**Tribulus Terrestris for Female Sexual Dysfunction: A Systematic Review**

**Tribulus terrestris para disfunção sexual feminina: Uma Revisão Sistemática**

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**Abstract**

**Objective** We performed a systematic review to assess the effectiveness and safety of *Tribulus terrestris* to treat female sexual dysfunction (FSD).

**Data sources** We performed unrestricted electronic searches in the MEDLINE, CENTRAL, EMBASE, LILACS, CINAHL, PsycINFO, WHO-ICTR, Clinicaltrials.gov and OpenGrey databases.

**Selection of studies** We included any randomized controlled trials (RCTs) that compared *T. terrestris* versus inactive/active interventions. After the selection process, conducted by two reviewers, 5 RCTs (n = 279 participants) were included.

**Data collection** Data extraction was performed by two reviewers with a preestablished data collection formulary.

**Data synthesis** Due to lack of data and clinical heterogeneity, we could not perform meta-analyses. The risk of bias was assessed by the Cochrane Risk of Bias (RoB) tool, and the certainty of evidence was assessed with Grading of Recommendations, Assessment, Development and Evaluations (GRADE).

**Results** After 1 to 3 months of treatment, premenopausal and postmenopausal women randomized to *T. terrestris* had a significant increase in sexual function scores. Three months of treatment with *T. terrestris* showed a significant increase in the serum testosterone levels of premenopausal women. There was no report of serious adverse events, and none of the studies assessed health-related quality of life. The certainty of the evidence was very low, which means that we have very little confidence in the effect estimates, and future studies are likely to change these estimates.

**Conclusion** More RCTs are needed to support or refute the use of *T. terrestris*. The decision to use this intervention should be shared with the patients, and the uncertainties around its effects should be discussed in the clinical decision-making process.

Number of Protocol registration in PROSPERO database: CRD42019121130

**Keywords**
- tribulus
- sexual dysfunction
- review
- evidence-based medicine

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Introduction

Female sexual dysfunction (FSD) is a common condition associated with physical, psychological, and sociocultural factors.¹⁻³ The International Society for the Study of Women’s Sexual Health (ISSWSH) classifies FSD in four categories: hypoactive sexual desire disorder (HSDD), sexual arousal disorders (genital and cognitive), orgasmic disorders, and sexual pain disorders.⁴ In a large epidemiological study conducted over 10 years ago, 12% of 31,581 American women reported a distressing sexual problem, and the percentages were higher among older (45–64 years) participants.⁵ Up to 30 to 50% of women will have FSD during their lifetime, and this rate is probably underestimated due to the social aspects associated with this condition.³⁻⁴ An estimated 20% of women in all age groups have orgasmic disorders, and 10 to 16% have HSDD, while up to 15% of premenopausal and 30% of postmenopausal women have arousal difficulties.²⁻⁶,⁷

Standard care for FDS usually involves a multidisciplinary approach, including hormonal therapy, psychotherapy and pharmacological therapy, to address all components of the disorder.³ Medicinal plants have been increasingly used by women with FDS, often without a medical prescription.² Tribulus terrestris L. (Zygophyllaceae) is a creeping herb, originally from India, which is used as a natural sexual stimulant. Tribulus extracts contain protodioscin, a steroidal sapinon that can influence hormonal activity and affect the production of endogenous androgen by increasing the release of luteinizing hormone.²,⁸,⁹ However, the effects of this intervention have not been established. Therefore, the objective of this systematic review was to evaluate the effectiveness and safety of T. terrestris for the treatment of FDS (►Table 1).

Methods

Study Design

We registered the protocol of the present study with the International Prospective Register of Systematic Reviews (PROSPERO) (CRD42019121130). The current systematic review of the literature followed the methodological recommendations of the Cochrane Handbook for Systematic Reviews of Interventions¹⁵ and the reporting recommendations of the Preferred Reporting Items for Systematic Reviews and Meta-Analyses (PRISMA) statement.¹⁶

Inclusion Criteria

Types of Studies

We included only randomized clinical trials (RCTs).
Table 1 Main characteristics of the included studies

<table>
<thead>
<tr>
<th>Study (year)</th>
<th>Participants</th>
<th>Interventions and comparators</th>
<th>Outcomes</th>
<th>Follow-up</th>
<th>Funding</th>
</tr>
</thead>
<tbody>
<tr>
<td>Vale et al (2018)¹⁰</td>
<td>N = 40 premenopausal women with HSDD Age 18 to 44 years</td>
<td>G1: <em>T. terrestris</em> (N = 20)* 250 mg orally 3 times/day, 120 days  G2: Placebo (N = 20)*</td>
<td>Sexual Function (FSFI and SQ-F); Serum testosterone level</td>
<td>Immediately after treatment (4 months)</td>
<td>No financial support</td>
</tr>
<tr>
<td>Souza et al (2016)¹¹</td>
<td>N = 46 postmenopausal women with HSDD Age 43 to 65 years</td>
<td>G1: <em>T. terrestris</em> (N = 20)* 250 mg orally 3 times/day, 120 days  G2: Placebo (N = 16)*</td>
<td>Sexual Function (FSFI); Serum testosterone level</td>
<td>Immediately after treatment (4 months)</td>
<td>A pharmacy provided <em>T. terrestris</em> used in the study</td>
</tr>
<tr>
<td>Postigo et al (2016)¹²</td>
<td>N = 60 postmenopausal women with HSDD Age: G1 54 ± 5.1 years G2 56 ± 5.8 years</td>
<td>G1: <em>T. terrestris</em> (N = 30)* 250 mg orally 3 times/day, 90 days  G2: Placebo (N = 30)*</td>
<td>Sexual Function (SQ-F)</td>
<td>Immediately after treatment (3 months)</td>
<td>The study received funding from a governmental fund and the main investigator had a research fellowship.</td>
</tr>
<tr>
<td>Guazzelli et al (2014)¹³</td>
<td>N = 66 postmenopausal women with HSDD Age: G1 56 ± 5.8 years G2 53 ± 3.9 years G3 54 ± 5.1 years</td>
<td>G1: <em>T. terrestris</em> (N = 22)* 250 mg orally 3 times/day, 90 days  G2: Tibolone (N = 24)* 1.25 mg/oral administration/day, 90 days  G3: Placebo (N = 20)</td>
<td>Sexual Function (SQ-F)</td>
<td>Immediately after treatment (3 months)</td>
<td>The study received funding from a governmental fund</td>
</tr>
<tr>
<td>Akhtari et al (2014)¹⁴</td>
<td>N = 67 premenopausal women with HSDD Age G1 36 ± 6.2 years G2 36.1 ± 5.8 years</td>
<td>G1: <em>T. terrestris</em> (N = 30)* 7.5 ml syrup, 2 times/day, 30 days (3.5 g of ethanolic extract per 5 ml of syrup)  G2: Placebo (N = 30)*</td>
<td>Sexual Function (FSFI)/p&gt;Adverse events</td>
<td>Immediately after treatment (1 month)</td>
<td>The study was supported by Tehran University of Medical Sciences; it is not clear if this was financial support</td>
</tr>
</tbody>
</table>

Abbreviations: FSFI, Female Sexual Function Index; g, grams; HSDD, hypoactive sexual desire disorder; mg: milligrams; ml, milliliters; N, number of participants; SQ-F, Sexual Quotient Female Questionnaire; T. terrestris, Tribulus terrestris.

¹Number of patients included in the analysis.

Types of Participants
We included trials that recruited women (aged 16 or over) with a clinical diagnosis of any type of female sexual dysfunction.

Types of Interventions
All RCTs that tested *T. terrestris* in any dose, regimen, route of delivery, and for any duration, were eligible for inclusion in the review. The studies had to compare this intervention versus placebo, no intervention, or any active treatment. Trials that administered *T. terrestris* combined with another intervention were eligible if the effects of *T. terrestris* could be isolated.

Outcomes
Primary Outcomes:

a) Sexual function assessed by validated tools, such as the Female Sexual Function Index (FSFI)¹⁷ and Sexual Quotient Female Version (SQ-F).¹⁸

b) Health-related quality of life assessed by any general or specific validated tool.

c) Serious adverse events defined as the proportion of patients who had at least one life-threatening adverse event that resulted in hospitalization, disability or incapacity. Secondary outcomes:

d) Serum testosterone levels measured by any laboratory exam.

e) Minor adverse events defined as the proportion of participants presenting at least one minor adverse event.

We considered all time-points reported in the RCTs. We intended to pool (in metanalyses) only similar time points: short term (up to 3 months), middle term (between 3 and 6 months) and long term (more than 6 months).

Search for Studies
We created a broad and sensitive search strategy, without language, date, or publication status restrictions, to identify all potentially relevant studies.

Electronic Search
We ran the search in the following electronic databases to identify studies published from inception to February 11, 2019: MEDLINE (via Pubmed), Cochrane Central Register of Controlled Trials - CENTRAL (via Wiley), EMBASE (via Elsevier), Literatura Latino Americana em Ciências da Saúde e do Caribe - LILACS (via Biblioteca Virtual em Saúde [BVS]), Cumulative Index to Nursing and Allied Health Literature – CINAHL (EBSCO host), and PsycINFO (via American Psychological Association). See complete search strategies and all terms used in the searches in – Supplementary Table S1.
Search for Ongoing Studies
We searched for ongoing studies in the World Health Organization (WHO) International Clinical Trials Registry Platform (apps.who.int/trialsearch) and in ClinicalTrials.gov (www.clinicaltrials.gov).

Hand Search and Search for Unpublished Studies
We searched for unpublished studies in Open Gray (http://www.opengrey.eu/). We contacted experts in the field to inquire about any additional ongoing or unpublished studies. We also screened the reference lists of all included studies to identify additional potentially relevant trials.

Process of Study Selection
We used the Rayyan software in the two phases of the study selection process. In the first phase, two authors (RLP and COCL) independently screened the titles and abstracts of all records retrieved through the search strategy. In the second phase, the same two authors independently read the full texts of the records coded as ‘potentially relevant’ and included those that fulfilled the aforementioned selection criteria. We created a table with reasons for exclusion of the studies in this phase of the selection process. When needed, a third review author (RR) solved disagreements.

Data Extraction
We used a data extraction form especially created for this review to collect relevant information from each included trial. Two independent review authors (RLP and ALCM) extracted data; a third author (RR) solved any disagreements.

Assessment of the Risk of Bias
We used the Cochrane risk of bias tool to assess the methodological quality of the included trials. This tool assesses seven domains of each RCT: random sequence generation, allocation concealment, blinding of participants and personnel, blinding of outcome assessors, incomplete outcome data, selective reporting of outcomes, and other potential sources of bias. Two authors (RLP and DVP) performed these assessments independently; a third author (RR) solved disagreements.

Heterogeneity Between Included Studies
We planned to assess the heterogeneity of the intervention effects by visual inspection of the forest plots. We planned to use the chi-squared test ($p > 0.1$) as indicative of statistical heterogeneity (inconsistency), and the I-squared test to measure the extent of heterogeneity ($I^2 > 50$ being indicative of significant heterogeneity). We also planned to examine the reasons for heterogeneity by conducting additional analyses. This was not possible due to lack of data.

Measures of Treatment Effect and Analysis
For dichotomous outcomes, we report results using risk ratios (RRs); for continuous outcomes, we used mean differences (MDs). We calculated the 95% confidence intervals (CI) for all reported outcomes. Where possible (availability and homogeneity of data), we planned to pool treatment effects of individual trials into metanalyses using a random effects model and the Review Manager 5.3 software (The Nordic Cochrane Centre/The Cochrane Collaboration, Copenhagen, Denmark). This was not possible.

Subgroup and Sensitivity Analyses
We planned to perform subgroup analyses for all primary outcomes comparing pre and postmenopausal women. We also planned to perform two sensitivity analyses for all primary outcomes: random-effects versus fixed-effect metanalyses, and trials with low versus high or unclear risk of selection, detection, performance, and attrition bias. However, due to lack of data, we could not perform these analyses.

Publication Bias Assessment
We planned to investigate publication bias using funnel plots in metanalyses with more than 10 studies. This was not possible due to lack of data.

Assessing the Certainty of the Evidence
We used the Grading of Recommendations, Assessment, Development and Evaluations (GRADE) approach to evaluate the certainty of the body of evidence for the comparison between T. terrestris versus placebo for the primary outcomes. We assessed the certainty of the evidence in all five GRADE domains (risk of bias, inconsistency, imprecision, indirectness, and publication bias). We report reasons to downgrade or upgrade the evidence. We present a summary of findings table using the software GRADEpro GDT (McMaster University, Hamilton, ON, Canada).

Results
Search Results
The search strategy retrieved 1,258 references. After the exclusion of 16 duplicates, we screened 1,242 unique references, excluded 1,236, and selected 6 for full text reading. One was an ongoing trial (IRCT201612131340N1) that may contribute data in future updates of this review (Supplementary Table S2). Thus, 5 RCTs were included in the review (Fig. 1). The 5-parallel design RCTs were published between 2014 and 2017 in Brazil ($N = 4$) and Iran ($N = 1$) and enrolled a total of 279 women with HSDD or loss of libido that caused distress. Three studies included only postmenopausal women ($N = 172$; age range 43–65 years), and 2 studies included only premenopausal women ($N = 107$; 18–44 years). Most trials excluded women with any psychiatric condition, smokers, with a history of breast or endometrial cancers, or with diabetes mellitus, cardiovascular or renal disease, and/or using any drugs that could interfere with sexual desire, including hormone therapy. All five RCTs compared T. terrestris versus placebo. One study had three groups: T. terrestris, tibolone, and placebo. Four trials administered the drug orally (250 mg 3 times daily for 90–120 days) and 1 gave the participants a syrup

Description of Studies
The five RCTs compared T. terrestris versus placebo. One study had three groups: T. terrestris, tibolone, and placebo. Four trials administered the drug orally (250 mg 3 times daily for 90–120 days) and 1 gave the participants a syrup
containing T. terrestris extract (twice daily for 30 days). All five studies assessed sexual function as one of their outcomes; two studies also assessed testosterone levels. Only one study reported adverse events.

**Risk of Bias of Included Studies**

We classified all trials as having an unclear risk for selection bias (random sequence generation and allocation concealment) because the authors did not provide sufficient information for judgement (► Fig. 2). All studies had a low risk of bias for blinding of participants and personnel. Three studies had a low risk of bias for blinding of outcome assessors; the other two had an unclear risk of bias for this domain. We classified two studies as having a high risk for attrition bias because of the large number of losses (20% and 37.5%). The five studies reported all the outcomes planned in their registered trial protocols; we, therefore, classified them as having a low risk for reporting bias. Two studies had an unclear risk for other biases because they did not report the baseline characteristics of the study participants.

**Effects of Interventions**

► Table 2 presents a summary of the results of the five trials. The results of the individual studies could not be combined in metanalyses due to clinical differences in the participants (premenopausal and postmenopausal women) and lack of data (mean and/or standard deviation). We contacted the authors of the studies to obtain additional data but only one replied.

**Sexual Function Assessment**

Three studies assessed sexual function using the Female Sexual Function Index (FSFI) (scores range from 2–36, higher values indicate better function), immediately after 1 to 4 months of treatment in 153 participants (46 post and 107 premenopausal women). Three studies used the Sexual Quotient Female Questionnaire (SQ-F) (scores range from 0–100, with higher values indicating better function), to assess sexual function after 3 to 4 months of treatment in 40 premenopausal and 126 postmenopausal women.

One of the three studies that used the FSFI assessed only premenopausal women and found significantly higher mean total scores in the T. terrestris group after 1 month of treatment (mean deviation [MD] 4.39; 95% confidence interval [CI] 2.90–5.88 points; 67 participants; very low certainty evidence). The authors also reported significant increases in arousal, lubrication, orgasm, and satisfaction scores, but not in desire and pain scores (► Fig. 3). The other two studies (46 post...
and 40 premenopausal women) that used the FSFI found non-significant differences in the mean overall scores of the *T. terrestris* and placebo groups after 3 months of treatment (p = 0.19 and p = 0.44, respectively). These two studies did not provide the standard deviations for these scores (► Table 2). One study12 (60 postmenopausal women) found a significant increase in SQ-F scores in the *T. terrestris* group after 3 months of treatment (MD 16.40; 95% CI 7.67–25.13; 60 participants; very low certainty evidence). The other studies10,13 (66 postmenopausal and 40 premenopausal women, respectively) reported only the results before and after treatment for each group and did not calculate the differences between them. One of those studies10 reported the presence of sexual dysfunction related to each domain of the SQ-F, and there was a significant improvement in all domains after treatment with *T. terrestris* (p = 0.001), but not after placebo (p = 0.07). The other study13 that assessed sexual function using the SQ-F score only reported the final mean scores in each of the three group and did not calculate the differences between them (mean final SF-Q scores: 56 points in the placebo group, 69 in the *T. terrestris* group and 84 in the tibolone group).

**Adverse Events**

Only Akhtari et al (2014)14 assessed adverse events. None of the 60 participants had serious adverse events; one participant had abdominal cramps, but the authors did not specify to which group she belonged (60 premenopausal women; very low certainty of evidence).

**Serum Testosterone Levels**

Two studies10,11 (96 participants) measured total serum testosterone levels after 3 months of treatment. One study11 involved only postmenopausal women and reported non-significant differences between the *T. terrestris* and placebo groups (MD 2.50; 95% CI -1.79–6.79; 46 participants). The other study10 involved only premenopausal women and reported a significant increase in testosterone levels in the *T. terrestris* group (MD 6.60; 95% CI 0.27–12.93; 40 participants).

![Fig. 2 Risk of bias summary.](image)

![Fig. 3 Forest plot of *Tribulus terrestris* versus placebo (only in premenopausal women). Outcome: Sexual function measured by Female Sexual Function Index (FSFI).](image)
None of the included studies assessed health-related quality of life.

**Certainty of the Evidence Assessment**
We assessed the certainty of the evidence for the primary outcomes of the main comparison (*T. terrestris* versus placebo). The certainty of the evidence is very low for sexual dysfunction and adverse events, after 1 and 3 months of treatment. The reasons to downrade the evidence were the risk of bias of the trials, and imprecision due to small sample sizes. We provide explanations for each judgment in the summary of findings table (*► Supplementary Table S3*).

**Discussion**
The present systematic review evaluated the effectiveness and safety of *T. terrestris* in the treatment of women with sexual dysfunction. We identified five RCTs that could not be pooled into metaanalyses due to lack of data and differences in study participants. We downgraded the certainty of the evidence to very low due to methodological limitations of the trials and imprecision attributed to small sample sizes. We had concerns about possible selection bias in all trials because the authors provided little information about the methods used for random sequence generation and allocation concealment. Based on these considerations, we advise caution when interpreting the results of this review.
on the findings of single studies, *T. terrestris*, when compared with placebo, showed an improvement in sexual function scores (FSFI and SQ-F) in premenopausal and postmenopausal women, after 1 to 3 months of treatment. Regarding serum testosterone levels, 3 months of treatment with *T. terrestris* showed a statistically significant increase in premenopausal women, but this effect was not seen in postmenopausal women. Only one study assessed adverse events and reported that one participant had abdominal cramps but did not specify to which group she belonged. Only one study compared *T. terrestris* versus another active intervention (tibolone), but the authors did not provide quantitative data to assess differences between these interventions.

The results of the current review should be interpreted with caution because of the very low certainty of the evidence. This means that we are very unsure about the effect estimates, and future studies are likely to change the magnitude and direction of these estimates. We cannot compare our findings to those of other reviews because, to the best of our knowledge, this is the first systematic review about this intervention.

Our study had several strengths starting with its strict adherence to the methodological recommendations of the Cochrane Handbook and the PRISMA reporting guidelines. We also conducted a broad and sensitive literature search, including gray literature and hand search, to try to identify all potentially relevant studies. The main limitation of the review was the lack of success in obtaining additional information from trial authors. These details would have been important to assess the risk of selection bias of all trials, and additional quantitative data could have allowed us to perform meta-analyses.

The findings of our review should alert clinicians and patients that there is very low certainty evidence regarding the effects (benefits and harms) of *T. terrestris* for the treatment of women with sexual disorders. Current evidence does not support the routine use of *T. terrestris* in clinical practice.

There is a need for additional, well designed, and well conducted RCTs to assess the effects of this intervention for FSD in pre and postmenopausal women. The authors of these trials should adhere to the Consolidated Standards of Reporting Trials (CONSORT) reporting guidelines. This will help to reduce the uncertainty of effect estimates and allow more robust conclusions.

**Conclusion**

The present systematic review found very low-certainty evidence, from small single studies, that *T. terrestris* increases sexual function scores (FSFI and SQ-F) in premenopausal and postmenopausal women. However, these results should be interpreted with caution since future studies are likely to change the magnitude and direction of our estimates. More trials are needed to support or refute the use of *T. terrestris* in clinical practice.

**Contributions**

Drafting the study protocol (all authors); development of the search strategy (COCL and RR); selection of studies and data extraction (RLP and ALCM and RR); assessment of risk of bias of the included studies (RLP and ALCM); statistical analyses (all authors); interpretation of the results (all authors); drafting the review manuscript (all authors). Revision of the content (RR). All authors read and approved the final manuscript.

**Conflict of Interests**

The authors have no conflict of interests to declare.

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