Natural Hair Supplement: Friend or Foe? Saw Palmetto, a Systematic Review in Alopecia

Evyatar Evron\textsuperscript{a}  Margit Juhash\textsuperscript{b}  Arash Babadjouni\textsuperscript{c}  Natasha Atanaskova Mesinkovska\textsuperscript{b}

\textsuperscript{a}Department of Medicine, Cedars-Sinai Medical Center, Los Angeles, CA, USA; \textsuperscript{b}Department of Dermatology, University of California, Irvine, Irvine, CA, USA; \textsuperscript{c}Arizona College of Osteopathic Medicine, Midwestern University, Glendale, AZ, USA

Keywords
Saw palmetto · Natural · Supplement · Hair loss · Alopecia · Androgenetic alopecia · Telogen effluvium

Abstract
Saw palmetto (SP), a botanical extract with antiandrogenic properties, has gained commercial popularity for its purported benefits on hair regrowth. To summarize published evidence on the efficacy, safety, and tolerability of supplements containing SP in the treatment of alopecia, we conducted a PubMed, Google Scholar, and Cochrane database search using the following terms: (saw palmetto and hair loss), (saw palmetto and androgenetic alopecia), and (saw palmetto and natural supplement and alopecia). Five randomized clinical trials (RCTs) and 2 prospective cohort studies demonstrated positive effects of topical and oral supplements containing SP in the treatment of alopecia, we conducted a PubMed, Google Scholar, and Cochrane database search using the following terms: (saw palmetto and hair loss), (saw palmetto and androgenetic alopecia), and (saw palmetto and natural supplement and alopecia). Five randomized clinical trials (RCTs) and 2 prospective cohort studies demonstrated positive effects of topical and oral supplements containing SP (100–320 mg) among patients with androgenetic alopecia (AGA) and telogen effluvium. Sixty percent improvement in overall hair quality, 27% improvement in total haircount, increased hair density in 83.3% of patients, and stabilized disease progression among 52% were noted with use of various topical and oral SP-containing supplements. SP was well tolerated and not associated with serious adverse events in alopecia patients. Although robust high-quality data are lacking, supplements containing SP may be a treatment option for patients with AGA, telogen effluvium, and self-perceived hair thinning. Further large-scale RCTs focusing on the sole contribution of SP to hair growth outcomes are needed to confirm efficacy and determine long-term adverse events.

Introduction
Saw palmetto (SP) is a botanical extract from the berries of the \textit{Serenoa repens} dwarf tree, native to the subtropical, Southeastern United States \cite{1}. The extract can be prepared as liquid via hot water or supercritical elicitation with carbon dioxide, or as powder via mechanical grinding of raw berries \cite{2}. Due to its antiandrogenic effects, SP has been used as an alternative treatment for benign prostatic hyperplasia (BPH) \cite{3–5}, dating back to the 15th century BCE, Egypt \cite{5}.

SP extract is primarily comprised of fatty acids (70–95\%) \cite{6}, phytosterols such as β-sitosterol (0.1\%) \cite{6}, β-carotene, vitamin E derivatives, and polysaccharides;
however, the exact ratios can vary depending on the specific preparation [7]. SP is a competitive, nonselective inhibitor of both 5α-reductase isoforms, blocking nuclear uptake of dihydrotestosterone (DHT) and decreasing DHT binding capacity to androgen receptors by nearly 50% [8, 9]. SP’s fatty acid components directly inhibit enzyme activity [10], aid in the enzyme’s selective hormone transformation processes [11], and influence access to cofactors by affecting the enzyme’s conformational state [8]. SP also increases the activity of 3α-hydroxysteroid-dehydrogenase, an enzyme converting DHT to its weaker metabolite, androstaneol [7]. However, also worth noting are the discordant views regarding SP’s actual effect on androgen-dependent parameters [12, 13] and debatable clinical efficacy in treating BPH [14, 15].

SP’s antiandrogenic properties, minimal side-effect profile [16], and low drug interaction potential have prompted its use as a complementary alopecia remedy [1, 16, 17]. The extract has been evaluated for the treatment of androgenetic alopecia (AGA) [18–21], telogen effluvium (TE) [22–25], seborrheic dermatitis [26, 27], and facial sebum [28], as a monotherapy or in combination with other supplements, in oral and topical formulations. In this review, we will systematically describe SP extract’s efficacy for the treatment of hair loss conditions and associated side effects.

**Materials and Methods**

A systematic review was completed searching the National Library of Medicine through PubMed, Google Scholar, and Cochran databases, without date limits, in January 2019, using the following terms: (saw palmetto and hair loss); (saw palmetto and androgenetic alopecia); (saw palmetto and natural supplement and alopec-
Results

The above search generated 9 articles using oral and topical supplements containing SP, with a total of 381 patients, for the treatment of alopecia (Fig. 1). Four RCTs [18–21], 2 prospective cohort studies [24, 29], and 1 case report [30] described the effects of topical or oral SP extract for the treatment of AGA. The efficacy of SP-containing products in treating TE was described in 2 case reports [25, 30], 1 RCT [21], and 1 prospective cohort study [24] (Table 1).

Androgenetic Alopecia

The first RCT tested the efficacy of a topical SP-containing lotion twice daily, alone and in combination with an oral tablet containing gelatin-cystine 4 times a day, in 60 men and women with AGA over a period of 50 weeks. Five arms were studied with varying combinations of lotion and oral agent versus vehicle and placebo. Hair samples were degreased, dried, counted on a grid, and weighed in an analytical balance at 22°C. The SP-containing topical demonstrated a time-dependent mean increase in hair count of 17% by week 10 and 27% by week 50, as compared to 6 and 14% among the vehicle group at weeks 10 and 50, respectively (p < 0.005). Hair mass and caliber increased approximately 20% by week 10 from baseline and 30% by week 50 using topical SP (p < 0.005), versus an approximately 10% decrease and an 8% increase amongst the vehicle group at weeks 10 and 50, respectively. The use of topical SP and oral gelatin-cystine supplement caused a further increase of approximately 50% in all hair growth parameters (p < 0.005) when compared to use of either agent alone. Both topical SP and gelatin-cystine supplement were well tolerated, and no adverse events (AEs) were reported [18].

A small RCT evaluated the efficacy of an oral tablet containing 200 mg of SP extract and an extra 50 mg of β-sitosterol (which has also been studied singularly for its antiandrogenic effects) versus placebo, taken twice daily in 26 men with mild to moderate AGA over 25 weeks. The primary endpoints of hair loss arrest and qualitative hair improvement were assessed subjectively by study participants through a standardized 7-point scale questionnaire. SP demonstrated 60% “improvement” in hair loss arrest and overall quality versus 11% among placebo. The active tablet also showed greater conservation of hair density and quality over time, as subjectively assessed by the study participants [19].

The largest RCT to date compared oral 320-mg SP-containing tablets versus 1-mg finasteride daily in 100 men with AGA over 2 years. Measurement of hair density was assessed by standardized global photography; the effectiveness of the treatment was evaluated with a 7-point clinical score scale as determined by 3 expert dermatologists experienced in the field of alopecia. Notably, 68% of patients treated with finasteride had higher hair density scores from baseline as compared to 38% of the SP group (p < 0.05), indicating that SP is inferior to finasteride. Although neither treatment was reported to be clinically effective in 10% of patients, SP had stabilized progression of AGA in 52% of cases [20].

A 24-week, prospective cohort tested the efficacy of a topical SP-containing lotion applied to the entire scalp daily for the treatment of AGA in 50 men. Study participants also applied a concentrated SP-containing serum dedicated to thinning areas of the scalp for the first 4 weeks. Increased hair count was assessed via macrophotographic techniques using the Canfield photography system; hair restoration (defined by average hair size and terminal hair count) along with change in AGA staging was assessed via the Norwood-Hamilton grading system of pictures. Investigator satisfaction and patient satisfaction were measured via photographic and self-perceived assessment questionnaires, respectively, using a 7-point scale. Total hair count increased at 12 and 24 weeks compared to baseline by 3.4 and 4.9%, respectively, while terminal hair count increased by 21.4 and 74.1%, respectively. Medium-sized and vellus hair counts decreased at 24 weeks compared to baseline by ~10.3 and ~25%, respectively. Median AGA stage changed from 4 at baseline to 3 at 24 weeks, with a slight increase of hair over the anterior and vertex scalp, as well as higher patient satisfaction scores noted at study completion [29].

The most recent RCT tested the efficacy of a 100-mg SP-containing food supplement (Lambdapil®, ISDIN, Barcelona, Spain) on 35 men with AGA, for 6 months. Outcomes were measured via phototrichogram, where digital images evaluating hair volume and appearance were assessed by dermatologists through a 7-point clinical score scale. Further outcomes were also subjectively evaluated by study participants via completion of self-as-
**Table 1. Summary of clinical investigations using SP extract for the treatment of AGA, TE, and self-perceived hair loss in male and female patients**

<table>
<thead>
<tr>
<th>Study</th>
<th>Disease</th>
<th>Patient demographics</th>
<th>Quality rating</th>
<th>Study design</th>
<th>Efficacy</th>
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<td>Morganti et al. [18]</td>
<td>Androgenetic alopecia (AGA)</td>
<td>60 M and F</td>
<td>1b</td>
<td>Double-blinded vehicle and placebo-controlled RCT</td>
<td>Significant improvement in hair count at 10 wk (17%) and 50 wk (27%) using SP-containing lotion versus vehicle (~6 and 14%, respectively)</td>
<td>No AEs reported</td>
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<td>5 tx arms: SP-containing topical lotion BID and/or an oral supplement containing gelatin-cystine QID versus vehicle and/or placebo for 50 wk</td>
<td>Hair mass and caliber significantly improved by 20% at 10 wk and 30% at 50 wk using SP-containing lotion versus vehicle (10% decrease to 8% increase, respectively)</td>
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<td>Prager et al. [19]</td>
<td>Mild to moderate AGA</td>
<td>26 M</td>
<td>1b</td>
<td>Double-blinded placebo-controlled RCT</td>
<td>60% improvement from baseline in hair loss arrest and overall quality for the SP group versus placebo (11%)</td>
<td>Nausea, constipation, diarrhea unlikely related to tx</td>
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<td>Oral soft-gel supplement (50-mg β-sitosterol, 200-mg SP) versus placebo BID for up to 25 wk</td>
<td>Patient-reported conservation of hair density and quality over time for the SP group versus placebo</td>
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<td>Rossi et al. [20]</td>
<td>AGA</td>
<td>100 M</td>
<td>1b</td>
<td>Head-to-head double-blinded RCT</td>
<td>Significantly higher hair density scores (determined as &gt; 0 indicating increased hair density from baseline) for finasteride (68%) versus SP (38%)</td>
<td>No AEs reported</td>
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<td>Oral supplement containing 320-mg SP versus 1-mg finasteride daily for 24 mo</td>
<td>SP stabilized AGA disease progression in 52%</td>
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<td>Neither tx effective in 10%</td>
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<td>Wessagowit et al. [29]</td>
<td>AGA</td>
<td>50 M</td>
<td>4</td>
<td>Prospective cohort</td>
<td>Increased total hair count at 12 and 24 wk from baseline (3.4 and 4.9%, respectively)</td>
<td>Cold sensation (16%)</td>
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<td>Topical SP-containing lotion to entire scalp daily for 24 wk; concentrated SP-containing serum to thinning areas daily for first 4 wk</td>
<td>Increased terminal hair count at 12 and 24 wk (21.4 and 74.1%, respectively)</td>
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<td>Decreased medium-sized and vellus counts at 24 wk from baseline (~10.3 and ~25%, respectively)</td>
<td>Unpleasant smell (2%)</td>
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<td>Median AGA stage change from 4 to 3 at 24 wk, with mildly increased hair over anterior and vertex scalp</td>
<td>Itchy scalp (2%)</td>
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<td>Higher patient satisfaction scores at 24 wk compared to baseline</td>
<td>Acne on the forehead (2%)</td>
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<tr>
<td>Narda et al. [21]</td>
<td>AGA</td>
<td>35 M</td>
<td>1b</td>
<td>Double-blinded placebo-controlled RCT</td>
<td>23.4% increase in the anagen/telogen ratio noted from baseline</td>
<td>No AEs reported</td>
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<td>Lambdapil® capsules (containing 100-mg SP) versus placebo for 6 mo</td>
<td>3.7% increase in total anagen hair percentage (vs. 0.8% decrease), and 3.7% decrease in total telogen hair percentage (vs. 0.8% increase) for SP groups versus placebo at 6 mo</td>
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<tr>
<td>Zanzottera et al. [24]</td>
<td>AGA</td>
<td>15 M</td>
<td>4</td>
<td>Prospective, noncomparative, open label pilot</td>
<td>Increased hair density at 6 mo in 83.3% of subjects 26.7% of men with “greatly increased” hair density from baseline</td>
<td>No AEs reported</td>
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<td>Nutritional supplement containing 300-mg SP, 2 tab daily for 6 mo</td>
<td>93.3% “general reduction” in hair loss reported by subjects, with 79.0% reporting “significantly high” reduction</td>
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assessment surveys regarding perceived treatment efficacy. There was a 23.4% increase from the baseline anagen/telogen ratio. An additional 3.7% increase in total anagen hair percentage (vs. 0.8% decrease), and a 3.7% decrease in total telogen hair percentage (vs. 0.8% increase), was noted among the active SP cohort as compared to placebo at 6 months. Patients reported increased hair volume, improved quality of life (QOL), and enhanced self-perceived efficacy using the SP-containing tablet [21]. A prospective, noncomparative cohort tested the efficacy of a nutritional compound containing 300-mg SP taken twice daily for 6 months in 15 males with AGA and 15 females with either female-pattern hair loss (FPHL), severe TE, or stress and food-deficiency induced hair loss.

Evaluation of hair loss and hair health parameters was performed via trichoscopy. Global photographs assessing hair density were evaluated by experts on a 7-point scale, and subjective participant efficacy was measured via standardized questionnaires. An increase in hair density at 6 months was noted among 83.3% of study participants,

<table>
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<th>Study</th>
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<th>Efficacy</th>
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<tr>
<td>Farris et al. [30]</td>
<td>Unspecified hair loss pattern (n = 3)</td>
<td>3 F</td>
<td>5</td>
<td>Case report</td>
<td>Subjective improvement in hair growth and temple area coverage; also decreased shedding</td>
<td>No AEs reported</td>
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<td>Pezza et al. [25]</td>
<td>Severe TE</td>
<td>1 M</td>
<td>5</td>
<td>Case report</td>
<td>Significantly improved hair growth and density (negative pull test)</td>
<td>No AEs reported</td>
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<td>Narda et al. [21]</td>
<td>Acute TE</td>
<td>35 F</td>
<td>1b</td>
<td>Double-blinded placebo-controlled RCT</td>
<td>Improved hair pull test at 1, 3, and 6 mo (9.8, 7.5, and 6.5%, respectively) for SP versus placebo (11.9, 10.7, and 8.8%, respectively)</td>
<td>Bloating (n = 1)</td>
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<td>Zanzottera et al. [24]</td>
<td>FPHL</td>
<td>15 F</td>
<td>4</td>
<td>Please see the text for discussion of study design and treatment</td>
<td>33.3% of women with “greatly increased” hair density from baseline</td>
<td>Please see the text for discussion of other results</td>
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<td>Ablon et al. [31]</td>
<td>Self-perceived hair thinning</td>
<td>40 F</td>
<td>1b</td>
<td>Double-blinded placebo-controlled RCT</td>
<td>Significant increase in terminal, vellus, and total hair counts at 3 mo for the supplement group (6.8, 10.1, and 7.1%, respectively) versus placebo (0.07, −2.9, and 0.4%, respectively)</td>
<td>No AEs reported</td>
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<td>At 6 mo, significant increase in terminal, vellus, and total hair counts for the supplement group (10.4, 15.7, and 10.8%, respectively) versus placebo (3.5, −2.2, and 0.3%, respectively)</td>
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<td>Higher GHA scores for hair growth (1.08 vs. 0.08) and hair quality (1.12 vs. 0.08) for the supplement group versus placebo, respectively</td>
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<td>Nutrafol® Women’s capsules versus placebo daily for 6 mo</td>
<td>Decreased anxiety and improved wellness for the supplement group versus placebo</td>
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</table>
with “greatly increased” density described by 26.7% of men and 33.3% of women. 93.3% of the subjects reported “general reduction” in hair loss, with 79.0% defining it as “significantly high,” 83.3% reported thicker and bulkier hair, 90.0% felt their hair was stronger with a better combing effect, and 88.5% reported reduced greasiness. At the end of the study, 93.0% of patients were “satisfied” with the results. The supplement was well tolerated [24].

A case series described 3 women with various hair loss patterns and 1 man with early onset AGA, who were successfully treated with Nutrafol® oral supplements (Nutraceutical Wellness Inc., New York, NY, USA) which include an undisclosed amount of SP extract. Subjective outcomes included enhanced hair growth, improvement in temple area coverage, decreased shedding, and self-reported satisfaction with treatment outcomes, as noted amongst all study participants [30].

**TE and Self-Perceived Hair Thinning**

The first case report of a patient with severe TE treated with an oral supplement containing SP, amino acids, vitamin E, and iron taken daily for 6 months demonstrated significant improvement in hair growth and density, as evidenced by a negative pull test and increased anagen hairs on trichoscopy [25]. In another study discussed above, besides the men with AGA, 35 women with acute TE were also treated with Lambdapil® capsules for 6 months. Outcomes measured via the hair pull test resulted in significantly less hair loss at 1, 3, and 6 months compared to placebo (9.8, 7.5, and 6.5% vs. 11.9, 10.7, and 8.8%, respectively; p < 0.05). The female patients also reported increased hair volume, improved QOL, and enhanced self-perceived efficacy with use of the SP-containing supplement. One patient experienced abdominal bloating, resulting in premature termination from the study [21]. In the study by Zanzottera et al. [24], 15 women with FPHL, TE, or stress-induced hair loss were treated with a nutritional supplement containing 300 mg of SP for 6 months with positive results using the abovementioned outcome evaluation techniques.

A recent RCT evaluated the use of an oral Nutrafol® Women’s capsule taken 4 times daily for 6 months, to strengthen and promote hair growth in 40 healthy adult women with self-perceived thinning. Increments in total hair counts, blinded investigator Global Hair Assessment (GHA) scales, changes in terminal hair diameter, and bundle measurements were all assessed via phototrichograms obtained though macrophotography analysis. A significant increase in terminal, vellus, and total hair counts was noted at 3 months in the supplement group (6.8, 10.1, and 7.1%, respectively) compared to placebo (0.07, −2.9, and 0.4%, respectively). Similar outcomes among the treatment group were also noted at 6 months (10.4, 15.7, and 10.8%, respectively). Higher GHA scores for hair growth (1.08 vs. 0.08) and hair quality (1.12 vs. 0.08), along with decreased anxiety and improved wellness parameters, were noted among those receiving SP-containing supplements [31].

**Adverse Events**

The majority of reported AEs with use of oral SP for alopecia are mild and most commonly gastrointestinal in nature, including nausea, constipation, and diarrhea [2, 16, 19, 26]. Additionally, likely associated with its hormonal effects, SP has been reported to cause vasomotor symptoms in premenopausal females and was suspected as a cause of early menarche in young girls [22, 32]. A report by Miroddi et al. [32] described hot flashes in an 11-year-old girl that appeared after 2 months of treating TE with a food supplement containing SP. The hot flashes ceased when the product was discontinued; however, 45 days after cessation the girl experienced menarche. Another report by Morabito et al. [22] also described similar vasomotor symptoms in a 10-year-old girl after 3 months of treating hirsutism with a food supplement containing 320 mg of SP extract. As in the prior report, here too the symptoms ceased upon discontinuation of the supplement and the girl experienced menarche shortly thereafter. Topical agents containing SP extract have also been associated with minor AEs including cold sensation, mild burning, unpleasant smell, itchy scalp, acne on the forehead, and allergic contact dermatitis [33, 34]. Although we are currently unaware of any recommendations against use of SP-containing supplements for hair loss among adult men or women with child bearing potential, as with any unregulated substance, patients are advised to seek specific recommendations from their primary care provider and/or dermatologist prior to initiation of use.

**Discussion**

Hair loss is associated with significant consequences affecting body image, self-esteem, emotional well-being, and QOL [35], possibly even leading to psychiatric morbidity such as anxiety and depression [36, 37]. While minoxidil and finasteride remain the only United States Food and Drug Administration (FDA)-approved and most widely used therapies for pattern hair loss [38], un-
Saw Palmetto, a Systematic Review in Alopecia

Saw Palmetto, a key ingredient in many over-the-counter supplements marketed for hair regrowth; however, limited data exist to support its efficacy in various alopecias or to better delineate its side-effect profile.

The vast majority of data pertaining to SP’s efficacy, antiandrogenic properties in vitro, and side-effect profile are derived from studies on BPH [8, 9] and treatment of urinary symptoms in vivo [3–5]. However, many of these findings have been called into question, given subsequent trials showing no effects on androgen-dependent parameters when SP is compared to traditional antiandrogenic agents such as finasteride [12, 13]. Despite numerous reports regarding SP’s clinical efficacy, comprehensive meta-analyses have failed to demonstrate significant improvement in BPH symptoms or objective disease parameters, thus casting doubt on SP’s previously attributed antiandrogenic properties [14, 15].

Clinical evidence regarding the efficacy of SP-containing products to treat hair loss is limited. SP has shown promising results in murine models, with hair regrowth mediated through transforming growth factor-β and mitochondrial signaling pathways [40]. Additionally, in vitro models have demonstrated SP’s ability to inhibit inflammatory gene expression in human keratinocytes, suggesting a multifaceted mechanism for the treatment of AGA, in addition to its proposed antiandrogenic properties [41]. The mechanism in which SP instigates hair growth in TE is not entirely clear. A possible explanation may be secondary to its anti-inflammatory properties, or to β-sitosterol’s purported angiogenic effects, stimulating vascular endothelial growth factor in vitro and promoting neovascularization in vivo [42].

Several human studies have demonstrated modest hair regrowth using oral and topical SP-containing products among patients with AGA, FPHL, TE, and self-perceived hair thinning. Improvement in total hair count ranging from 3.4 to 27% [18, 29], increased hair density in up to 83.3% of patients [24], 60% improvement in overall hair quality [19], and stabilized disease progression among 52% of patients [20] were noted with use of various SP-containing oral supplements, topical lotions, and/or serums.

However, almost all current formulations containing SP also contain other vitamins, minerals, or chemical additives, making it challenging to discern the exact extent of SP’s contribution to the observed clinical findings. Additionally, among the RCTs presented there were several methodological flaws, such as inadequate statistical analyses [18], limited sample size [19], lack of appropriate comparison groups [24, 29], qualitative rather than quantitative measures of hair regrowth [30], and conflicts of interest with sponsorship by the pharmaceutical industry [21]. In the only head-to-head RCT comparing SP-containing tablets to systemic finasteride, moderate efficacy in hair regrowth was demonstrated among the SP group; however, overall outcomes were inferior to finasteride [20]. Clinical trials involving patients with TE were only conducted using female patients, and results were described qualitatively. Currently, there is a trial (NCT03052413) assessing the efficacy of an SP-containing supplement in women with mild to moderate hair loss; however, results are pending [43].

SP is well tolerated, and although it has not been associated with serious AEs in alopecia patients [16], more grave effects in other clinical settings have been described. Due to its hormonal effects, SP has the potential to cause gynecomastia [16], decrease libido [16], and reduce prostate serum antigen (PSA) levels, thereby necessitating vigilance not to miss early prostate cancer diagnosis [26]. Other reported AEs include rhinitis [16], intraoperative hemorrhage [44], hepatitis and pancreatitis [45, 46], loss of iris tone and intraoperative floppy-iris syndrome (IFIS) [47], and even 1 case of hypokalemia leading to cardiac arrest and death [16]. Further long-term studies are required for reliable safety assessment.

Limitations of this review include lack of meta-analysis, which was not performed owing to absent standardization of results between studies, and a low number of high-quality RCTs. Additionally, the exclusive impact of SP could not be adequately assessed, as many supplements were comprised of multiple, active ingredients, and several studies failed to adequately specify the precise amount of SP or the exact formula content used. The absence of standardized measurement tools used to assess hair growth in each study is also a considerable limitation given the significant expected variability in outcomes directly correlating with the preferred method of evaluation.

Conclusion

Although robust efficacy data are lacking, SP extract in either topical or oral formulations may have a role in the treatment of hair loss disorders such as AGA or TE, demonstrating modest improvement in hair regrowth. However, as many of the current trials are lacking in quality (descriptive, qualitative measures, small sample sizes, inadequate product content description, long-term results,
Conflict of Interest Statement
The authors have no conflicts of interest to declare.

References

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Author Contributions
This is the original work of Dr. Evyatar Evron, who was involved in reviewing the literature, subsequent analysis of the collected data, and writing of the final report. Dr. Natasha Atanaskova Mesinkovska, Dr. Margit Juhasz, and Mr. Arash Babadjouni were involved in the development of the initial study design and protocol as well as in the editing of this paper.


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