from the first follow-up at six weeks and continued to be significant at 26 weeks [10].

These 5 studies of HESr vs. a prescription drug comparator show consistent findings relating to the efficacy of LSESr vs. LUTS. Together, they involve a total of 2,573 patients, and with the clear clinical endpoints of IPSs, QoL and Qmax. This represents a comparable degree of improvement in the absolute values and percentages between HESr, the alpha-blocker (tamsulosin), and a 5-alpha reductase inhibitor (finasteride). While it is true that these are not placebo-controlled studies, we would have to conclude that LSESr is either an active agent vs. LUTS or that tamsulosin and finasteride are no better than a placebo. In my half-century in medicine, I would invoke the "duck principle."



2.2. LSESr Efficacy is Greater in Severe vs. Moderate LUTS.

The greatest improvement in IPSS using a hexanic extract of Serenoa repens (HESr) was observed in those patients with severe lower urinary tract symptoms (LUTS), defined as an IPSS of 20 to 35. In a follow-up to the 2002 PERMAL study [12], a subset of patients with a baseline IPSS of >19 had a decrease of -7.8 points in the Permixon group (65 patients) vs. a -5.8 decrease in the tamsulosin group (49 patients). The corresponding mean percentage decrease in the total IPSS was 35.2% vs. 25.0%, respectively [12]. Further analysis showed that patients with the higher baseline IPSS (>21) had still greater improvement with Permixon and tamsulosin with average IPSS of -9.3 and -6.0 compared to -6.9 and -5.5 found in patients with a baseline IPSS of 20-21, respectively. Flow symptoms (aka obstructive, voiding) symptoms improved more than storage symptoms (aka irritative, filling) symptoms for groups having an IPSS of 20-21 as well as an IPSS >21 with a similar pattern of response for Permixon and tamsulosin. These findings have been extracted from the text and graphs in the Debruyne 2002 and 2004 papers and are shown in Table 4.

Table 4. Comparative changes in IPSS with Permixon vs. tamsulosin from PERMAL (adapted after Debruyne 2002 and Debruyne 2004).

	PERM	AL 2002	PERM	AL 2004
IPSS Value	Δ (%) IPSS	Δ (%) IPSS
_	Permixon	Tamsulosin	Permixon	Tamsulosin
>10	-4.4 (28%)	-4.4 (29%)		
>19a			-7.8 (35%)	-5.8 (25%)
= 20-21 (all)			-6.9	-5.5
= 20-21			-2.5	2.0
irritative			-2.5	-2.0
= 20-21		_	-4.4	2.5
obstructive			-4.4	-3.5
>21 (all)			-9.3	-6.0
>21 irritative			-3.5	-1.9
>21 obstructive		_	-5.8	-4.1

^a For patients with a baseline IPSS >19, Permixon was superior to tamsulosin (p = 0.051). When comparing Permixon to tamsulosin, the greatest change in IPSS (e.g., \geq 9 points) was seen in patients receiving Permixon (41.5%) vs. 25.4% of patients receiving tamsulosin.

 Δ , mean change; -, negative change; %, percent change; +, positive change; IPSS, International Prostate Symptom Score.

Brown and Emberton endorsed the findings of the Debruyne 2004 paper with the following editorial comments. "There is no reason not to take this study seriously. It was part of a European multicentre large-scale study that was well designed and thought out." "Overall, it appears that phytotherapy is as valid a pharmacotherapy as α -blockers and 5α -reductase inhibitors in the management of men with BPH/LUTS. Indeed, it may have less adverse effects, be better tolerated, and cheaper." Emberton noted that in the UK phytotherapy cannot be prescribed and that urologists should be aware and informed about phytotherapy as it will inevitably become part of the standard medical therapy for men with BPH/LUTS. "The previous lack of standardization of herbal remedies that once prevented doctors from recommending these products is now much improved" [12].

The Breza study in 2005 from Slovakia evaluated 596 patients who received an ethanolic extract of Serenoa repens (EESr) at a dose of 320 mg/d [13]. In the total population, the change in IPSS over one year was -5.89 (35.9%), with 84% of patients experiencing more than a 3-point drop. In a subgroup of 150 patients with a mean baseline IPSS of 23.3 (range 20-33), the mean IPSS decreased to a post-treatment value of 15.5 or -7.8 points reflecting a 33.5% improvement. However, as Barry et al. pointed out, the change in absolute values in the IPSS may not be as valid as the percentage change in measurements of the IPSS [14]. This appears to be a logical criticism, and if true, the percentage changes seen in the Breza study are possibly not of statistical significance. In the Debruyne 2004 study, percentage changes could not be calculated for the IPSS subgroups 20-21 and >21, while they could for the IPSS subgroups >10 vs. >19. Therefore, for Debruyne 2004, only in these subgroups (>10 vs. >19) can it be concluded that the greatest changes in IPSS percentage and absolute values occur in those patients with the more severe baseline values.

The 2020 follow-up to the QUALIPROST study was mentioned earlier. An analysis of response to Permixon vs. tamsulosin vs. the combination of the two in men presenting with IPSS baseline values of > 19 indicated a significant improvement in the combination arm over either monotherapy arms in men with an IPSS of >10 but < 19. In this cohort of severe LUTS, the absolute change in IPPS was -7.8 for HESr, -8.0 for tamsulosin, and -10.1 for the combo. The percentage improvements, respectively, were 32.5%, 34.2% and 42.1%. For all patients with any IPSS >10, the respective results were -5.4, -5.7 and -7.2, with percentage improvements of 29%, 30.5% and 36.9%. In this study and others, it would be important to compare the changes in IPPS in the cohort of patients with severe LUTS (20-35 IPSS) vs. those with moderate LUTS (8-19 IPSS), rather than comparing the severe cohort with the total group of patients. The study performed by Eickenberg et al. in 1997 made a comparison between men with a baseline IPSS of ≤ 18 vs. ≥ 19. In that study, 6,967 patients were treated with an EESr (Sita®) at a dose of 320 mg/d for 6 months. At the study end, the IPSS was -8.0. A subgroup analysis based on the IPSS categories ≤ 18 vs. ≥ 19 revealed the same mean percentage improvement of 41% at study end [15]. This issue of degree of efficacy of LSESr based on the severity of the baseline IPSS warrants further scrutiny, and hopefully the corresponding author of the QUALIPROST study will clarify this issue by comparing the severe with the moderate LUTS cohorts. Notwithstanding, the 2020 QUALIPROST publication [9] and its supplement are valuable contributions and raise the possibility of a synergistic effect of LSESr with the alpha-blocker tamsulosin.

3. Peer-reviewed Evaluable Studies of LSESr vs. LUTS

3.1 Fifty-five of fifty-eight Evaluable Studies Indicate Efficacy

31 English-language and 27 non-English-language peer-reviewed publications were identified and were evaluable. Considering only peer-reviewed evaluable English-language papers on LSESr vs. the endpoints of IPSS, QoL, and Qmax, data was extracted (SBS) from 31 publications for analysis. Not all studies reported all three endpoints. Of the 31 studies, Bent 2006 (STEP) [1], Barry 2011 (CAMUS) [2], and Willetts 2003 [16] represent

the three negative publications. The remaining 28 studies (90%) show results that are not consistent with a placebo effect (Table 5).

Table 5. Thirty-one LSESr vs. LUTS English-language studies meeting evaluability criteria. The three negative studies are shown in red font. Robert 2015 & Latil 2015 are two separate papers, but both report the same data. Therefore, the data from Robert was arbitrarily eliminated from the final analysis.

Author	Year	Extraction	Pt. #	Duration	IP	SS	Q	oL	Qn	nax	% fatty
(senior)	rear	(method)	11.#	(months)	Δ	%	Δ	%	Δ	%	acids
Carraro (1)	1996	Hexane	467	6	5.8	37.0	1.38	38.0	2.7	25.0	80.7
Stepanov (2)	1999	Hexane	92	3	6.4	33.3	1.0	26.0	1.6	17.9	80.7
Al-Shukri	2000	Hexane	57	2.25	2.2 ‡	26.8	0.6	18.2	0.7	6.0	80.7
Debruyne (3)	2002	Hexane	704	12	4.4	28			1.9	17.0	80.7
Giannakopo ulos (4)	2002	Hexane	100	6	8.0	40	0.56	16.5	3.67	40.0	80.7
Pytel (5)	2002	Hexane	116	24	5.3	42	1.31	40.0	1.19	10.0	80.7
Debruyne (6)	2004	Hexane	124	12	7.8	35.0	1.2	29.0	1.2	11.0	80.7
El-Demiry (7)	2004	Hexane	190	6	11.4	51.0			4.4	45.4	80.7
Djavan*	2005	Hexane	88	24	1.0 ‡	17.0	0.4	19.0	1.75	15.0	80.7
Hizli (8)	2007	Hexane	20	6	6.1	34.0	2.6	62.0	3.2	34.0	80.7
Giulianelli	2012	Hexane	591	6	5.6	31.5			3.0	28.0	80.7
Latil (9)	2015	Hexane	83	3	4.5	25	0.9	23.0	1.65	15	80.7
[Robert] (10)	2015	Hexane	[102]	[2]	[4.5]	[25.4]					80.7
Alcaraz	2016	Hexane	1713	6	3.8	25					80.7
Totals Hexane		n=13	210	8.0	-5.6	32.7	-1.1	30.2	+2.3	22.0	80.7
Romics	1993	Ethanol	31	12					4.3	39.0	54.8
Bach (11)	1996	Ethanol	315	36		73.0			6.1	45.5	54.8
Kondas (12)	1996	Ethanol	38	6					4.08	39.0	54.8
Gerber (13)	1998	Ethanol	46	6	7.6	37.0			-0.7	- 5.0	40.0
Barry **	2011	Ethanol	151	18	2.2 ‡	15.0					54.1
Gerber (14)	2001	Ethanol	85	6	4.4	26.0	0.7	21.0	1.0	10.0	40.7
Sinescu (15)	2011	Ethanol	120	24	5.5	40.0	1.8	50.0	5.6	54.0	59.3
Argirovic (16)	2013	Ethanol	265	6	6.1	33.9	1.6	38.0	3.2	34.0	59.3
Cai	2013	Ethanol	47	3	3.1 ‡	18.3			+0.5	4.0	
Suter	2013	Ethanol	69	2	7.5	52					95
Saidi	2019	Ethanol	40	12	2.1 ‡	18.1			0.76	5.8 n.s	59.3
Vinarov	2019	Ethanol	30	180	6.0	50	3.0	60.0	5.0	45.0	59.3
Ye	2019	Ethanol	159	6	4.4	29	1.11	26.0	4.09	36.0	68.4
Totals ETOH		n=13	488.6	24	-5.0/-5.3	37/39	-1.8	44	+3.2	28	58.4
Braeckman	1994	Carbon dioxide	305	3	6.6	34.7	1.54	41.6	2.41	26.4	74.0
Braeckman ‡	1997	Carbon dioxide	67	12	10.2	60.0	1.5	41.7	2.6	23.8	74.0
Braeckman (17)	1997	Carbon dioxide	125	3		64.0				29.8	74.0
Willetts	2003	Carbon dioxide	46	3	1.1	7.8	0.49	13.0	2.35		
Bent	2006	Carbon dioxide	102	12	0.7	4.3			0.42	3.7	92.1
Totals CO2		n=5	260.8	5.5	4.7/8.4	39/48	1.2/1.5	32/42	2.7/3.3	25/30	69/66
Totals All		31									

^{*} Djavan study on prevention of progression of LUTS from mild to greater than mild; Permixon vs. WW ** Barry study is a negative study and the placebo group had Δ in IPSS of -2.99 or 20% improvement.

[‡] Braeckman study used Prostaserene® as LSESr. QoL not from IPSS but only a rating scale. Only 67 patients completed the study, with 34 patients receiving LSESr at 160 mg bid and 33 patients receiving 320 mg/d.

⁽¹⁾ Carraro study of Permixon vs. finasteride. HESr showed equivalent efficacy to 5ARI with fewer side effects (2) Stepanov study comparing Permixon at 160 mg bid vs. 160 mg x2 once a day. Average results used.

⁽³⁾ Debruyne 2002 study of Permixon vs. tamsulosin study with 4-week run-in phase. No significant differences in the effect of Permixon vs. tamsulosin 0.4 mg/d.

- (4) Giannakopoulos study compared 160 mg bid vs. 160 tid. The results shown are the average of both findings. Qmax with 480 mg/d +4.54 vs. +2.8 for 320 mg/d.
- (5) Pytel reported that 46-69% of patients reported improvement in obstructive and irritative symptoms from month-6 to the study's end at 2 years.
- (6) Debruyne 2004 subset analysis of high > 19 IPSS patients with randomization between Permixon vs. tamsulosin.
- (7) El-Demiry is an abstract but with solid data.
- (8) Hizli 2007 study comparing Permixon vs. tamsulosin vs. Permixon + tamsulosin. All groups with no significant differences in efficacy; Permixon + tamsulosin did not increase efficacy.
- (9) Latil study comparing Permixon vs. tamsulosin and correlations with inflammation.
- (10) Robert 2015 data is the same as Latil 2015, so not counted as a separate study, but shown in the table.
- (11) Bach 1996 3-year study quantitated nocturia, frequency, and incomplete emptying. Nocturia improved 73%, and no nocturia or nocturia x1 increased from 33% to 85%. Improvements in frequency and incomplete emptying of 54% and 76%, respectively.
- (12) Kondas used Strogen Forte. The authors stated they measured IPSS but did not report results.
- (13) Gerber 1998 noted improvement at 2 months. At 6 months, 46% of patients with ≥ 50% (21/46) improvement.
- 14) Gerber 2001 study a with one-month placebo run-in for all patients.
- (15) Sinescu used Prostamol Uno.
- (16) Argirovic study compared Prostamol Uno 320 mg/d vs. tamsulosin vs. tamsulosin + Prostamol uno; percentage improvements were 33.9% vs. 28.4% vs. 31.4%, respectively for IPSS. Results were 38% vs. 40% vs. 37% for QoL; and for Qmax they were 34% vs. 35% vs. 44.5%, respectively.
- (17) Braeckman 1997 study with calculations done by SBS. For placebo, IPSS improved 25% & Qmax improved 10%.

The Hutchison 2007 study was not shown because it was a group analysis, but it is a valuable study.

If the non-English-language papers are considered, 27 evaluable publications from the peer-reviewed non-English-language literature can be identified (Table 6). All studies indicated the efficacy of LSESr vs. LUTS. The majority of these non-English-language studies were not cited by authors of the major English-language literature on LSESr. The publication dates of these non-English-language publications range from 1983 to 2013. It would seem improbable for the beneficial effects of LSESr to be the result of a placebo effect given the consistency regarding efficacy across so many studies published in different countries over close to 40 years.

Table 6. Evaluable non-English-language papers (27 studies) categorized by extraction method. The mean number of patients, duration of the study, and the key clinical outcome assessments are detailed.

Senior	Ref.	Year	Extraction	Serenoa	Study Duration	IPS	SS	Qo	οL	Qm	ax
Author	(#)	rear	Method	Patients (#)a	(mo.)	Δ	% b	Δ	%	Δ	%
Cirillo-	[17]	1983	Hexane	47	4		56 §			+ 4.55	50 §
Marucco											Ü
Cukier ψ	[18]	1985	Hexane	73	2		33 §§				
Tosto	[19]	1985	Hexane	20	3	-4.95	28 ***				
Pannunzio	[20,21]	1986	Hexane	30	2					+ 5.10	74
Pescatore	[21]	1986	Hexane	30	3					+ 2.50	27
Authie	[22]	1987	Hexane	500	3		78 *				
Ollé Carreras	[23]	1987	Hexane	40	2		68 ‡‡				
Orfei	[24]	1988	Hexane	30	3		50 ^^	-2.17		+ 0.03	0.2
Dathe	[25]	1991	Hexane	49	6					+5.90	49
Aliaev	[26]	2002	Hexane	26	60	-8.80	76	-1.31	53	+4.30	35
Foroutan	[27]	1997	Hexane	592	3	-6.48	38	-1.49	45	+5.85	66
Medeiros †	[28]	2000	Hexane	130	3	-6.50	37	-1.37	39	+1.95	22
Totals (12)											
Derakhshani	[4]	1997	Ethanol	1,047	3	-7.4	40	-1.61	46	+3.70	31
Eickenberg	[15]	1997	Ethanol 96%	6,967	6	-8.0	44	-1.80	38	+ 3.00	23
Redecker	[29]	1998	Ethanol 90%	50	3		48 ^^^			+ 3.40	24
Ziegler ЖЖ	[30]	1998	Ethanol 90%	109	3				36	+3.72	29
Breza	[13]	2005	Ethanol	596	12	-5.89	36	-1.7	54	+ 2.31	19
Aliaev	[31]	2007	Ethanol	50	6	-2.98	26	-1.8	43	+ 1.73	14
Razumov	[32]	2007	Ethanol	30	6	-6.9	43	2.73	68	+2.80	23

Aliaev Ж	[33]	2009	Ethanol	50	24	-4.18	37	-2.2	52	+ 2.66	21
Vinarov	[34]	2010	Ethanol	50	36	-6.0	50	-2.0	50	+ 4.50	39
Aliaev	[35]	2013	Ethanol	38	120	-1.3	12	-1.05	35	+ 3.25	26
Totals (10)											
Mattei ψ	[36]	1990	CO ₂	20	3		55 ^				
Vahlensieck	[37]	1993	CO ₂	1,334	4		47 §§§				
Vahlensieck	[38]	1993	CO ₂	312	3					+ 5.8	52
Fabricius XX	[39]	1993	CO ₂	176	6		39/59				
Bauer‡ψ	[40]	1999	CO ₂	101	6		38				16
Totals (5)											
Mean Across Al	l Studie	s (n=27)									
Hexane extracti	on (n=12	2)		460	12	F 0	-: 24	1.2	477	+3.5	30
Ethanol extracti	on (n=10	0)		400	12	-5.8 ≈ 24 -1.3 47					30
Carbon dioxide	extracti	on (n=5)									

^a The number of patients at study end, or as reported.

 ψ are placebo-controlled studies.

§§ Study done before IPSS; raw data on nocturia.

§§§ IPSS not used. The numbers shown are based on the change in frequency and nocturia (urinations) pre-versus post-S. repens.

**XX Ziegler did not use IPSS, so his reported symptoms were based on % improvement involving weak stream, hesitancy, incomplete emptying, frequency, and nocturia.

XX The Fabricius 1993 study reported decreases in frequency and nocturia of 39%, and 59%, respectively. This resulted in a mean percent change from 23.5% to 24.4% for IPSS from these surrogate assessments, yielding a mean improvement across all 27 studies of about 24%

- † QoL scale 6 (worst) to 1 (best) rather than 6 (worst) and 0 (best).
- ≈ approximately; Δ mean change; negative change; # number; % percent change; + positive change; CO₂, carbon dioxide; IPSS International Prostate Symptom Score; mo. month; QoL quality of life; Qmax peak urinary flow (ml/s); Ref. citation reference.

In addition, head-to-head studies, placebo-controlled and non-placebo-controlled studies of LSESr, showed similar, if not superior, efficacy when compared with the 5α -reductase inhibitor finasteride or with the α -blocker tamsulosin as shown earlier in Table 3. If LSESr is a placebo, then so too are finasteride and tamsulosin. This is what Frater-Schröder concluded in the 2009 editorial entitled "when a=b and a=c, then b=c" [41]. This editorial was directed at the Cochrane 2012 meta-analysis [3] and its criticism of the Carraro 2006 [10] and Debruyne 2002 [6] studies. In Frater-Schröder's opinion, "Quasi-scientifically-based reports like this Cochrane review weaken the importance and value of phytotherapy in the awareness of experts and the general public" [41]. At the time this editorial was published, Frater-Schröder was the co-secretary and an active member of the scientific committee of ESCOP, an organization that described the results of some Serenoa studies, but whose committee report never came to conclusions about the efficacy of LSESr [42].

3.2. Previous Key Assessments of the Literature (Novara and Vela-Navarrete

Two key meta-analyses reviewed the efficacy and safety of the HESr (Permixon) in the treatment of LUTS [43,44]. The Novara 2016 meta-analysis identified seven randomized and controlled clinical trials each conducted with Permixon and concluded that

^b The clinical endpoints of IPSS, QoL and Qmax are rounded off to two significant digits.

[‡] Placebo-controlled and double-blinded, randomized.

^{*} Study before IPSS use; nocturia, frequency, and urgency improvements were 82%, 67%, and 85.3%, respectively (average improvement 78.1%); average complete resolution of these symptoms was 43.5%.

^{**} Study done before IPSS; symptoms evaluated pre- and post-treatment included frequency, nocturia, urgency, weak stream.

^{***} Study done before IPSS; authors used a unique point scoring to evaluate frequency, nocturia, incomplete emptying, weak stream. § Study done before IPSS; raw data on nocturia; the study also included Qmax results.

[^] Using scores from frequency, nocturia, and incomplete emptying.

^{^^} Using scores from frequency, nocturia, urgency, weak stream, and straining at the beginning and end of the study.

^{^^^} Only able to evaluate nocturia with before and after scores.

 $[\]ddagger$ IPSS not used. The number shown is based on the change in frequency with complete resolution in 27/40 patients.

Ж Aliaev 2009 is a 2-year extension of the 6-month 2007 paper.

Permixon improved peak urinary flow rate (Qmax) and decreased nocturia compared with placebo. However, 6 of 7 studies cited by Novara were considered (SBS) non-evaluable for the following reasons. Three studies had less than 20 patients at the end of the study [45-47], two studies had a duration of only four weeks [48,49], and one study presented unclear data [50]. Two additional studies reviewed by Novara concluded that Permixon relieved LUTS comparable to tamsulosin [6,8]. These latter two studies did meet the requirements for evaluability (Debruyne 2002 and Latil 2015, Table 3). Finally, in two other studies reviewed by Novara, Permixon and tamsulosin were used as therapy in combination, but Permixon was not evaluated as monotherapy, so efficacy could not be concluded [51,52]. In contrast to tamsulosin and finasteride, Permixon had little impact on sexual function and the safety profile of HESr was comparable to placebo [43].

Similar to the Novara meta-analysis, Vela-Navarrete reviewed 27 studies using HESr as monotherapy in patients with LUTS at the standard dose of 320 mg/d [44]. This 2018 meta-analysis included 15 randomized and controlled studies and 12 observational studies conducted under conditions of routine clinical practice. The authors concluded that the standardized HESr was well-tolerated and effective for the long-term treatment of LUTS/BPH and that HESr reduced nocturia and improved peak urinary flow rate vs. placebo. Moreover, patients receiving the standardized HESr had a statistically significant mean improvement in the IPSS, decreasing from baseline by -5.73 points (p < 0.0001), and well above the minimum 3-point improvement cited by Barry 1995 as a threshold for clinical significance [53]. The Vela-Navarrete 2018 review included the 1997 open-label study by Foroutan, et al. [27] conducted in Austria, with 592 patients evaluated over 3 months. This study showed an improvement in IPSS (-6.48; 38%), in QoL (-1.49; 45%), and in Qmax (+5.85 ml/s; 66%). This study was initially missed in this author's (SBS) search of the Serenoa literature; it is an important evaluable study (see Table 6).

An additional non-English-language study not easily discoverable with standard search approaches is the open, multicenter study by Medeiros, et al., published in 2000 in Portuguese [28]. They evaluated 130 patients from 17 urology centers over 3 months. The IPSS was significantly improved (-6.54, 37.5%, p < 0.0001), as was QoL (-1.37, 38.6%, p < 0.0001) and Qmax (+1.95ml/s, 22%, p < 0.0001; Table 6). Medeiros, et al. used a different scale for QoL assessment, scoring 6 (worst) to 1 (best), rather than the established scoring of 6 (worst) and 0 (best).

An English-language trial conducted in Egypt by El-Demiry and published in 2004 as an abstract in the British Journal of Urology International was identified late in our exhaustive search of the Serenoa literature [54]. El-Demiry evaluated 200 patients over 6 months using Permixon 160 mg bid after a 2-week washout period. IPSS, QoL, Qmax, residual volume, prostate volume, and PSA assessments were made after 1, 3, and 6 months of treatment. A total of 190 patients completed the study. Significant improvement was seen in IPSS (-6.6, 30%) at 1 month, and further improvement at 6 months (-11.4, 51%, p < 0.0001). QoL improved by a mean value of 73% at 6 months, and Qmax increased significantly, by +2.8 ml/s at 1 month, up to +3.7 ml/s at 3 months, and further improvement by +4.4 ml/s (45.4%) at 6 months (p < 0.0001; Table 6) [54].

4. In the Final Analysis, the Effect of LSESr vs. LUTS is Not a Placebo Effect

There is no question that the lack of efficacy of LSESr per the STEP and CAMUS double-blind placebo-controlled trials resulted in diminishing physician acceptance of LSESr in the US. This negative impact has been and continues to be compounded by the absence of tight regulations concerning quality requirements and the commercial prevalence of non-standardized Serenoa products in the United States. In other words, the marketplace in the US is flooded with saw palmetto products that range from good to inferior quality. In contrast, LSESr products from hexane, ethanol, or carbon dioxide extraction processes that meet a standardized profile are widely used in Europe, and accordingly, the physician perception of LSESr vs. LUTS is of a significantly higher degree in Europe, Asia, and South America than in the US.

Adding to the complexity of the above issue concerning the quality of Serenoa repens products is that the carbon dioxide LSESr product used in STEP was never evaluated in another trial. If this had occurred, it would have confirmed or refuted the conclusions reached in STEP. Were the results of STEP a fluke in contrast with the results of many other published studies? With the evaluability criteria detailed in Part I, a total of 58 evaluable peer-reviewed studies of LSESr were found (SBS) using the three different extraction methodologies. Ten of these 58 used a lipidosterolic product from carbon dioxide extraction (sCESr). The mean clinical endpoints of IPSS, QoL, and Qmax in these 10 studies were -4.7, -1.2, and +2.7, respectively. It is important to note that these results include both negative studies using carbon dioxide extraction (STEP [1], and Willetts [16]). If these two negative studies are eliminated from analysis, the mean results are -8.4 for IPSS, -1.5 for QoL, and +3.3 ml/s for Qmax, with percentage improvements of 48%, 42%, and 30%, respectively (see Table 7).

Table 7. Summary of mean outcome for evaluable studies grouped by extraction technology for IPSS, QoL, and Qmax. Data are shown for all studies and also for only the positive studies. All 25 HESr studies were positive. The three negative studies were Willetts 2003 (CO₂), Bent 2006 (CO₂), and Barry 20119 (Ethanol).

Extraction	Mean	Mean	Included -]	PSS	Q	oL	Qma	ax	Typical FFA	
Technology	Patients #	Study Duration	Studies	(Δ)	%	(Δ)	%	(ml/s)	%	%	
Hexane n =25	236	8.0 mo.	All Positive	-5.8	35	-1.3	34%	+2.9	29	Min. ≈ 80	
Ethanol n =23		_	All †	-5.0	37	-1.8	44%	+3.0	27	_	
(1 negative)	469	23 mo.	Positive Only	-5.3	39					Min. ≈ 70	
60 . 10			All	-4.7	39	-1.2	32%	+2.7	25		
CO ₂ n =10 (2 negative)	259	5.5 mo.	Positive Only	-8.4	48	-1.5	42%	+3.3	30	Min. ≈ 65-70	

[†] The one negative ethanol study does not significantly alter the IPSS outcome. The free fatty acid (FFA) minimums are typical values for lipidosterolic products (data on file 2021, Valensa International).

The CAMUS 2011 study publication used Prosta-Urgenin Uno, an EESr. In an earlier paper by Derakhshani in 1997 involving 1,047 men, this same product was evaluated over a treatment period of three months and resulted in a mean change in IPSS of -7.4, QoL improvement by 46%, and Qmax increase of 3.7 ml/s [4]. Of the 58 evaluable studies reviewed by this author (SBS), 22 (37.9%) used an ethanol extract product and all showed positive results except for the CAMUS study. The mean clinical endpoints of IPSS, QoL, and Qmax, including the negative CAMUS study, were -5.0, -1.8, and +3.2, respectively. The percentages of mean improvement were 37%, 44% and 58.4%, respectively. If the CAMUS data for IPSS are removed, the IPSS mean results further improve from -5.0 to -5.3 (see Table 7). After examining all extraction processes, including the outcome from negative studies, and including all evaluable peer-reviewed articles published in every language that the benefit of LSESr vs. LUTS is undeniable.

Cochrane 2012 presented results from studies comparing Serenoa repens vs. placebo [3]. Some of these references were considered non-evaluable (SBS) because the patient number at the end of the study was less than 20, and/or the study duration was less than 2 months. Contrary to the position taken by the authors of Cochrane 2012, those non-evaluable studies (SBS) did indicate that Serenoa repens was effective vs. LUTS. Data from these studies considered negative by Cochrane 2012 included a Qmax +3.4 ml/s, a decrease in nocturia with 55% improvement), and a decrease in urgency by 65%. In contrast, the 2018 Vela-Navarrete review and meta-analysis assessed the efficacy of the HESr product Permixon vs. placebo (or comparator) in randomized clinical trials. Of the 15 trials considered in the meta-analysis, seven are placebo-controlled [44]. Vela-Navarrete 2018 used

^{≈,} approximately; ∆, mean change; ¬, negative change; %, percent; +, positive change; CO₂, carbon dioxide; FFA, free fatty acid; IPSS, International Prostate Symptom Score; ml/s, milliliters per second; mo., month; QoL, quality of life; Qmax. peak urinary flow (ml/s).

a random effect model and considered publication bias. The outcome of their review determined that HESr compared to placebo was associated with 0.64 fewer voids/night (95% confidence interval [CI] -0.98 to 0.31, p < 0.001) and an increase in Qmax of +2.75 ml/s (95% CI 0.57 to 4.93; p = 0.01). Figure 2 in the Vela-Navarrete 2018 paper showed a forest plot for Qmax from four studies (Boccafoschi 1983, Emili 1983, Tasca 1985, and Descotes 1995) comparing HESr (n=122) to placebo (n=133). The studies that supported efficacy for a decrease in nocturia and improvement in Qmax were not impacted by study heterogeneity, and no publication bias could be found in the 2018 review and meta-analysis by Vela-Navarrete, et al [44].

A table of the 17 English and non-English-language placebo-controlled studies for Serenoa repens vs. LUTS/BPH is presented in Table 8. For the HESr studies, Vela-Navarrete 2018 did not cite the Mandressi 1983 study [49], which also included a separate Pygeum intervention arm. Boccafoschi 1983, Emili 1983, Mandressi 1983, Champault 1984, Tasca 1985, and Descotes 1995 were all considered by Cochrane 2012. Including clinical studies of LSESr using hexane, ethanol, and CO_2 extraction processes, a robust set of literature for placebo-controlled trials for Serenoa repens exists and confirms that the body of evidence for the efficacy of Serenoa repens does not represent a placebo effect.

Table 8. Summary of 17 placebo-controlled LSESr clinical trials. Studies with italicized senior author did not meet the criteria for evaluability for LSESr for LUTS/BPH.

Senior Author	Year	Study	Extraction	Product	Serenoa Patients (#)	Placebo Patients (#)	Study Duration (mo.)
Boccafoschi	1983	D, P	Hexane	Permixon	11	11	2
Emili	1983	D, P	Hexane	Permixon	15	15	1
Mandressi	1983	D, P	Hexane	Permixon	19	15	1
Champault	1984	D, P	Hexane	Permixon	50	44	1
Cukier	1985	D, P	Hexane	Permixon	71	76	2.5
Tasca	1985	D, P	Hexane	Permixon	14	13	2
Reece Smith	1986	D, P	Hexane	Permixon	33	37	3
Mattei	1990	D, P	CO ₂	Talso Sanofi	20	20	3
Löbelenz	1992	P	Ethanol	Sabal Extract	30	30	1.5
Descotes	1995	D, P	Hexane	Permixon	82	94	1
Braeckman	1997	D, R, P	CO ₂	Prostasere ne	125	113	3
Bauer	1999	D, R	CO ₂	Talso Uno	101 pa	atients*	6
Gerber	2001	D, R	Ethanol	Solaray	39	40	6
Willetts	2003	R, C	CO ₂	Proseren	46	47	3
Bent	2006	D, P	CO ₂	Indena product	102	104	12
Barry	2011	D, P	Ethanol	Prosta- Urgenin Uno	151	170	18
Ye	2019	D, P	Ethanol	Prostess Uno	159	169	6

^{*}Individual numbers not given.

CO₂, carbon dioxide; D, double-blind; mo., months; R, randomized; P, placebo-controlled. Studies with italicized senior author did not meet the criteria for evaluability.

The Boyle 2004 meta-analysis involving 8 randomized clinical trials presented findings consistent with the data presented above. Boyle et al. found that HESr vs. LUTS was associated with a 5-point reduction in IPSS and significant improvements in Qmax and nocturia compared to placebo [55]. As mentioned earlier, of these 17 placebo-controlled

studies, eight were categorized as non-evaluable due to a short study duration (Champault, Descotes, Emili, Mandressi) [46,48,49,56], too few patient numbers (Boccafoschi, Tasca), [45,47] or unclear data (Reece Smith) [50]. Excluding these studies did not alter findings from our review of the evaluable studies in confirming the efficacy of LSESr in improving the key study endpoints: IPSS, QoL, and Qmax. Moreover, of the six studies that were non-evaluable due to low patient number (<20 patients) or short duration (<2 months), clinical endpoints such as urgency decreased by 65%, nocturia decreased by 55%, and Qmax improved by 3.4 ml/s (Table 9). This is further support to the body of literature attesting to LSESr efficacy vs. LUTS.

Table 9. Efficacy of hexanic extract of Serenoa repens (HESr) vs. placebo in six non-evaluable studies. Six of the non-evaluable studies in Cochrane 2012 assessed the efficacy of HESr (Permixon) and reported clinical improvement in symptoms vs. placebo.

Senior Author	Year	Ref. (#)	Serenoa Patients (#)ª	Study Duration (mo.)	Key Results for Serenoa vs. Placebo or Comparator
Boccafoschi	1983	[45]	11	2	Qmax + 4.2 (42%) vs. placebo + 2.1 (20.6%)
Emili	1983	[46]	15	1	Qmax +3.56 (34.5%) vs. placebo + 0.20 (2.2%)
Mandressi	1983	[49]	19	1	Serenoa vs. Pygeum vs. placebo; ↓ urgency 70% vs. 62% vs. 24%; ↓ frequency 30% vs. 22% vs. 10%; ↓ nocturia 42% vs. 38% vs4%
Champault	1984	[56]	50	1	Qmax +2.7 (50.5%) vs. placebo +0.25 (5%); nocturia - 1.53 (49%) vs. placebo -0.48 (15%)
Tasca	1985	[47]	14	2	Qmax +3.3 (25.6%) vs. placebo -0.6 (-5%); nocturia 74.3% vs. 38.7%; urgency 60% vs. 20%; weak stream 50% vs. 16.6%
Descotes	1995	[48]	82	1	Qmax +3.4 (28.9%) vs. placebo + 1.1 (8.9%)
	Mean Acro	oss All St	udies for Clinica	al Outcome	Qmax +3.4; ↓ nocturia 55%; ↓ urgency 65%

^a The number of patients at study end, or as reported.

5. Extract Quality May Affect LSESr Efficacy

Major differences between extraction technology and composition of finished Serenoa products have been identified, and which have been stated to significantly impact the ability of the supplement to ameliorate LUTS [57-60]. Research suggests that there is a "fingerprint" of saw palmetto that represents a quality standardized profile. A key element of this quality standardized profile is the ratio and content of fatty acids, which can vary dramatically across products [59]. Both the EU monograph and USP standards established the minimum level of total fatty acids (TFA) that are needed for a quality Serenoa repens extract. The USP also established ratios of the key fatty acids compared with lauric acid that are required to meet the established chemical profile [61]. The USP stated the chemical profile for a quality Serenoa repens extract would have a minimum of 80% total fatty acids and have a fatty acid composition of oleic acid (30-35%), lauric acid (26-32%), myristic acid (10-12%), palmitic acid (8.5-9.2%), and linoleic acid (4.3-6.0%) [61]. This fatty acid profile distinguishes quality saw palmetto extracts from vegetable oils, adulterated products, and dried saw palmetto berry powders that are deficient in fatty acid amount and/or composition. Key issues remain whether or not the content of total vs. free fatty acids (FFA) of LSESr correlates with efficacy in treating LUTS, or does a particular fatty acid account for LSESr activity, and also, whether or not one extraction process is better than another.

6. The Extraction Process Does Not Correlate with the Efficacy of LSESr Products vs. LUTS

[,] decreased; †, increased; -, negative change; #, number; +, positive change; mo., month; Qmax, peak urinary flow (ml/s); Ref., reference citation; vs., versus.

An analysis of 20 commercially available Serenoa repens products using a gas chromatography-flame identification detector (GC-FID) and gas chromatography-mass spectrometry (GC-MS) showed considerable variability in TFA and phytosterol content among preparations[60]. In another gas chromatography study involving 19 different Serenoa repens mono-preparations, the fatty acid content varied from one-tenth to greater than 4.6-times the fatty acid mg/d dose stated on the supplement product package insert [57]. The mean FFA content in 14 different Serenoa repens products available in Europe has ranged from as high as 80.7% to as low as 40.7% [59]. Additionally, only 9 of the 19 mono-preparations evaluated contained the recommended daily dosage of 320 mg LSESr per day, consisting of 70% to 95% fatty acids (range, 224-304 mg) [57]. With consideration of this huge variability in the quality of Serenoa repens products, the hexanic lipidosterolic extract Permixon has been found to have the highest percentage of FFA, and this finding has been attributed to the therapeutic efficacy of this LSESr.

Studies suggest that hexane, supercritical CO2, and ethanol extraction technologies lead to different fatty acid and phytonutrient profiles. But commercial Serenoa repens extract-containing products made from any of the extraction technologies are said to have demonstrated activity against 5α -reductase and/or an impact on symptoms of LUTS/BPH [43,62]. At a biological level, the pharmacologic activity of 10 lipidosterolic extracts of Serenoa repens differed in the degree to which they inhibited fibroblast proliferation and 5α -reductase Types 1 and 2 [63]. It was the hexanic lipidosterolic extract that most actively inhibited enzyme activity and fibroblast-induced cell proliferation. Data on supercritical CO2 extracts are more limited, but it is recognized that LSESr extracts using ultrahighpressure supercritical CO2 have a fatty acid profile similar to hexanic lipidosterolic extracts. Because ethanol has a different polarity than hexane, this may contribute to the differences in the extract profiles of hexane vs. ethanol lipidosterolic products. Do these differences in fatty acid profiles or biochemical actions translate to marked differences in the clinical efficacy of LSESr vs. LUTS? Our review of the peer-reviewed Serenoa repens literature, with evaluability requirements for LSESr monotherapies, and known extraction modality yielded findings that failed to show any obvious relationships between extract type and the degree of clinical effect. The pooled results of the IPSS, QoL and Qmax for twenty-five, ten and twenty-three evaluable studies using hexane, CO2 and ethanol extraction, respectively, show very similar results. These findings were presented previously (Table 7) but are reproduced below with the mean values of free fatty acid percent, to emphasize that neither FFA content nor extraction modality have any bearing on clinical efficacy (Table 10).

Table 10. All evaluable LSESr studies analyzed by extraction modality and mean free fatty acid (FFA) percent show no clinical correlation with IPSS, QoL or Qmax.

Extraction	Patient #	Mo.	Δ IPSS & Percent		Δ QoL &	Percent	Δ Qmax &	& Percent	FFA %
Hexane (n=25)	236	8.0	-5.8	35%	-1.3	34%	+2.9	29%	80.7
ETOH (n=22)	469	23	-5.0 (-5.3)	37% (39%)	-1.8	44	+3.0	27%	58.4
CO ₂ (n=10)	259	5.5	-4.7 (-8.4)	39% (48%)	-1.2 (-1.5)	32% (42%)	+2.7 (+3.3)	25% (30%)	80 (69)

The upper bolded and italicized numbers include both positive and negative studies, vs. numbers in parentheses indicating only positive results. The negative studies for CO_2 were Bent 2006 [1], and Willetts 2003 [16]. The negative study for ethanol extraction was Barry 2011 [2].

Commented [SS1]: Ms. Wang: I cannot seem to properly align this table. Please assist. Thank you.

Patient #, mean number of patients; Mo., month; Δ, mean change; IPSS, International Prostate Symptom Score; QoL, quality of life; Qmax. peak urinary flow in milliliters per second; FFA, free fatty acid; -, drop in IPPS or improvement in QoL; %, percent; +, improvement change; CO₂, carbon dioxide; ml/s, milliliters per second;

Unfortunately, none of the clinical studies of LSESr products have involved head-to-head comparisons of one extraction process vs. another [64]. Although it would seem apparent that every peer-reviewed study should detail the extraction process and the details of dosing, it is disappointing that some publications omitted such crucial information. Concerning the clinical endpoints of IPSS, QoL and Qmax, when the extraction method is one of the standard LSESr processes, be it hexane, ethanol, or carbon dioxide, there is no evidence of the superiority of one process vs. another. The biologic differences seen in non-clinical studies dealing with various pharmacologic actions in vitro seem to have minimal relevance to what is seen in vivo in human clinical studies. The major challenge is to educate both physicians and the lay public that products labeled as "saw palmetto" or "Serenoa repens" are not equivalent to a standardized lipidosterolic extract of Serenoa repens (LSESr) with a product profile that meets an established definition, and that only the use of the latter is acceptable.

7. Milieu Factors Are Important When Assessing LSESr vs. LUTS

The demographics of the population under study cannot be ignored when evaluating clinical trial results. Strong epidemiologic data demonstrate that lifestyle factors such as obesity, diet, alcohol intake, stress, and physical activity, play a role in LUTS etiology and progression [65-69]. In fact, such lifestyle issues interact with each other and are consequential in the processes of inflammation, aging, and cancer. The patient's medical history, the presence of co-existent chronic inflammation, the proper assessment of diabetes, metabolic syndrome, and details about medications and supplements and possible drug interactions need to be taken into account when determining the clinical efficacy of LSESr vs. LUTS [70-77]. Study design should stratify patients into subsets of those who may have lingering pharmacologic effects of alpha-blockers vs. those who were never on them. Inflammation is of such paramount importance that a more substantive assessment of the patient's inflammatory status must be routine in any analysis of LUTS. Despite significant technological advances in the biological sciences, the current testing of inflammation remains inadequate and should be addressed in future trials of LSESr given the widely recognized role of inflammation in the development of LUTS [78-85]. The anti-inflammatory properties of Serenoa repens are detailed in many reports [8,85-90]. Lifestyle modifications aimed at reducing inflammation should help modulate LUTS symptoms and possibly prevent progression [65]. Investigators should consider stratifying the patient population using an "inflammation index," and further interpreting clinical data by comparing an "inflammation index" with the BPHII [11].

Simple measures such as restricting fluid intake 4 hours before bedtime, routinely attempting to void prior to sleep, limiting or omitting caffeinated beverages, and avoiding salt in the diet can significantly affect a key symptom such as nocturia [91-94]. Principal investigators should evaluate study participants based on these lifestyle factors to clarify the potential beneficial effects of LSESr relative to the possible lifestyle modifications known to affect LUTS. In an article published 25 years ago [48], Descotes, et al. referred to an article written by Castro still 23 years earlier. In that publication from nearly a half-century ago, Castro remarked on the challenges faced when evaluating patients undergoing treatment for LUTS. "The clinical symptoms of BPH are also labile, and can vary with time, seasons, stress, medication, changes in sympathetic activity, bladder training, sedentary activity, and irregular voiding. Spontaneous variation in disease symptoms and the degree of dynamic obstruction, coupled with a pronounced placebo effect, clearly complicate any assessment of drug efficacy in BPH" [48,95]. Forty-two years after the 1972 publication by Castro, Vaughan shared his views about the clinical lability of LUTS. "To this, I would add my 47 years in frankly discussing LUTS with thousands of patients. Not

only is there variability in nocturia, but also in symptoms of hesitancy, weak stream, incomplete emptying, urgency, and terminal dribbling" [96]. At age 78 years (SBS), and despite being a non-smoker, non-drinker, following a low sodium diet, and not having sleep apnea or obesity as problems, I too echo these observations about the day-to-day variations in LUTS. Such variable symptomatology, combined with issues such as the quality of the Serenoa repens product, and patient compliance with medications, contribute to the difficulty in our understanding of LUTS and its optimal approach to prevention and treatment.

8. Clinical Perspective

A review and network meta-analysis (NMA) of randomized placebo-controlled trials on Serenoa repens vs. placebo vs. alpha-blockers in the treatment of LUTS was reported by Russo, et al [97]. Twenty-two trials were identified by the authors for data investigation using this NMA methodology. The outcomes of IPSS and peak flow were considered across the 22 studies, including 10 randomized trials comparing LSESr to a placebo (five studies), or an alpha-blocker (five studies). For the LSESr studies, two used a HESr product that was compared to tamsulosin, and eight other studies used a non-hexane Serenoa product, with five being placebo-controlled and three studies using a prescription drug as a comparator. From the NMA, Russo concluded that HESr and non-HESr did not demonstrate clinically meaningful improvement in LUTS and peak flow over placebo. De Nunzio, et al [98] published a response to Russo, criticizing the NMA methodology for being inappropriate to ascertain clinical efficacy, and that patient cohorts were unbalanced and affected the validity of the conclusions reached. De Nunzio, et al. also noted that a large number of randomized clinical trials were either not identified or were excluded. They concluded that these shortcomings could lead to false conclusions [98]. Many criticisms similar to those voiced by De Nunzio et al. of the Russo review are to be found in our global review presented herein. This includes failure to retrieve all eligible publications for analysis, including early science that represents the body of literature relevant to modern medicine, and failure to establish strict evaluability criteria for the studies to be reviewed as opposed to relying on methodology for analysis at the expense of clinical relevance. In fact, of the 22 studies cited by Russo, four did not indicate the extraction process, and three of those four were combinations of Serenoa repens with other products, and one study involved only 13 patients. If clinical data are used to present an opinion only, rather than to enhance medical practice, then the data do not have value. The same could be said for the methodology for both reviews and meta-analyses, and the failure to consider non-native language publications. Dated science that is well done builds a foundation for clinical practice and allows patient care to be improved.

The clinical literature on Serenoa repens for the treatment of LUTS is extensive. In some studies, important variables are often inadequately controlled. This has resulted in inconsistent findings and controversy concerning what benefit may result from the use of commercially available Serenoa products. The most important variable repeatedly presented in this multi-part report is whether the product is a lipidosterolic extract of Serenoa repens (LSESr) vs. a crude product such as crushed dried saw palmetto berry powder. In addition to the use of a high-quality standardized LSESr, other factors to consider when evaluating the Serenoa clinical literature are the dosage of LSESr, the criteria used to select patients, the exclusion of products combining Serenoa with other agents, and the clinical study design. Based on the evidence presented, a standardized LSESr, given as monotherapy, and that has an established profile defined by the EMA or USP, at a dosage of 320 mg/d, either in divided doses or as a single daily dose, may contribute to the alleviation of LUTS. The clinically significant endpoints include a decrease in the IPSS score, an improvement in the QoL score, and an increase in peak urinary flow (Qmax), with all parameters achieved in association with a high therapeutic index. LSESr's very favorable safety profile includes a negligible impact on sexual function [43,44,99-105]. Despite achieving the desired endpoints mentioned, the American Urological Association (AUA) and the European Association of Urology (EAU) treatment guidelines have downplayed the efficacy of LSESr therapy [106,107]. In contrast, meta-analyses published in 2016 and 2018 support the use of LSESr in men with mild-to-moderate LUTS/BPH [43,44].

In this clinician's perspective, based upon nearly 40 years of clinical data, some conclusions are clear. First, LSESr has a definite role in the treatment of LUTS. It has a high safety profile with relatively few adverse side effects. It does not cause sexual dysfunction such as ejaculatory disorders seen with 5α -reductase inhibitors, nor hypotension with some α -blockers such as tamsulosin (Flomax®). LSESr does not alter PSA expression and therefore, does not interfere with the monitoring of men at risk of developing prostate cancer. LSESr using hexane, ethanol, or carbon dioxide extraction all have shown efficacy in published studies. The onset of action may be as early as two weeks but is clearly established by 3 months. Of importance is the durability of efficacy seen with long-term treatment of LUTS with LSESr. And of greater significance is the finding of slowing and even halt in the progression of LUTS/BPH during prolonged studies using LSESr, with some trials extending 10 to 15 years. Such studies may indicate that LSESr is affecting the pathologic processes i.e., pathobiology or etiopathogenesis, that lead to LUTS/BPH. Patient selection is important, and those patients with severe LUTS, and at high risk for acute urinary retention are not optimal candidates and warrant careful observation relating to the need for surgical intervention. The most critical issue in the use of Serenoa repens in treating LUTS is the need to educate physicians that crude herbal products are never to be equated with standardized LSESr products that have a profile established by the EMA or USP and that the former products have no role to play in LUTS treatment. An unresolved issue in the use of LSESr relates to the lack of head-to-head studies to ascertain any difference in the hexane vs. ethanol vs. carbon dioxide extracts, but the results presented in this report would indicate that no particular extraction process is superior to another. An additional issue relates to regulatory agencies and their role in monitoring the quality of products such as LSESr. Why is LSESr available by prescription in some countries, OTC in others, and inadequately regulated concerning product quality in others, the latter especially the situation in the United States? We should never confuse the business of medicine with the practice of principled medicine. The former has led to the deterioration of medicine as a profession and has diminished the quality of care to patients while increasing the risk of adverse events.

In summary, LSESr (lipidosterolic extract formulations of Serenoa repens) show efficacy in treating LUTS ± BPH, and the results discussed herein provide a rationale for conducting larger, better-controlled studies using such formulations in men with mild-to-moderate LUTS. These studies should quantitate change in IPSS, QoL, Qmax. An evaluation of the inflammatory status of the patient and the effect of long-term use of LSESr on halting the progression of LUTS/BPH should provide further confirmation that LSESr alters the natural history of this affliction of great "bother" in the adult male.

Declaration of Conflicting Interests: Dr. Strum serves as a paid consultant for Valensa International. He receives a consulting fee from Life Extension.

Funding: Dr. Strum received a consulting fee for researching and compiling this review from U.S. Nutraceuticals, dba Valensa International and funding by an education grant from the Life $Extension^{\otimes}$ group. .

Ethical Approval/Patient Consent: This is a review of the literature and did not require ethical board approval.

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