Valerian Root in Treating Sleep Problems and Associated Disorders—A Systematic Review and Meta-Analysis

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Abstract
Sleep problems are widely prevalent and associated with various comorbidities including anxiety. Valerian (Valeriana officinalis L.) is a popular herbal medicine used as a sleep aid, however the outcomes of previous clinical studies are inconsistent. This study was conducted to update and re-evaluate the available data in order to understand the reason behind the inconsistent outcomes and to provide a broader view of the use of valerian for associated disorders. PubMed, ScienceDirect, and Cochrane Library were searched to retrieve publications relevant to the effectiveness of valerian as a treatment of sleep problems and associated disorders. A total of 60 studies (n=6,894) were included in this review, and meta-analyses were performed to evaluate the effectiveness to improve subjective sleep quality (10 studies, n=1,065) and to reduce anxiety (8 studies, n=535). Results suggested that inconsistent outcomes were possibly due to the variable quality of herbal extracts and that more reliable effects could be expected from the whole root/rhizome. In addition, therapeutic benefits could be optimized when it was combined with appropriate herbal partners. There were no severe adverse events associated with valerian intake in subjects aged between 7 and 80 years. In conclusion, valerian could be a safe and effective herb to promote sleep and prevent associated disorders. However, due to the presence of multiple active constituents and relatively unstable nature of some of the active constituents, it may be necessary to revise the quality control processes, including standardization methods and shelf life.

Keywords
valeriana officinalis, herbal medicine, sleep, insomnia, anxiety, systematic review

Introduction
Sleep Problems
Sleep plays a crucial role in maintaining brain functions and systemic physiology, and chronic sleep problems could have a significant impact on our health. Insufficient sleep leads to reduced stress resilience, decreased quality of life, mood disorders, and cognitive, memory, and performance deficits. It can also contribute to metabolic disorders including hypertension, dyslipidemia, cardiovascular disease, and type 2 diabetes mellitus. In addition, it is associated with significantly increased dementia risk. Problems with sleep are widely prevalent, affecting 70 million (9-20%) of adults in the US and 45 million (7%) of adults in Europe. Typical manifestation of sleep disorders include sleep deprivation or fragmentation and events that occur during sleep. The major sleep disorders are insomnia, restless leg syndrome (RLS), obstructive sleep apnea syndrome, and narcolepsy, among which insomnia is the most common. Sedative-hypnotic medications commonly prescribed for insomnia are GABA_A receptor agonists, antidepressants, and antihistamines. However, the long term use of most sedative-hypnotic drugs is limited due to various side effects, such as cognitive and daytime performance impairments. In recent years,
herbal supplements, such as valerian (Valeriana officinalis L.), sour jujube seeds (Ziziphus jujuba Miller var. spinosa Hu ex H. F. Chou), and kava (Piper methysticum G.Forst.), have gained popularity as alternatives to prescription drugs to improve sleep quality without side effects.15

Valerian for Sleep Problems and Associated Disorders

Recorded medicinal use of valerian dates back to the first century AD. In recent years, it is popular as a sedative and hypnotic. Historically, however, its metabolic stimulating features such as diuretic, carminative, and menstrual stimulant properties had been more valued.16 It was in the middle ages that the use of valerian in treating nervous disorders and insomnia was recorded.16,17 There are over 200 valerian species worldwide, including V. wallichii DC., V. edulis Nutt., and V. fauriei Briq., among which Valeriana officinalis L. is the most well known in Europe and North America as “valerian.” In the United States, valerian is regulated by the Food and Drug Administration (FDA) as a dietary supplement (https://ods.od.nih.gov/factsheets/Valerian-HealthProfessional/). According to the European Medicine Agency (EMA), the well-established uses of V. officinalis root include the relief of mild nervous tension as well as sleep disorders. For relief of nervous tension, recommended oral dosages are 400-600 mg dry hydroalcoholic extract or comminuted herbal substance (root) 0.3-3 g up to 3 times daily.18 Valerian is considered relatively safe and well-tolerated, however EMA monograph notes gastrointestinal symptoms (e.g. nausea, abdominal cramps) as undesirable effects. Although hydroalcoholic extracts of valerian root in the recommended dosage improve sleep latency and quality, it is uncertain what constituents contribute to the efficacy.18

The effectiveness of valerian as a sleep aid has been the major research focus, and several systematic reviews were conducted previously. A systematic review published in 2000, which analyzed 9 randomized clinical trials, found contradictory results and significant inconsistency in terms of patients, experimental design and methodology among the trials.19 Another systematic review and meta-analysis, published in 2006, analyzed 16 studies and this study also found significant methodological problems.20 Taibi et al. (2007) conducted a systematic review on 37 studies, of which 29 were controlled trials and 8 were open-label trials. They concluded that, although it is a safe herb, evidence did not support the clinical efficacy of valerian as a sleep aid for insomnia.21 A meta-analysis of 18 randomized placebo-controlled trials, published in 2010, concluded that valerian’s effectiveness had not been demonstrated with quantitative or objective measures although valerian could improve subjective sleep quality.22 So far, these inconsistent outcomes have not been fully explained. In addition, it is not clear if valerian is effective in treating other disorders associated with, and possibly contributing to, sleep problems. The aim of the current study is to up-date the available data, to evaluate the effectiveness of valerian as a treatment of sleep problems and associated disorders, and to discuss possible reasons behind the inconsistent research outcomes, by particularly focusing on the herbal preparations used in the studies.

Summary

In summary, this study will focus on:

- The herbal materials – extracts or whole root/rhizomes – that have been used in trials.
- Potential applications of valerian, including but looking beyond its use for sleep problems.
- Popular and effective herbal combinations.

Methods

In order to evaluate the therapeutic effects of valerian, systematic reviews and meta-analyses were conducted following PRISMA guidelines.23 Searches A-C were conducted using PubMed (https://www.ncbi.nlm.nih.gov/pubmed), ScienceDirect (https://www.sciencedirect.com/search/advanced), and Cochrane Library (https://www.cochranelibrary.com/search).

Search Strategy

Search A: Clinical studies on valerian, published between January 1980 and November 2019, were retrieved using search term “Valeriana” in PubMed database under article type: Clinical Trial.

Search B: ScienceDirect advanced search was conducted using search term “Valeriana clinical trial.” Articles published between January 2000 and November 2019 were sought.

Search C: To identify clinical trials using valerian, the Cochrane Library was searched (up to December 2019) using the search term ‘Valeriana officinalis’.

Study Selection

Studies were screened initially for relevance based on the title and abstract. Reviews, unrelated studies, and works without available full text were excluded. Studies evaluated the effectiveness of valerian alone or in combination on sleep or related health problems were considered relevant. The full texts of potential studies were assessed following the exclusion and listed below.

Exclusion criteria

1. Articles published in any non-English language.
2. Studies using unknown substance.
3. Studies on non-human subjects.

The quality of randomized trials were assessed using Jadad scale.24 In order to extensively collect currently available information, both controlled trials and observational studies were included for reviewing, irrespective of potential risk of bias (Figure 1).
Meta-Analysis

The effectiveness of *V. officinalis* in promoting sleep and reducing anxiety were evaluated by meta-analyses. Due to the scarcity of data, all placebo-controlled studies reporting values that are convertible to effect sizes were included for statistical analysis, irrespective of trials’ quality (Jadad scale). The outcome variables were extracted from *V. officinalis* treatment group and placebo group by 1 author, which was checked by 2 authors. In order to evaluate the effectiveness as a sleep aid, data on subjective ‘sleep quality’ improvement by repeated administration (ranging between 5 days to 8 weeks) were included in the analysis. Due to the variable measures, both numerical scores and binary scores were converted to effect sizes for the analysis.25 To evaluate the anxiolytic effect, a study reporting the overall emotional symptoms including anxiety and a study reporting binary outcomes, in addition to the studies reporting standard anxiety test scores, were included. When several measures were reported over time in the same study, the one with the longest follow-up period was selected. When more than 1 dose were tested, the result of the most effective dose was selected. Adjusted effect sizes (Hedges’ *g*) were calculated from summary measures such as means and standard deviations or confidence intervals, odds ratio (for binary outcomes), and sample sizes, using reported formula.22-27 Meta-analyses were performed using Meta-Essentials.28 I² statistic was used to estimate heterogeneity. Publication bias was evaluated using the funnel plots.29

Results

Based on the inclusion and exclusion criteria described above, a total of 60 articles were selected for reviewing (Table 1 and 2). Both quantitative and qualitative studies with human subjects were included. Of 40 articles using valerian as a single herb, 36 studies were randomized controlled trials (RCT)30-65 and 2 studies were observational.66,67 Two studies addressed the potential induction of liver enzyme cytochrome P450 (CYP) isoforms.68,69 In addition, 2 RCTs were conducted using valepotriates (80% didrovaltrate, 15% valtrate, and 5% acevaltrate)68% (Table 1). Nineteen articles assessed the effectiveness of herbal combinations, of which 13 studies were RCT65,72,73-80,81-83 and 6 were observational studies84-89 (Table 2).

**V. Officinalis as a Single Herb**

**Sleep problems.** To evaluate the effectiveness of valerian for sleep problems, 40 articles using *V. officinalis* as a single herb were assessed. As a result, it was found that 23 studies addressed the effectiveness of valerian for sleep problems. Among them, 13 studies found valerian effective as a sleep aid. Three studies were performed with healthy subjects,37,40,41 4 studies with insomnia patients,36,42,52,66 and 1 study in RLS patients.30 In addition, studies conducted with mental health patients,67 cancer patients,56 HIV-positive patients,35 postmenopausal women,57 and the elderly,38 also found that valerian could improve sleep quality.35,38,56,57 On the other hand, 10 studies found that valerian was not significantly effective compared to control25,36,38,40,48,50,51,53,54,65 at least for the outcome measures used in those studies. Study outcomes were assessed using various measures, such as sleep questionnaires and diaries, such as Pittsburgh Sleep Quality Inventory (PSQI),90 Insomnia Severity Index (ISI),91 and Epworth Sleepiness Scale (ESS), to polysomnography.92

Six studies measured immediate responses after single dose administration,36,38,40,42,50 while most studies evaluated the effectiveness of repeated doses.30,35,54,56,57,65,67,36-38,44,48,51-53 The intervention periods ranged from 5 days to 8 weeks.

In order to extract possible associations between herbal preparations and study outcomes, the reports were classified into 4 groups (Table 1): 8 studies using hydroalcoholic extracts,36,42,44,50,52-54,65 3 studies using aqueous extracts,38,40,41 3 studies using extracts using unspecified solvents,37,48,51 and 5 studies using herbal substance (the whole root / rhizome).30,35,56,57,67 Among the 3 studies using aqueous extracts, 2 studies with single dose found improved subjective sleep quality and latency in healthy volunteers40,41 (RCTs [Jadad scale 2 and 5]), while there was no significant difference in polysomnography40 (RCTs [Jadad scale 2]). Schulz et al. found that repeated administration of *V. officinalis* aqueous extract (for 1 week) increased deep sleep (slow-wave sleep: SWS) in elderly subjects, although single dose of the same preparation did not have any significant impact on polysomnography38 (RCT [Jadad scale 2]). As a single dose, *V. officinalis* hydroalcoholic extracts did not improve sleep quality36,50 (RCT, Jadad scale 2-3), however improved REM sleep was observed in insomnia patients42 (RCT [Jadad scale 2]). Repeated administration of hydroalcoholic extracts also led to inconsistent outcomes: sleep quality was improved in 4 studies using extract 600 mg (2 to 6 weeks)36,37,44,52 (RCTs [Jadad scale 2-5]), whereas 3 studies using extract 300-600 mg (5 days to 4 weeks) found no improvement53,54,65 (RCTs [Jadad scale 3-5]). Donath et al. found improved sleep latency and deep sleep after 2 weeks, while no improvement was observed after a single dose36 (RCTs [Jadad scale 2]), which suggests that repeated administration is needed to exert observable effects. There were 3 studies using extracts with unspecified procedures.38,51,66 The outcomes from those studies were also inconsistent: negative outcomes for 2 studies38,51 (RCTs [Jadad scale 5]) and positive for an observational study.66 On the other hand, all 5 studies using the dried root/rhizome showed that the interventions led to improved sleep at least in 1 subgroup30,35,56,57,67 (RCTs [Jadad 3-5] and 1 observational study), suggesting that maximum efficacy could be expected from the root / rhizome before extraction.

It has been suggested that valerian may be useful to improve subjective sleep quality22 and that repeated administration is required to obtain significant effects.36 In order to evaluate if repeated administrations could consistently improve sleep quality, a meta-analysis was carried out (Figure 2). Studies using repeated valerian administrations were assessed to extract numerical values for the meta-analysis, and 10 randomized placebo-controlled trials reporting quantifiable results30,34,36,38,39,54,56,57 were included in the analysis. The combined effect size was 0.36 (95% CI: -0.08 to 0.81) (Figure 2A). The funnel plot indicated missing data (Figure 2B) and high
Table 1. Valeriana spp. as a Single Herb.

| First Author | Study design | Intervention and measures | Outcomes | Adverse events |
|--------------|--------------|---------------------------|----------|----------------|
| **Hydroalcoholic extract** | 1 Roh D 2019 | Randomized, placebo-controlled, double-blind.  
*Population*: nonclinical volunteers suffering psychological stress (n=52) | **Intervention**:  
*V. officinalis 70%EtOH* extract capsule 300 mg (0.8% valerenic acid) per day (100 mg t.i.d.) for 4 weeks.  
**Measures**:  
1. Psychiatric variables (HAM-A, HAM-D, PSQI).  
2. Functional brain connectivity changes (EEG)  
**Summary measures for meta-analysis**:  
- difference in means |  
- 1. Improvement in both treatment and control groups. No statistically significant difference between the groups.  
- 2. Greater increases in frontal brain region alpha coherence in valerian group, which was correlated with anxiolysis | No moderate or serious adverse reaction.  
Two adverse effects (one from each group): atypical chest pain (unrelated to the medication) in treatment group and gastrointestinal problems in the placebo group. |
| | 2 Taibi DM 2009 | Randomized, placebo-controlled, single-blind, pilot study.  
*Population*: Arthritis patients with mild sleep disturbance (n=15) | **Intervention**:  
*V. officinalis 70%EtOH* extract (0.8% valerenic acid) 600 mg for 5 nights  
**Measures**: Sleep outcomes (sleep diaries and wrist actigraphy)  
**Summary measures for meta-analysis**:  
- difference in means |  
- No significant difference between the groups. | Dizziness and sleepiness  
(1) Self-rated  
(2) n/a  
(3) 5 |
| | 3 Taibi DM 2009 | Randomized, placebo-controlled, double-blind, cross-over  
*Population*: Older women with insomnia (n=16) | **Intervention**:  
*V. officinalis 70%* extract 300 mg (0.8% valerenic acid), 30 min before bedtime for 2 weeks  
**Measures**: (1) polysomnography (PSG); (2) self-rated sleep  
**Summary measures for meta-analysis**:  
- difference in means |  
- No significant difference between the groups in sleep latency, wake after sleep onset, sleep efficiency, and self-rated sleep quality. | No severe adverse events.  
No difference between valerian and placebo  
(1) Self-rated  
(2) n/a  
(3) 5 |
| | 4 Oxman AD 2007 | Randomized, placebo-controlled, double-blind (Web-based trial)  
*Population*: Volunteers with insomnia without treatment (n=405) | **Intervention**:  
*V. officinalis 60%EtOH* extract 600 mg per day, 1 h before bedtime for 14 days.  
**Measures**: Sleep diary (sleep onset latency, number of night awakenings, sleep duration, sleep quality, energy level the following day).  
**Summary measures for meta-analysis**:  
- difference in means  
- Global self-assessment of change was better for valerian group (p=0.04).  
- Similar trends favoring the treatment group for sleep quality, night awakenings, and sleep duration. |  
- No serious adverse events. No statistically significant differences in minor adverse events between the groups. |  
(1) Self-rated  
(2) n/a  
(3) 5 |
| | 5 Koetter U 2007 | Randomized, placebo-controlled, double-blind  
*Population*: Patients suffering from non-organic sleep disorders (n=27) | **Intervention**:  
*V. officinalis 45%MeOH* extract 500 mg compared to placebo. For 4 weeks.  
**Measures**: (1) the reduction of the initially prolonged sleep latency; (2) WASO, sleep efficiency, relative proportion of the sleep stages, REM latency and CGI |  
- No significant difference between valerian extract and placebo. | No adverse events.  
(1) n/a  
(2) n/a  
(3) 3 |
| First Author | Study design | Intervention and measures | Outcomes | Adverse events |
|--------------|--------------|----------------------------|----------|----------------|
| Diaper A 2004 | Randomized, placebo-controlled, double-blind, cross-over | Intervention: (1) *V. officinalis* 70%EtOH extract (LI 156) 300 mg; (2) *V. officinalis* 70%EtOH extract 600 mg; (3) placebo. Single dose, separated by 6 days washout. Measures: Sleep EEG, CFF, CRT, SMT, LARS, LSEQ, SRTI, and STAI. | No significant differences between the 3 groups. | Mild or moderate adverse events: Drowsiness was 4 under 600 mg valerian, 12 under 300 mg valerian, and 8 in placebo. |
| Donovan JL 2004 | Open-label, fixed treatment, cross-over | Intervention: *V. officinalis* 70%EtOH extract 1000 mg (containing 11.02 mg valerenic acid) nightly for 14 days. Measures: CYP2D6 and CYP3A4 activities. | *V. officinalis* has a moderate effect on CYP3A4 (increase in maximum probe drug concentration), however there were no significant differences in other pharmacokinetic parameters. | No treatment-associated adverse events |
| Hallam KT 2003 | Placebo-controlled, double-blind, cross-over | Intervention: Single doses of *V. officinalis* 70%EtOH extract (0.25% valerenic acids) 500 mg or 1000 mg. Compared to triazolam 0.25 mg. Measures: Cognitive and psychomotor performance (CFF, CRT, DSST, SST, DST) and mood (VAS). Summary measures for meta-analysis: difference in means. | Single doses of valerian were without effect on either cognitive or psychomotor performance, while triazolam had detrimental effects on cognitive processes, in healthy volunteers. Although not statistically significant, valerian 500 mg may have tranquilizing / calming effect. | |
| Ziegler G 2002 | Randomized, double-blind, comparative | Intervention: *V. officinalis* 70%EtOH extract (LI 156 sedonium) 600 mg for 6 weeks. Compared with oxazepam. Measures: Sleep questionnaire. | Both Valerian and oxazepam improved sleep quality. | Mild or moderate adverse events in both oxazepam (36%) and Valerian (28%) groups. |
| Herrera-Arellano A 2001 | Randomized, controlled, double-blind, cross-over | Intervention: Single doses of (1) *V. officinalis* hydroalcoholic extract 450 mg or (2) *V. edulis* (hydroalcoholic extract) 450 mg. Prior to lights out. Compared to basal levels. Measures: Hypnotic effects (polysomnography), anterograde memory (measured by D. Wechsler Memory Scale). | *V. officinalis* and *V. edulis* increased the REM sleep (better improved by *V. officinalis*), and morning sleepiness. *V. edulis* reduced the number of awakening episodes. | Slight dyspepsia (2 for *V. officinalis* and one for *V. edulis*). |

(continued)
| First Author | Study design | Intervention and measures | Outcomes | Adverse events *
---|---|---|---|---
11 Donath F 2000 | Randomized, placebo-controlled, double-blind, cross-over | **Intervention:** V. officinalis 70%EtOH extract (Sedonium) 600 mg, 1 h before bedtime. (1) Single dose, (2) 14 days. **Measures:** Objective sleep efficiency (SE) and sleep onset latency (SOL), NREM, REM, and slow wave sleep (SWS). Subjective sleep quality (VAS). **Summary measures for meta-analysis:** difference in means | (1) There was no significant effect after a single dose. (2) SE was significantly higher in both groups at the end of the trial period. SWS latency was significantly shorter and SWS increased in valerian group. A tendency toward reduced subjective sleep latency was observed despite there was no significant difference in objective SOL between the groups. | Fewer adverse events for Valerian group than placebo. (1) Self-rated (2) n/a (3) 2
12 Kuhlmann J 1999 | Randomized, placebo-controlled, double-blind | **Intervention:** V. officinalis 70%EtOH extract (600 mg, L1 156). Single dose in the evening (compared to placebo and Flunitrazepam (FNZ)), and repeated evening administration for 2 weeks (compared to placebo). **Measures:** (1) Sleep quality and latency (self-assessment). (2) Reaction time (Vienna Determination Test), alertness, and concentration (2-handed co-ordination) at next morning. | (1) Sleep quality and latency improved after single doses of valerian and FNZ. After 2 weeks with Valerian, sleep quality had a trend toward improvement, compared to placebo. (2) Neither single dose nor repeated administrations of valerian extract had a negative impact on reaction time, alertness, and concentration. | No serious adverse events. Headache and weakness were most frequently reported, which had not inter-group difference (no causal relationship to valerian). Note: Single dose of FNZ caused hangover effect in 59.4%, while only 32.4% of placebo and 30.3% of valerian-treated groups reported this effect. (1) n/a (2) n/a (3) 3

**Aqueous extract**

13 Pakseresht S 2011 | Randomized, placebo-controlled, double-blind | **Intervention:** V. officinalis aqueous extract 750 mg per day (250 mg t.i.d.) for 8 weeks **Measures:** Yale-Brown scale for OCD (Y-BOCS) | Significantly reduced Y-BOCS score in the treatment group compared to placebo. | No significant difference in observed side effects, however increased somnolence in the treatment group (53.3% in treatment group and 18.75% in placebo group) (1) n/a (2) n/a (3) 5
14 Schulz H 1994 | Randomized, placebo-controlled, double-blind | **Intervention:** V. officinalis aqueous alkaline extract (Valdispert forte): (1) 405 mg single administration 1 h before sleep recording; (2) 405 mg t.i.d. for 7 days. **Measures:** Objective (polysomnography) | (1) Acute administration had no effect. (2) V. officinalis increased SWS and K-complex density, and decreased sleep stage I. (SWS increased in each single subject on valerian.) No effect on sleep onset time, time awake after sleep onset, or self-rated sleep quality. | Not noted. (1) n/a (2) n/a (3) 2

(continued)
| First Author | Study design | Intervention and measures | Outcomes | Adverse events |
|--------------|--------------|--------------------------|----------|----------------|
| 15 Balderer G | Randomized, placebo-controlled, double-blind, cross-over | **Intervention**: Single doses of *V. officinalis* aqueous extract (450 or 900 mg) or placebo 30 min before bedtime. At home (n=10) and in the laboratory (n=8). Measures: Subjective: sleep questionnaires, self-rating scale, night-time motor activity (n=18). Objective: polysomnography (n=8). | Home: *V. officinalis* dose-dependently reduced subjective sleep latency and wake time after sleep onset. Self-rating of subjective quality showed no difference (data not shown). Laboratory: No statistically significant difference between valerian and placebo. | (1) n/a (2) n/a (3) 2 |
| 16 Leathwood PD | Randomized, placebo-controlled, blinded | **Intervention**: Single doses of (1) *V. officinalis* aqueous extract 400 mg; (2) over-the-counter preparation containing valerian and hops (Zyma, Zyma S.A.); (3) placebo. 1 h before bedtime. Each treatment 3 times, randomly on non-consecutive nights. Measures: Sleep latency and quality, dream recall, night awakenings, somnolence the next morning (sleep questionnaire). | *V. officinalis* produced significant decrease in subjective sleep latency and improved sleep quality. Sleep quality improvement was most notable among poor or irregular sleepers and smokers. Proprietary preparation produced 'hangover effect' the next morning, which was absent after valerian intake. | No notable side effects (1) n/a (2) n/a (3) 5 |
| Unknown extract | Randomized, split-mouth, controlled, double-blind, cross-over | **Intervention**: *V. officinalis* extract 100 mg. Control drug: midazolam 15 mg. Both in an identical-looking capsules. 60 min before the surgical procedures. Measures: Oxygen saturation, HR, systolic and diastolic BP, RR, sedation (Ramsay scale). Self-assessment of anterograde amnesia. | More patients with valerian were rated as calm. No difference between valerian and midazolam (benzodiazepine) in oxygen saturation. HR and BP were lower with midazolam. Midazolam caused somnolence and anterograde amnesia, while valerian did not. | The most commonly reported side effect was somnolence. No statistically significant difference between the 2 drugs. (1) n/a (2) n/a (3) 5 |
| 17 Farah GJ | Randomized, placebo-controlled, double-blind, cross-over. | **Intervention**: Single dose of *V. officinalis* extract capsule (300 mg with 0.8% valerenic acid) | Significant reduction in intracortical facilitation (ICF) at 1 h after Valerian extract intake, without affecting other TMS parameters. ICF returned to normal at 6 h. | No adverse events (1) n/a (2) n/a (3) 4 |
| 18 Mineo L | Randomized, placebo-controlled, double-blind, cross-over. | **Intervention**: *V. officinalis* extract 630 mg twice daily, in the last 7 days of menstrual period for 3 cycles. Measures: Severity of emotional, behavioral, and physical premenstrual symptoms (questionnaire) | Valerian improved emotional, behavioral, and physical premenstrual symptom severity, whole placebo had no effects. | Not noted (1) n/a (2) self-rated emotional state (3) 5 |
| Behboodi Moghadam Z | Randomized, placebo-controlled, double-blind | **Intervention**: *V. officinalis* extract 400 mg | | | (continued) |
| First Author | Study design | Intervention and measures | Outcomes | Adverse events | * |
|--------------|--------------|---------------------------|----------|----------------|---|
| 20 Jacobs BP 2005 | Randomized, placebo-controlled, double-blind (internet-based) | Population: Volunteers with anxiety and insomnia (n=391). Recruited via e-mail and banner advertisements on websites | Intervention: (1) *V. officinalis* extract (6.4 mg (1%) valerenic acid); (2) *Piper methysticum* extract (100 mg total (30%) kavalactones); (3) placebo. For 28 days | Similar decrease in anxiety and insomnia symptoms among the 3 groups. | Diarrhea (valerian 18%, Kava 11%, control 8%) | (1) ISI (2) STAI (3) 5 |
| 21 Gurley BJ 2005 | Cross-over | Population: Healthy volunteers (n=12) | Intervention: (1) *V. officinalis* extract (125 mg 3 times daily); (2) *Hydrastis canadensis* (900 mg 3 times daily); (3) *Piper methysticum* (100 mg twice daily); (4) *Cimicifuga racemosa* (1090 mg twice daily). For 28 days separated by 30 days washout. | No significant changes in phenotypic ratios were observed for *V. officinalis*. (H. canadensis inhibited CYP2D6 and CYP3A4/5, P. methysticum reduced CYP2E1, and C. racemosa inhibited CYP2D6.) | No serious adverse events | (1) n/a (2) n/a (3) n/a |
| 22 Gutierrez S 2004 | Randomized, placebo-controlled, double-blind, cross-over | Population: Healthy volunteers (n=10) | Intervention: Single doses of *V. officinalis* standardized extract 600, 1200, and 1800 mg | No significant difference from placebo. | | |
| 23 Coxeter PD 2003 | Randomized, placebo-controlled, double-blind, n-of-1 | Population: Patients with chronic insomnia (n=24) | Intervention: *V. officinalis* extract 450 mg equivalent to 2000 mg dry root / rhizome (5.88 mg valerenic acids, 0.96 mg valeranal, and 2.46 mg valtrates), 30 min before bed for 3 weeks. 5 days wash-out. | Fair response for energy level in the previous day, but only modest response for other variables. | There was no significant difference in the number, distribution, or severity of side effects between valerian and placebo. | (1) n/a (2) n/a (3) 5 |
| 24 Cropley M 2002 | Randomized, controlled | Population: Young healthy volunteers (n=54) | Intervention: (1) *V. officinalis* standardized extract 600 mg (Lichtwer-Pharma UK Ltd); (2) *Piper methysticum* (120 mg) For 7 days. Compared to non-placebo control. | *V. officinalis* (and *Piper methysticum*) reduced psychological pressure and systolic blood pressure responsivity during stress task. *V. officinalis* reduced heart rate reaction to mental stress. | | (1) n/a (2) n/a (3) 1 |
| First Author | Study design | Intervention and measures | Outcomes | Adverse events |
|-------------|-------------|----------------------------|----------|---------------|
| Wheatley D 2001 | Observational, cross-over | **Intervention:** (1) *Piper methysticum* 120 mg; (2) *V. officinalis* 600 mg (Lichtwer-Pharma UK Ltd). For 6 weeks. 2 weeks wash-out. *P. methysticum* first. **Measures:** Severity of stress (VAS) and sleep disturbance (subjective). | **Severity of stress:** Significant reduction during the first 6 weeks on *P. methysticum*. There was no further change during the second 6 weeks on *V. officinalis*. **Severity of insomnia:** Score fell after he first 6 weeks on *P. methysticum*, and further fell after the final 6 weeks on *V. officinalis*. | 1. Vivid dreams, daytime drowsiness, heavy sleep and depression for *V. officinalis*. 2. Dry mouth, gastric disturbance, diarrhea, dizziness, craving for trial drug, and depression for *P. methysticum*. |
| Kohnen 1988 | Randomized, placebo-controlled, double-blind | **Intervention:** Single doses of (1) Valerian extract 100 mg, (2) propranolol (20 mg), (3) placebo, and (4) valerian + propranolol. **Measures:** Pulse frequency, test performance, psychological strain, and somatic arousal under stress conditions after intervention. | Valerian induced a slight improvement in the concentration task, while propranolol and valerian-propranolol combination reduced the performance values. Valerian reduced the feelings of somatic arousal. Valerian had no effect on psychological strain. | Not noted |
| Samaei A 2018 | Randomized, placebo-controlled, double-blind, cross over | **Intervention:** *V. officinalis* root/rhizome 530 mg, 60 min before bedtime for 1 month. **Measures:** Cognitive brain functions (Mini-Mental State Examination: MMSE) and EEG | Valerian significantly improved cognitive status (MMSE), although no significant changes were observed in EEG. | Not noted. |
| Ahmadi M 2017 | Randomized, placebo-controlled, double-blind | **Intervention:** *V. officinalis* root/rhizome 530 mg (> 0.05% valerenic acid), nightly 1 h before sleep for 4 weeks **Measures:** Neuropsychiatric status (HAM-D, HAM-A, PANSS, PANSI, and PSQI) Summary measures for meta-analysis: difference in means | Sleep and anxiety significantly improved in Valerian treatment group. | There was no significant difference between the groups. |
| Jenabi E 2017 | Randomized, placebo-controlled, triple-blind | **Intervention:** *V. officinalis* root/rhizome 225 mg 3 times per day for 8 weeks **Measures:** Severity and frequency of hot flashes (Kupperman index) | Significantly lower severity after 1 and 2 months and reduced frequency after 2 months in *V. officinalis* treatment group. | No side effects |

(continued)
| First Author | Study design | Intervention and measures | Outcomes | Adverse events |
|--------------|--------------|--------------------------|----------|----------------|
| Thomas K 2016 | Randomized, placebo-controlled, double-blind, cross-over | **Intervention:** Single dose of *V. officinalis* root/rhizome 1600 mg (containing 0.8% valerenic acid)  
**Measures:** SVRT, subjective sleepiness scales (Karolinska Sleepiness Scale), SFST performance scores, driving simulator performance | There were no significant differences in SVRT, sleepiness scales, SFST, and driving simulator performance | (1) n/a  
(2) n/a  
(3) 5 |
| Hassani S 2015 | Randomized, placebo-controlled, double-blind | **Intervention:** *V. officinalis* root/rhizome 1,060 mg per day (530 mg twice daily, every 12 h) for 8 weeks  
**Measures:** Cognitive brain function (MMSE test) | MMSE score was restored on the 60th day in valerian treatment group, whereas reduced in placebo group. | (1) n/a  
(2) n/a  
(3) 5 |
| Pinheiro ML 2014 | Randomized, placebo-controlled, double-blind, split-mouth | **Intervention:** *V. officinalis* comminuted root/rhizome 100 mg, 1 h prior to each surgical procedure  
**Measures:** Anxiety (assessed by surgeons and researchers) and physical parameters | Valerian significantly reduced anxiety. Valerian was effective in maintaining systolic blood pressure and heart rate after surgery. | (1) n/a  
(2) Surgeon-assessed anxiety  
(3) 5 |
| Mirabi P 2013 | Randomized, placebo-controlled, double-blind | **Intervention:** *V. officinalis* root/rhizome 675 mg per day (225 mg t.i.d), for 8 weeks  
**Measures:** Severity and frequency of hot flashes | Valerian reduced the severity and frequency of hot flashes | –  
(1) n/a  
(2) n/a  
(3) 4 |
| Mirabi P 2011 | Randomized, placebo-controlled, double-blind | **Intervention:** *V. officinalis* root/rhizome 675 mg per day (225 mg t.i.d) for 3 days from the first day of menstruation, continued for 2 menstrual cycles.  
**Measures:** Pain severity (VAS, 3 times per day), the severity of associated systemic symptoms (fatigue, diarrhea, syncope, nausea and vomiting, lack of energy, headache, and mood swings) | The pain severity was reduced in both groups, but the extent of the reduction was larger in the treatment group. There was no difference between the groups in the total scores of associated systemic manifestations, except for syncope (improvement in *V. officinalis* treatment group) | No adverse effects  
(1) n/a  
(2) n/a  
(3) 5 |
| Taavoni S 2011 | Randomized, placebo-controlled, triple-blind | **Intervention:** *V. officinalis* root/rhizome 530 mg (Sedamin) twice daily for 4 weeks  
**Measures:** PSQI | Significant improvement in PSQI score and improved percentage in the intervention group compared to placebo. | No reported adverse effects  
(1) PSQI  
(2) n/a  
(3) 4 |
| First Author | Study design | Intervention and measures | Outcomes | Adverse events |
|--------------|--------------|---------------------------|----------|---------------|
| **36 Barton DL 2011**<sup>56</sup> | Randomized, placebo-controlled, double-blind | **Intervention:** *V. officinalis* root/rhizome 450 mg (containing 0.8% valerenic acid) 1 h before bedtime for 8 weeks **Measures:** (1) PSQI. (2) Functional outcomes of sleep questionnaire, fatigue (BFI), and mood (POMS). **Toxicity** | (1) No difference in PSQI score. (2) Significantly better fatigue end points (BFI and POMS), less trouble with sleep, and less drowsiness in the treatment group over control. | No difference in toxicities, except for alkaline phosphatase increase (slightly more common in the placebo group). |
| **37 Cuellar NG 2009**<sup>30</sup> | Randomized, placebo-controlled, triple-blind | **Intervention:** *V. officinalis* root/rhizome 800 mg (containing 1.16 mg valerenic acid) per day 60 min before bedtime for 8 weeks **Measures:** Sleep disturbances (PSQI and ESS) and severity of RLS | Although there was no statistically significant difference valerian group showed a greater improvement in the majority of sleep quality and RLS scores. The most obvious changes were seen in participants with ESS score of ≥10. | GI disturbances, fatigue, vivid dreams, agitation/restlessness, headache, dizziness, rash |
| **38 Glass JR 2003**<sup>45</sup> | Randomized, placebo-controlled, double-blind, cross-over. | **Intervention:** Single doses of *V. officinalis* root/rhizome 400 or 800 mg (containing 0.683% valerenic acids). Compared to temazepam, diphenhydramine, and placebo. **Measures:** Psychomotor performance (Manual Tracking Test, DSST, manual tracking and digit symbol substitution test); sedation and mood (VAS) | Valerian was not different from placebo on any measure of psychomotor performance, while temazepam and diphenhydramine caused psychomotor impairment. No sedation or any side effects as a result of valerian treatment. | (1) n/a (2) n/a (3) 2 |
| **39 Francis AJ 2002**<sup>46</sup> | Randomized, placebo-controlled, double-blind, cross-over. | **Intervention:** *Valeriana edulis* (dried and crushed whole root, containing 1.1% of valtrate/isovaltrate) 20mg/kg body weight, nightly 1 h before bedtime for 2 weeks. Wash-out 7 days. **Measures:** Sleep latency, time spent awake during the night, and sleep quality (VAS). | *V. edulis* treatment significantly reduced sleep latencies and nocturnal time awake, lengthened total sleep time, and improved sleep quality. The treatment was most effective in children with hyperactivity. | No significant adverse effects. |
| **40 Dominguez RA 2000**<sup>57</sup> | Observational | **Intervention:** *V. officinalis* root/rhizome (Nature's Way) 470 mg each night 30-60 min prior to retiring (possibly increased up to max. 1410 mg after 1 week) for 4 weeks. **Measures:** Sleep questionnaire | Week 1: 16 patients rated moderately improved. Week 2: 15 patients rated more than moderately improved. Global improvement at week 2 was significantly better at week 2 than week 1. | (1) n/a (2) n/a (3) n/a |

**Isolates**
| First Author | Study design | Intervention and measures | Outcomes | Adverse events |
|--------------|--------------|---------------------------|----------|---------------|
| **41 Andreatini R 2002** | Randomized, placebo-controlled, double-blind | **Population:** Outpatients with generalized anxiety disorders (n=36)  
**Intervention:** (1) Valepotriates (80% dihydrovaltrate, 15% valtrate, and 5% acevaltrate) (average dose: 81.3mg/day); (2) placebo; (3) diazepam (average dose:6.5mg/day). Flexible doses according to the patient's therapeutic response. 4 weeks.  
**Measures:** Anxiety (HAM-A and STAI). | Total HAM-A score: significant reduction in all 3 groups. (No significant difference among the 3 groups.)  
Psychological factor of HAM-A: Reduced in valepotriates and diazepam groups. | No moderate or serious adverse reaction was observed.  
(1) n/a  
(2) n/a  
(3) 5 |

| **42 Poyares DR 2002** | Randomized, placebo-controlled, double-blind | **Population:** (A) Primary insomnia patients with poor sleep despite chronic use of BZDs (n=19); (B) Healthy individuals (n=18)  
**Intervention:** valepotriates: 80% didrovaltrate, 15% valtrate, and 5% acevaltrate (from *Valeriana wallichii* root). Three times daily in 100 mg doses for 15 days. After BZD withdrawal (over 2 weeks + 48 h without BZD).  
**Measures:** Sleep diaries and sleep parameters (sleep latency, total sleep time, sleep efficiency, number of arousals, WASO, REM and NREM; sleep EEG) | Valerian group reported significantly better sleep quality than placebo during the second week of treatment.  
Sleep EEG data showed a significant increase in alpha count during Sleep stage 3 and 4 NREM in valerian group.  
Valerian group showed a significant decrease in WASO compared to placebo, while placebo group presented shorter sleep latency. | Gastrointestinal symptoms (diarrhea, stomachache, and bitter taste in their mouth) in the first week of valerian treatment.  
(1) n/a  
(2) n/a  
(3) 3 |

* (1) data included in meta-analysis for sleep quality; (2) data included in meta-analysis for anxiety; (3) Jadad scale.

Abbreviations: ARCI-Addiction Research Center Inventory, CFF-Critical Flicker Fusion, CRT-Choice Reaction Time, DSST-Digit Symbol Substitution Test, DST-Digit Span Test, EEG-Electroencephalogram, HAM-A-Hamilton Anxiety Rating Scale, HAM-D-Hamilton Depression Rating Scale, ISI-Insomnia Severity Index, LARS-Line Analogue Rating Scales, LSEQ-Leeds Sleep Evaluation Questionnaire, MMSE-Mini-Mental State Examination, PANSS-Positive and Negative Syndrome Scale, PANSI-The Positive and Negative Suicide Ideation, PSQI-Pittsburgh Sleep Quality Index, SFST-Standardized Field Sobriety Test, SMT-Short term Memory Test, SST-Symbol Search Test, STDI-State Trait Depression Inventory, SVRT-Simple Visual Reaction Test, WASO-Wake Time After Sleep Onset.
### Table 2. Herbal Combinations.

| First Author | Study design | Intervention and measures | Outcomes | Adverse events | Combined with | Jadad scale |
|--------------|--------------|---------------------------|----------|----------------|---------------|-------------|
| 1 Abdellah SA 2019 | Open-label, observational | Intervention: Combination of *V. officinalis* extract 32 mg and *Eschscholtzia californica* extract 80 mg per tablet, Max. 4 tablets per day. For 4 weeks | ISI score decreased by approx. 30%, night sleep duration increased, the number of awakening decreased, and anxiety score significantly decreased by 50%. Sleep latency did not improve. | Mild nocturnal pollakiuria in 1 patient. No serious adverse events. | *E. californica* | n/a |
| 2 Meier S 2018 | Randomized, placebo-controlled, double blind | Intervention: Combination of *V. officinalis* 45%MeOH extract 90 mg, *Passiflora incarnata* 50%EtOH extract 90 mg, *Melissa officinalis* 20%EtOH extract 60 mg (Ze 185) | Attenuated self-reported anxiety. There was no difference between the groups in salivary cortisol levels. | No difference between the groups. | *P. incarnata* | 5 |
| 3 Toolika E 2015 | Observational | Intervention: *V. wallichii* root/rhizome 4 g with milk after meal, 3 times daily (12 g per day) for 1 month. | V. wallichii with milk improved the initiation and duration sleep, reduced sleep disturbance, and improved daily functioning. | Not mentioned | Milk | n/a |
| 4 Gromball J 2014 | Observational study | Intervention: Combination of *V. officinalis* 62%EtOH extract 640 mg and *Melissa officinalis* 30%EtOH 320 mg per day, for 7 weeks. | Doctors’ rating: The treatment improved concentration and reduced hyperactivity and impulsiveness. Parents’ rating: Social behavior, sleep and symptom burden improved. | Two patients had transient moderate adverse reactions, however causal relation with the medication was judged unlikely. | *M. officinalis* | n/a |
| 5 Taavoni S 2013 | Randomized, placebo-controlled, triple-blind | Intervention: Valerian essence 160 mg and lemon balm 80 mg per day for 1 month | Significantly reduced sleep disorder score after 1 month of Valerian/Lemon balm treatment | No adverse events | *M. officinalis* | 3 |

(continued)
| First Author | Study design | Intervention and measures | Outcomes | Adverse events | Combined with | Jadad scale |
|--------------|--------------|---------------------------|----------|---------------|---------------|-------------|
| Maroo N 2013 | Randomized, controlled, double-blind | Population: Primary insomnia patients (n=78) | Intervention: 1. NSF-3: Polyherbal extract of *V. officinalis* (300 mg with 0.8% valerenic acid), *Passiflora incarnata* (80 mg with 4% isovitexin), and *Humulus lupulus* (30 mg with 0.35% rutin) at bedtime for 2 weeks. 2. Control: zolpidem 10 mg | Significant improvement in total sleep time, sleep latency, number of nightly awakenings and severity index score in both groups. No statistically significant difference between the groups. | Mild adverse events (e.g. drowsiness): 12 with herbal treatment (1), 16 with zolpidem (2) | *P. incarnata* |
| Trompeter I 2013 | Observational (multicentre) | Population: Children (6-12 years) suffering from nervous agitation due to affective disorders (n=115) | Intervention: Herbal triplet containing dried extracts of *Valeriana officinalis* (40%EtOH extract 28 mg), *Hypericum perforatum* (38%EtOH extract 60 mg), and *Passiflora incarnata* (60%EtOH extract 32 mg) per tablet. Average dose: 1-3 tablets per day. Duration: 2-4 weeks. | 81.6-93.9% of affected children had no or just mild symptoms (depression, anxiety, sleeping problems, and different physical problems) at the end of observation. | A good tolerability for 97.4% of the children. Moderate to poor tolerability in 7 cases. Paradox reactions: restlessness, weepiness, increased irritability, or aggressiveness; redness in cheekbone region (1 case); stomach pain (1 case) | *H. perforatum* |
| Chen JH 2012 | Randomized, controlled | Population: Stable and less critical ICU patients (n=95) | Intervention: Valerian acupressure (application of 2.5% *V. officinalis* essential oil 2 drops per point before acupressure). Compared to regular treatment | Valerian acupressure increased sleeping hours, reduced wake frequency and SSS grades. Heart rate variability indicated immediate relaxation response. | Acupressure | n/a |
| Melzer J 2009 | Randomized, controlled, double-blind | Population: Somatoform disorder patients (n=167) | Intervention: (1) Ze185 (4-combination) containing 90 mg of *V. officinalis* (45%MeOH extract), 90 mg of *Petasites* (90%EtOH extract), 90 mg of *Passiflora incarnata* (50%EtOH extract), and 60 mg of *Melissa officinalis* (20%EtOH extract); (2) 3-combination without *Petasites*; (3) placebo. 3 times daily for 2 weeks | Significantly reduced anxiety in both treatment groups (3- and 4-combination), but not in placebo. Effectiveness: 4-combination > 3-combination > placebo | Adverse events: (1) 4-combination: vomiting and flatulence, (2) 3-combination: nausea and constipation | *P. incarnata* |

(continued)
| First Author | Study design | Intervention and measures | Outcomes | Adverse events | Combined with | Jadad scale |
|--------------|--------------|---------------------------|----------|---------------|---------------|-------------|
| Dimpfel W    | Randomized, placebo-controlled, double-blind | **Intervention:** V. officinalis (tinct. 1:10) 460 mg and Humulus lupulus (tinct. 1:12) 460 mg. Single administration, 15 min before the lights were turned off at 22:00. **Measures:** (1) Depth of sleep (EEG); (2) Sleep quality (sleep inventory SF-A sub-score) | Time spent in sleep and deeper sleep was significantly higher in the treatment group, which was correlated with the difference in sleep quality. | H. lupulus | 3 |
| Koetter U    | Randomized, placebo-controlled, double-blind | **Intervention:** (1) V. officinalis 45%MeOH extract 500 mg; (2) V. officinalis 45%MeOH extract 500 mg and Humulus lupulus 45%MeOH extract 120 mg (fixed combination Ze91019); (3) placebo. For 4 weeks. **Measures:** (1) the reduction of the initially prolonged sleep latency; (2) WASO, sleep efficiency, relative proportion of the sleep stages, REM latency and CGI | The fixed combination was significantly superior to placebo in reducing the sleep latency, whereas the single valerian extract did not differ significantly from placebo. | No adverse events. | H. lupulus | 3 |
| Müller SF   | Observational study | **Intervention:** V. officinalis 62%EtOH extract 160 mg and Melissa officinalis 30%EtOH extract 80 mg per tablet (Euvégal forte, Schwabe Pharmaceuticals, Germany). Max. 2×2 tablets/day (average: 3.5 tablets/day) for 4 ± 1 weeks (average: 31.9 days). **Measures:** Severity of symptoms and improvement efficacy. Scored by the investigators and the parents. | Dyssomnia and restlessness were reduced from moderate/severe to mild or absent in most of the patients. (80.9% of dyssomnia patients and 70.4% of restlessness patients improved.) Both parents and investigators assessed efficacy to be very good or good (60.5% and 67.7%). Tolerability was good or very good in 96.7% of the patients. | No adverse events. | M. officinalis | n/a |
| Kennedy DO  | Randomized, placebo-controlled, double-blind, cross-over | **Intervention:** 600 mg, 1200 mg, and 1800 mg of a standardized V. officinalis and Melissa officinalis extracts (V. officinalis: M. officinalis 3: 2). Three single doses, separated by a 7 day wash out period. **Measures:** (1) Modulation of mood and anxiety (Bond-Lader scales, STAI) after DISS battery exposure; (2) DISS performance | The 600 mg dose ameliorated the negative effects of DISS on anxiety, however 1800 mg increased anxiety. All 3 doses led to decrements in performance on the Stroop task module, and the 2 lower doses led to decrements on the overall score generated on DISS. | M. officinalis | 3 |
| First Author | Study design | Intervention and measures | Outcomes | Adverse events | Combined with | Jadad scale |
|--------------|--------------|----------------------------|----------|---------------|---------------|-------------|
| 14 Morin CM 2005⁷⁹ | Randomized, placebo-controlled | Population: Mild insomnia patients (n=184) | Intervention: (1) Combination of V. officinalis (45%MeOH extract) 187 mg and Humulus lupulus (45%MeOH extract) 41.9 mg, nightly for 28 days; (2) placebo for 28 days; (3) diphenhydramine 25 mg for 14 days followed by placebo for 14 days. Measures: Sleep parameters measured by daily diaries and polysomnography, clinical outcome ratings from patients and physicians, and quality of life. | Modest improvements of subjective sleep parameters in valerian-hops (1) and diphenhydramine (3). Valerian-hops produced slightly greater reductions of sleep latency relative to placebo and diphenhydramine after 14 days, and greater reductions than placebo after 28 days. No group difference on polysomnography. Quality of life was significantly more improved in valerian-hops group than the placebo group after 28 days. | No serious adverse events | H. lupulus 3 |
| 15 Schellenberg R 2004⁸⁰ | Randomized, placebo-controlled | Population: Healthy volunteers (n=32) | Intervention: Single doses of V. officinalis / Humulus lupulus 45%MeOH extract combination (500 mg / 120 mg, or 1500 mg / 360 mg). Valerian extract contained 0.388% if valerenic acids. Measures: competition between caffeine and valerian/hop on the central nervous system (EEG) | Valerian 500mg/hops 120 mg reduced, and valerian 1500mg/hops 360 mg inhibited caffeine-induced arousal, 60 min after oral administration. | Not mentioned | H. lupulus 3 |
| 16 Farag NH 2003⁸¹ | Randomized, placebo-controlled, double-blind, cross-over | Population: Sleep onset insomnia (n=25) | Intervention: Combination of V. wallichii 320 mg, Rosa centifolia, Nardostachys jatamansi, Tinospora cordifolia, Withania somnifera, Piper nigrum, Zingiber officinalis, Convolvulus pluricaulis, and Glycyrrhiza glabra. Compared to placebo. 4 nights each with 10 days washout. Measures: Sleep latency (questionnaire) | The supplement reduced sleep latency. The effect was greater in subjects with worse insomnia. | None | Ayurvedic herbs 4 |
| 17 Müller D 2003⁸⁵ | Observational | Population: Patients suffering from mild to moderately severe depression (n=2,462) | Intervention: (1) V. officinalis 70%EtOH extract (Euvégal Balance) 500 mg or 1000 mg and Hypericum perforatum 60%EtOH extract (Neuroplant) 600 mg per day. For 6 weeks. Measures: Rating of 16 symptoms during each doctor’s visit (ICD-10 and HAM-A). | Core symptoms of depression and the main symptoms of anxiety disorder receded significantly. Efficacy was ‘very good’ or ‘good’ in 87.2%, and ‘poor’ in 1.6%. Tolerability was ‘very good’ or ‘good’ in 96.0%. Higher Valerian dosage could be more effective. | No significant side effects | H. perforatum n/a |

(continued)
| First Author | Study design | Intervention and measures | Outcomes | Adverse events | Combined with | Jadad scale |
|--------------|--------------|---------------------------|----------|---------------|---------------|-------------|
| Vonderheid-Guth B 2000 | Randomized, placebo-controlled, single-blind, cross-over | Population: Young healthy male volunteers (n=12) | Intervention: Single dosages of *V. officinalis* and *H. lupulus* 45%MeOH extract combination (Ze91019). Low dosage: Valerian 500 mg and hops 120 mg. High dosage: valerian 1500 mg and hops 360 mg. Measures: Quantitative topographical EEG (qEEG). Performance Test (CPT) according to Duerker and Lienert (1965). Prior to, 1, 2, and 4 hours after drug intake. | qEEG: High dosage led to power increases in delta, decreases in alpha and beta. CPT: Concentration and performance capability were hardly influenced. (A minimal increase of mean answer time and mean time for correct answer was observed after 4 h after high dosage intake and 1 h after low dosage intake. – More pronounced after the low dosage.) | | |
| Lindahl O 1989 | Randomized, placebo-controlled, double-blind, cross-over | Population: Volunteers with sleep difficulties and fatigue. (n=27) | Intervention: Single doses of (1) test preparation: *V. officinalis* extract (equivalent to 400 mg of the root), *H. lupulus* extract (375 mg equivalent) and *M. officinalis* extract (160 mg equivalent) or (2) Control: same amounts of hops and lemon balm and *V. officinalis* 4 mg equivalent. Measures: Subjective rating of sleep quality | 78% rated high valerian preparation better than the control. | No side effects. | |

Abbreviations: BDI-Beck Depression Inventory, DIS-Defined Intensity Stress Simulator, DSST-Digit Symbol Substitution Test, DST-Digit Span Test, EEG-Electroencephalogram, HAM-A-Hamilton Anxiety Rating Scale, ICD-10-International Classification of Diseases 10th Revision, ISI-Insomnia Severity Index, PSQI-Pittsburgh Sleep Quality Index, TSST-, WASO-Wake Time After Sleep Onset
heterogeneity was observed ($I^2 = 85.33\%$). As described above, differences in herbal preparation could potentially contribute to the heterogenous outcomes. In fact, subgroup analysis for the whole root$^{30,35,56,57}$ (RCTs [Jadad 3-5]) and the extract$^{34,36,51-54}$ (RCTs [Jadad 2-5]) revealed that the combined effect size for the whole root was considerably higher (0.83 (95\% CI: 0.03 to 1.62), compared to the extract (0.10; 95\% CI: -0.02 to 0.22) (Figure 3), further supporting the use of the whole root rather than extract.

**Anxiety.** Anxiety was assessed by Hamilton Anxiety Rating Scale (HAM-A): a clinician-rated scale that considers psychological and somatic factors,$^{93,94}$ State-Trait Anxiety Inventory (STAI): a patient-rated scale that measures state and trait anxiety,$^{95,96}$ mental stress test using the color/word interference task,$^{97}$ or other questionnaires. Seven studies examined valerian’s anxiolytic potential, and positive outcomes were observed in 6 studies.$^{33,35,43,60,63,71}$ Valerian standardized extract 600 mg per day for 1 week reduced psychological and physiological stress reactivity under stress in healthy adult subjects$^{43}$ (RCT [Jadad scale 1]). Average 81.3 mg per day of valepotriates for 4 weeks reduced the score of HAM-A in generalized anxiety disorder patients$^{31}$ (RCT [Jadad scale 5]), suggesting that valepotriates are the active anxiolytic constituents. Valerian root / rhizome 530 mg per day for 4 weeks reduced anxiety in HIV-positive patients receiving efavirenz$^{35}$ (RCT [Jadad scale 3]), suggesting valerian could prevent neuropsychiatric adverse effects caused by the antiretroviral medication. Valerian extract 100 mg$^{33}$ or root / rhizome 100 mg, 60 single dose 60 min before the initiation of surgical operation, reduced anxiety in anxious patients undergoing dental operation$^{33,60}$ (RCTs [Jadad scale 5]). In addition, valerian extract 1,260 mg per day for 7 days for 3 menstrual cycles effectively reduced premenstrual anxiety$^{63}$ (RCT [Jadad scale 5]). However, an internet-based study with volunteers with non-clinical anxiety and insomnia found no significant difference in STAI compared to placebo$^{51}$ (RCT [Jadad scale 5]). Figure 4 shows the result of meta-analysis for valerian’s anxiolytic effects based on 7 randomized placebo-controlled trials that reported quantitative outcomes$^{34,35,47,49,51,60,63}$ (Figure 4A). The funnel plot indicated significant publication bias, missing data in the area of negative outcome, and high heterogeneity ($I^2 = 93.13\%$) (Figure 4B). Subgroup analysis revealed high variability for valerian extracts$^{34,47,49,51,63}$ (RCTs [Jadad scale 3-5]), whereas positive outcomes were observed for the whole root$^{35,60}$ (RCTs [Jadad scale 3 and 5]). However, due to the scarcity of data (2 studies for the whole root), it is difficult to draw a conclusion.

Two studies investigated the effects of *V. officinalis* extracts on anxiety-associated brain activities.$^{31,34}$ Single dose of 300 mg extract reduced intracortical facilitation, possibly via GABA_A enhancement$^{31}$ (RCT [Jadad scale 4]), and hydroalcoholic extract 300 mg per day for 4 weeks increased frontal brain region alpha coherence, which was correlated with anxiolysis$^{34}$ (RCT [Jadad scale 5]). These findings suggest that *V. officinalis* could induce immediate as well as long-lasting effects on anxiety-associated brain activities.

**Other therapeutic effects.** In addition to insomnia and anxiety, several other therapeutic benefits were suggested (Figure 5). Valerian aqueous extract 750 mg per day for 8 weeks was
effective in reducing the symptoms of obsessive-compulsive disorder (OCD)\(^{55}\) (RCT [Jadad scale 2]). Valerian root / rhizome 530 mg per day for 1 month improved cognition in haemodialysis patients\(^{32}\) (RCT [Jadad scale 2]), and valerian root / rhizome 1,060 mg per day for 8 weeks prevented cognitive dysfunction after coronary bypass surgery\(^{61}\) (RCT [Jadad scale 5]). Valerian root / rhizome 675 mg or extract 1,060 mg per day for 8 weeks reduced severity and frequency of hot flashes in menopausal and postmenopausal women\(^{59,64}\) (RCTs [Jadad scale 4]). In addition, valerian root / rhizome 765 mg or extract 1,060 mg per day for 3 days for 2 cycles reduced pain severity and syncope in young females with dysmenorrhea\(^{56}\) (RCT [Jadad scale 5]), and valerian extract 1,260 mg for 7 days for 3 cycles reduced premenstrual symptoms\(^{63}\), supporting the traditional use of valerian as a menstrual stimulant.\(^{16}\)

Other Valerian Species

Three studies investigated the effectiveness of other *Valeriana* spp. as a sleep aid: 2 studies on *V. edulis\(^{42,46}\) and 1 study on *V. wallichii\(^{70}\). *V. edulis* 20 mg/kg for 2 weeks reduced sleep latencies and improved sleep quality in children with intellectual deficit and primary sleep problems\(^{46}\) (RCT [Jadad scale 5]), and *V. edulis* hydroalcoholic extract 450 mg single dose increased REM sleep and reduced the number of awakenings in insomnia patients\(^{42}\) (RCT [Jadad scale 5]). A concoction of

\[\text{**Figure 2.** Sleep quality improvement by repeated administration of *V. officinalis*. Ten studies were included in meta-analysis. Positive values indicate enhanced sleep quality. (A) Forest plot for sleep quality improvement. Effect sizes (Hedges' g) for 10 studies (blue circles) and combined effect size (green circle) are shown. Black horizontal bars indicate 95% CI. (B) Funnel plot. Included data points are presented as filled blue circles and imputed data are shown as open orange circles.} \]
Valepotriates from *V. wallichii* 300 mg per day for 15 days reduced Wake Time After Sleep Onset (WASO) and improved sleep quality in insomnia patients (RCT [Jadad scale 3]). Of note, chemical characteristics of the 3 species (*V. officinalis*, *V. edulis*, and *V. wallichii*) significantly differ; valerenic acids (valeric acid, acetoxyvalerenic acid, hydroxyvalerenic acid) are specific to *V. officinalis*, whereas valepotriates (valtrate and isovaltrate) are common among the 3 species, suggesting that valepotriates at least in part contribute to valerian’s sleep promoting activity. Due to the scarcity of data, it was not possible to perform meta-analysis for *V. edulis* and *V. wallichii*.

**Valerian in Herbal Combinations**

In addition to the effectiveness of valerian alone, 19 studies evaluated the effectiveness of valerian in combination with other herbs or other agents, among which 16 studies used *V. officinalis*, 2 studies used *V. wallichii*, and 1 study used an unknown valerian species. One study treated menopausal women with valerian essence in combination with lemon balm and observed sleep improvement, however the species and extraction methods were unidentified. Among 16 reports using herbal combinations with *V. officinalis*, 8 studies examined the effects on sleep quality and/or latency and all those interventions led to positive outcomes. Six studies evaluated anxiolytic potential, and all those studies found positive outcomes at least for one of the measures at one of the tested dosages (RCTs [Jadad scale 3-5] and 3 observational studies). However, a study using the combination of *V. officinalis*, *Passiflora incarnata* (syn. *P. edulis*), and *Melissa officinalis* extracts (single dose) found that the intervention attenuated anxiety under stress, although it did not alter the level of salivary cortisol, a physiological parameter of stress reactions (RCT [Jadad scale 5]). In another study, *V. officinalis* and *M. officinalis* combination (extract weight ratio 3:2) at 1,800 mg (single dose) increased anxiety while the same formulation at 600 mg reduced anxiety (RCT [Jadad scale 3]). Both of these negative outcomes were observed for single doses in healthy subjects. On the other hand, the other 4 studies, conducted with subjects with adjustment insomnia, somatoform disorder patients (RCT [Jadad scale 4]), depression patients (observational), and children with affective disorders (observational), found that the herbal combinations with *V. officinalis* were effective in reducing anxiety. Of note, when depression patients were treated with *V. officinalis* (500 or 1,000 mg extract) and *Hypericum perforatum* (600 mg extract) for 6 weeks, higher valerian dose (1,000 mg) reduced anxiety more effectively (observational). In addition, a study found that a combination of *V. officinalis* and *M. officinalis* extracts was effective in reducing the symptoms in children with hyperactivity and concentration difficulties (observational). In a study with Intensive Care Unit (ICU) patients, *V. officinalis* essential oil combined with acupuncture effectively induced relaxation and improved sleep (observational). *V. wallichii* with 8 ayurvedic herbs improved sleep in sleep onset insomnia patients (RCT [Jadad scale 4]), and *V. wallichii* with milk improved sleep in primary insomnia patients (observational).
With respect to the potential herbal partners of *V. officinalis*, *Humulus lupulus* was the most frequently used in the clinical trials. All 7 reports that examined the effects on sleep or associated parameters observed sleep improvement\(^6,5,72,79,83\) (RCTs [Jadad scale 2-3]), EEG power alteration toward sleep\(^82\) (RCT [Jadad scale 2]), and reduced arousal\(^80\) (RCT [Jadad scale 3]) were observed. *M. officinalis* used in 5 trials\(^73,77,78,84,89\) was the second most frequently combined herb, followed by *P. incarnata* used in 4 trials.\(^73,75,77,86\) The 2 herbs were used in combination with *V. officinalis* for sleep problems and anxiety, resulting in positive outcomes.\(^73,75,77,78,84,86,89\) Two studies were conducted on the combination of *V. officinalis* and *H. perforatum* with depression patients\(^85\) (observational) and children with affective disorders\(^86\) (observational), and the
intervention reduced anxiety, improved sleep, and reduced depression symptoms. Two studies used *M. officinalis* and *P. incarnata* (RCTs [Jadad scale 4-5]), and 1 study used *H. lupulus* and *P. incarnata* (RCT [Jadad scale 4]) in combination with *V. officinalis*. These 4 frequently used herbal partners and the study outcomes are summarized in Figure 6. It was not possible to perform meta-analysis for herbal combinations due to the highly variable outcome measures.

**Safety**

There were no serious adverse events reported in the studies included in this review (60 studies, total n=6,894). Paradoxical stimulation, such as agitation and restlessness, was experienced only in minority. In older women with insomnia, valerian extract increased WASO, suggesting it could have stimulatory effects. However, a study found that the same formulation at a higher dose increased anxiety while a lower dose was anxiolytic, suggesting that paradoxical stimulation could be avoided by cautious dosing. Six studies investigated possible adverse effects on cognitive performance after *V. officinalis* extract intake (single dose) in healthy adults and elderly volunteers. *V. officinalis* standardized extracts at 100-1,600 mg did not impair cognitive or psychomotor performance and were proven safer compared to triazolam, temazepam (benzodiazepines), and diphenhydramine (antihistamine). Valerian did not cause any adverse events in postmenopausal women with insomnia, older women with insomnia, children with intellectual deficits and primary sleep problems, and psychophysiological insomnia patients. Mild adverse events were reported in RLS patients (vivid dreams and fatigue), arthritis patients with sleep disturbance (dizziness and sleepiness), sleep-disturbed subjects (drowsiness), insomnia patients (gastrointestinal symptoms), and outpatients with stress-induced insomnia (vivid dreams, drowsiness, heavy dream and depression), however there was no clear association with the treatments. Potential herb-drug interaction was addressed by examining the effects on cytochrome p450 (CYP) expression levels, and no significant impact was detected at least for CYP1A2, CYP2D6, CYP2E1 and CYP3A4/5. As a whole, valerian is safe in all ages.

**Discussion**

Concurring sleep problems and anxiety are frequently observed and their onset and course are interrelated. In addition, these disorders are associated with a number of comorbid conditions including depression, dementia, OCD, and hot flashes (Figure 6). Our results demonstrated that sleep promotion and anxiolytic effects were the major therapeutic benefits expected of valerian (*V. officinalis*), and this herb could be also useful in treating OCD, cognitive dysfunction, menopausal hot flashes, as well as menstrual problems. However, the study outcomes considerably differed particularly when herbal extracts were used. We address this issue by discussing potential mechanisms underlying the therapeutic actions of valerian.
Key Constituents and Mechanisms of Actions

Although the use of valerian extracts as a sedative and sleep aid dates back to the 18th century and a number of constituents have been identified in the last 120 years, the active constituents and underlying mechanisms for the reported activities remain obscure. The identified constituents include iridoids known as valepotriates (valtrate, isovaltrate, didrovaltrate and acevaltrate), essential oil constituents including monoterpenes (e.g. borneol, bornyl acetate), sesquiterpenes (e.g. valerenal, valerenic acid), and carboxylic compounds (valeric / isovaleric acid), lignans, flavonoids, and low levels of γ-aminobutyric acid (GABA).

Valeriana edulis (Mexican valerian) and Valeriana wallichii (Indian valerian) contain higher levels of iridoids, while valerenic acid and acetoxylvalerenic acid are unique to V. officinalis.

Valerian and GABAergic Signaling

GABA is an inhibitory neurotransmitter within the central nervous system and is a key target of pharmacotherapies in the treatment of anxiety and sleep disorders. GABA acts via 3 subclasses of receptors termed GABA\textsubscript{A}, GABA\textsubscript{B}, and GABA\textsubscript{C}, each of which has distinct characteristics. GABA\textsubscript{A} and GABA\textsubscript{C} receptors are ligand-gated ion channels, while GABA\textsubscript{B} members are G-protein coupled receptors. GABA\textsubscript{A} receptors are heteropentameric transmembrane protein complexes made up of α1-6, β1-3, γ1-3, δ, ε, θ, π subunits, and have numerous allosteric binding sites. Although it is not very likely that GABA in valerian, taken orally, directly exerts any therapeutic action, valerenic acid and valerenol can allosterically modulate GABA\textsubscript{A} receptors to enhance the response of GABA\textsubscript{A} to GABA in vitro, and both valerenic acid and valerenol exerted anxiolytic activity in a GABA\textsubscript{A} β3 subunit-dependent manner in mice. On the other hand, valerenic acid derivatives (e.g. acetoxy valerenic acid and hydroxy valerenic acid) bind the identical sites without inducing allosteric modulation, suggesting that these constituents can potentially compete with valerenic acid. In fact, in an anxiety model in rodents, a valerian extract containing valerenic acid at a high level and acetoxy valerenic acid at a low level exhibited higher anxiolytic potency compared to an extract with acetoxy valerenic acid at a high level, indicating that standardization with respect to the total amount of valerenic acids (both valerenic acid and acetoxy valerenic acid) is inappropriate. In addition, an aqueous extract of valerian induced GABA release in rat brain synaptosomes in vitro, suggesting the presence of hydrophilic constituents potentially promotes GABAergic signaling. Expected mechanisms of actions are summarized in Figure 5. Of note, the expression patterns of GABA receptors in humans are sex- and age-dependent, and GABA\textsubscript{A} β3 subunit levels in the superior temporal gyrus were lower in older women than old men, suggesting that valerian’s actions via GABAergic signaling could depend on the sex and age.

Valerian and Serotonergic Signaling

Serotonin (5-HT) plays essential roles in the regulation of sleep and mood, and serotonergic system is a promising therapeutic target for psychiatric disorders including anxiety, depression, and sleep disorders. Both valerian extract and valerenic acid exhibit partial agonist activity at 5-HT receptor 5A (5-HT\textsubscript{5A}) in vitro, suggesting the involvement of serotonergic system in the actions of valerian. There are 7 families of its receptors (5-HT\textsubscript{1-7}). Although the functional significance of 5-HT\textsubscript{5A} remains obscure, 5-HT\textsubscript{5A} is present at high levels in the regions that regulate circadian rhythms (the suprachiasmatic nucleus, the intergeniculate leaflet, the median raphe...
nuclei, and dorsal raphe nucleus) and colocalized with serotonin, suggesting the auto-receptor role of 5-HT2A in circadian regulation. In addition, 5-HT2A is expressed in the brain areas such as the cerebral cortex, hippocampus, amygdala and hypothalamus, suggesting that it is potentially involved in stress reactivity and resilience. Indeed, a study demonstrated that 5-HT2A antagonists could have different sedative, anxiolytic, and anti-depressant properties (Figure 6). These data suggest that valerian, as a 5-HT2A partial agonist, may act differently depending on what circumstances the subjects were placed and this could possibly explain the inconsistent outcomes among the trials.

Valerian and Adenosine Signaling

Adenosine receptors play important roles in mood and anxiety disorders. Adenosine A1 receptor activation reduces anxiety-like behavior in mice and partial allosteric modulators of adenosine A1 receptor could be potential anxiolytic therapy. In addition, adenosine helps maintain brain homeostasis by preventing over excitation and regulating the sleep-wake rhythm. Sleep stages are divided into 1 stage of rapid eye movement (REM) sleep and 4 stages of non-rapid eye movement (NREM) sleep (Stages 1-4) that are characterized by increasing sleep depth. The deeper stages (Stages 3 and 4) are believed to be the most restorative and collectively called SWS. Adenosine acting through A1 receptors facilitates sleep and increases slow wave activity (SWA) in SWS, whereas A1 receptor antagonism suppresses SWA. Importantly, 2 studies found that *V. officinalis* could increase SWS, possibly contributing to the improved subjective sleep quality while objective sleep latency remained unchanged. These observations suggest that valerian may promote the sleep quality by inducing deep sleep via adenosine A1 receptor signaling. In fact, a polar extract of valerian exhibited partial agonistic activity at A1 receptors and valerian contains hydrophilic constituents that interact with adenosine signaling, such as lignans of a olivil derivatives that is a partial agonist of adenosine A1 receptors. Thus, valerian’s hydrophilic constituents could exert the anxiolytic and sleep-inducing effects via adenosine signaling. Importantly, while the hydrophilic extract was agonistic, isovaltrate in the hydrophobic extract acted as an inverse agonist at adenosine A1 receptor. These findings indicate the hydrophilic and hydrophobic components of valerian act in opposite directions at adenosine receptors, supporting the traditional use of water-based extract as a sleep aid, and possibly explaining the contradictory findings regarding the efficacy of valerian hydroalcoholic extracts.

Valepotriates

Valepotriates are unstable thermolabile compounds that rapidly decompose. The characteristic odor of dried valerian is due to isovaleric acid released from decomposing valepotriates. Although the mechanistic roles of valepotriates in the therapeutic effects are yet to be clarified, pre-clinical evidence suggests that valepotriates exert anxiolytic and antidepressant-like effects, possibly via the dopaminergic and noradrenergic neurotransmission but not via the serotonergic system. The monoaminergic neurotransmitters (dopamine, noradrenaline, and 5-HT) play overlapping roles in the etiology of anxiety and depression, and medications targeting these systems are proved to be effective in treating both anxiety and depression. Moreover, those monoamines play a role in sleep-wake regulation. Thus, the presence of valepotriates is likely to be essential in treating these disorders. In addition, valtrate reduced the serum glucocorticoid level in a rat anxiety model, suggesting hypothalamus-pituitary-adrenal axis modulatory potential of valepotriates. Considering the significant association between neuroendocrine-immune axis imbalance and stress vulnerability and resilience, the potential impact on the hypothalamus-pituitary-adrenal axis is particularly important. Stress can disturb the sleep, and stress reactivity is implicated in vulnerability to insomnia and other stress-related disorders such as anxiety and depression. Furthermore, glucocorticoid imbalance could lead to amyloid formation and tau accumulation, hallmarks of Alzheimer’s disease, and elevated glucocorticoid levels are associated with neurodegenerative disorders. Thus, valepotriates may prevent stress-induced neuropsychiatric disorders including anxiety and depression, as well as cognitive dysfunction, by restoring neuroendocrine-immune balance. Expected mechanisms of action are summarized in Figure 6.

Quality and Shelf Life

As described above, multiple constituents, including valerenic acids, lignan, and valepotriates, differentially contribute to the therapeutic effectiveness of valerian. However, most of the commercially available valerian extracts are standardized to the levels of valerenic acids, while the contents of valepotriates are often unknown. Importantly, Bos et al. compared freshly prepared tincture (70% EtOH extract) and the same extract stored at room temperature for 2 months, and found that the levels of valepotriates (valtrate and isovaltrate) after 2 months were less than 5% of the original, whereas the levels of valerenic acids (valerenic acid and acetoxyvalerenic acid) were not affected. This finding suggests that the shelf life of valerian extracts may be significantly shorter if the levels of valepotriates are considered. It is possible that the negative outcomes observed in the trials using valerian extracts, rather than the raw materials, could have been caused by the loss of valepotriates in those extracts.

In the United States, valerian-containing products are widely available as dietary supplements, in various forms under various trade names. Examples are Valerian (NOW FOODS, IL, US; Nature’s Way, WI, USA), Standardized Valerian (root extract standardized to 0.8% valerenic acids; Nature’s Way), and Amantilla (hydroalcoholic extract; NutraMedix, FL, USA), to name a few. In Europe, the Committee on Herbal Medicine Products (HMPC) concluded that valerian-containing...
medicinal products have to be assessed by the responsible national authorities. Available products include Baldrian (85% ethanol extract; Kneipp, Wu¨rzburg, Germany), and Baldrianwurzel (70% ethanol extract; Hofapotheke St. Afra, Augsburg, Germany). Herbal combinations are also available (e.g. Valeirana-Melissa extract [standardized to 0.8% valerenic acids; Bonusan, Rotterdam, Netherlands]). Of note, the recommended storage temperatures for most of these products are at or below room temperature (around 25°C).

Limitation
The current study has reviewed the effectiveness of valerian without specifying target populations. This raises a limitation as the etiology underlying sleep problems and associated health problems could be diverse depending on age, sex, and general health conditions. Consequently, there could be potential differences in the outcomes depending on the target populations, however such possibilities were not addressed in this review. In addition, heterogeneity in the outcome measures limited the number of data sources for meta-analysis, which in turn limits the generalizability of the results. Further research is needed to assess the effectiveness for specific populations and outcomes.

Conclusion
According to the 1905 edition of King’s American dispensatory,99 ‘valerian excites the cerebro-spinal system.’ and ‘in medicinal doses it acts as a stimulant-tonic,...’
suggests that valerian is not a simple hypnotic or anxiolytic agent. In addition, the text continues that ‘The extract of valerian is worthless, but the fluid extract has been found to possess all the medicinal virtues of the root’, suggesting that extraction methods could play a crucial role in preserving the therapeutic potency. Valerian has diverse chemical properties. As summarized in Table 3, the differences in polarity can lead to variability in the quality of valerian extracts depending on the extraction solvents. Our study has found that the hydroalcoholic extracts used in the clinical trials are variable with respect to the solvents (Table 4), and their phytochemical properties are not characterized in most cases. Furthermore, there is no information regarding the storage conditions such as temperature and storage period. Considering the low stability of some of the important active constituents, it is unknown to what extent the active constituents had been preserved at the end of those trials. The absence of such information limits the discussion as to why some extracts were ineffective while others exhibited effectiveness in those clinical trials. Due to the multiple active constituents and complex mechanisms of actions (Figure 5), it is imperative to use well characterized plant materials and extracts in the future clinical trials, in order for the evaluation of valerian’s true effectiveness to be possible.

This study demonstrated that valerian could be a safe and useful herb alone and also in combination in treating sleep problems, anxiety, and associated comorbidities. One possibility for exploration that arises is whether valerian is particularly useful in treating insomnia where there are higher levels of anxiety. In fact, the observation that *V. edulis* reduced sleep problems most effectively in children with hyperactivity may suggest that valerian is more useful in treating insomnia associated with particular psychiatric conditions such as anxiety. The safety has been proven in a wide range from childhood to old age. Sleep problems are associated with a number of comorbidities including anxiety, depression, and dementia in both adulthood and childhood. As we age, the quantity and quality of sleep decrease and insomnia and daytime drowsiness are frequently reported in the older population. Thus, valerian may help improve the quality of life in all ages by improving sleep quality, thereby preventing a number of psychiatric and cognitive dysfunctions. However, due to the presence of multiple active constituents and the unstable nature of some components, it may be useful to revise the standardization methods and shelf life of the extracts. Repeated treatments with the whole root / rhizome consistently promoted sleep quality at 450-1410 mg per day for 4-8 weeks, whereas valerian extracts 300-600 mg per day for 5 days-4 weeks resulted in inconsistent outcomes. Evidence is poor with respect to the effectiveness of single dose intervention as a sleep aid. Developing effective standardization methods would be desirable. Meanwhile, the usage of whole herbal substances (root / rhizome), rather than extracts, may be the way to obtain optimal efficacy for the time being.

Lastly, due to the subjective nature of sleep problems and associated mood disorders, there remains a difficulty in determining the best measure for the efficacy of interventions. The ultimate goal of any therapies, whether it be traditional, conventional or integrative, beyond disease ‘cure’ or modulation, is to improve the quality of life for patients. While objective measures in a laboratory setting provide valuable information and mechanistic understanding, it may be useful to consider additional and more important factors that need to be addressed specifically in each individual. Such measures and tailor-made approaches would necessarily involve sufficient interaction and communication between patients and practitioners. Developing effective trial strategies to address this issue is a future challenge.

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**Author Contributions**
NS wrote this systematic review with input and contributions of GW and JG. JS, GW and JG planned the search in databases and defined criteria for the selection of articles. NS carried out the search, screen the materials and proposed a selection. All authors contributed to the evaluation of the studies. All authors contributed to the interpretation of data and discussion. All authors reviewed the whole manuscript, read and approved the submitted version.

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