No Interacting Influence of Lavender Oil Preparation Silexan on Oral Contraception Using an Ethinyl Estradiol/Levonorgestrel Combination

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Abstract

Purpose  Silexan is an oral Lavender oil preparation with proven anxiolytic efficacy. Given the high prevalence of anxiety and restlessness in younger women, oral contraceptives and Silexan will likely be co-administered.

Methods  A double-blind, randomised, 2-period crossover study was performed to investigate the effects of Silexan on the pharmacokinetics and pharmacodynamics of Microgynon®, a combination oral contraceptive containing ethinyl estradiol 0.03 mg (EE) and levonorgestrel 0.15 mg (LNG) in healthy, fertile, adult females. During 2 consecutive cycles of 28 days, oral contraception was given for 21 days combined with 1 × 160 mg/day Silexan or placebo. Plasma concentration–time profiles of EE and LNG were obtained on day 18 ± 1 up to 24 h after dosing. The primary outcome measure was the area under the concentration–time curve over a dosing interval of $\tau = 24$ h (AUC$\tau$) for EE and LNG plasma levels. An interaction with Silexan was formally excluded if the 90 % confidence interval for the AUC$\tau$ ratio during co-administration with Silexan or placebo was included within the range of 0.80–1.25. Secondary outcomes included EE and LNG peak concentration ($C_{max}$) and time to $C_{max}$ ($t_{max}$), follicle size, endometrial thickness, the Hoo gland score, and serum levels of estradiol, progesterone, and sex hormone-binding globulin.

Results  A total of 24 women (mean age 27.3 years; mean body mass index 22.2 kg/m$^2$) participated. The confidence intervals for the EE and LNG AUC$\tau$ and $C_{max}$ ratios fell within the pre-specified limits, indicating no interaction (point estimates [Silexan/placebo] AUC$\tau$ EE 0.97, LNG 0.94; $C_{max}$ EE 0.99, LNG 0.96). For LNG, $t_{max}$ was slightly delayed. No secondary outcome indicated any impairment of contraceptive efficacy.

Conclusions  Co-administration of Silexan did not affect the efficacy of a combination oral contraceptive containing EE and LNG and was well tolerated.

Trial registration: ISRCTN52706881, International Standard Randomised Controlled Trial Number Register.

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Key Points

Due to the high prevalence of anxiety and restlessness in younger women, the herbal anxiolytic drug Silexan and oral contraceptives are likely to be co-administered. Therefore, it is important to investigate the interaction potential of Silexan with oral contraceptives.

This randomised, double-blind, placebo-controlled crossover trial showed that Silexan did not relevantly affect the pharmacokinetic properties of a combination oral contraceptive containing ethinyl estradiol and levonorgestrel.

Silexan therefore does not decrease the efficacy of currently used combination oral contraceptives.
1 Introduction

Silexan\(^1\) is a preparation from \textit{Lavandula angustifolia} for oral use that is obtained from the flowering tops of the plant by steam distillation, and that complies with the monograph Lavender oil of the European Pharmacopoeia [1] with respect to all quality parameters. It exceeds the quality definition of the pharmacopoeial monograph with respect to items that are important for efficacy and tolerability. Randomised controlled clinical trials have demonstrated that Silexan is a potent anxiolytic drug with superior efficacy compared with placebo in subsyndromal anxiety disorder [2]. In generalised anxiety disorder (GAD), Silexan was found to be comparably efficacious as lorazepam [3] and at least as efficacious as paroxetine. The drug has a marketing authorisation in Germany for the symptomatic treatment of restlessness related to anxious mood, with a recommended dosage of 80 mg/d. While Silexan is a complex, multi-ingredient mixture, its anxiolytic properties have been ascribed in particular to linalool and linalyl acetate [4].

Anxiety disorders are by far the most prevalent psychiatric conditions in Western Europe and the USA [5, 6]. Their prevalence in women is at least twice as high as in men [7, 8]. Since the prevalence of anxiety disorders tends to decrease rather than to increase with age [9], a substantial proportion of the patients who seek treatment for anxiety are women of childbearing potential who may be practising oral contraception.

The steroid hormones contained in oral contraceptives are metabolised mainly through hydroxylation in the liver. Drugs that induce the cytochrome 
P450 (CYP) enzyme system, notably CYP3A4, reduce the bioavailability by increasing the first-pass metabolism of the oral contraceptive steroids and may therefore impair their efficacy, leading to breakthrough bleeding and, possibly, unwanted pregnancy [10, 11]. A comprehensive list of drugs that induce CYP has been published by Rendic [12]. They include several anticonvulsants and broad spectrum antibiotics, rifampicin, griseofulvin, and anti-retroviral drugs.

Bioavailability is the rate and extent to which the active substance or active moiety is absorbed from a pharmaceutical form and becomes available at the site of action [13]. As the use of herbal preparations increases in the general population, the potential for herb–drug interactions also increases. Therefore, herbal products must be assessed for their efficacy and safety, including their potential for interactions with other drugs [14]. Since women of child-bearing potential who practise oral contraception represent an important part of the target population for which Silexan is intended, it is important to investigate whether the herbal drug may have an effect on the bioavailability and efficacy of oral contraceptives. Although a clinical study showed no induction of the CYP enzymes 1A2, 2C9, 2C19, 2D6 and 3A4 [15], suggesting that no increased activity in metabolism of contraceptive ingredients occurs during co-administration of Silexan, some examples of inhibition of phase-II enzymes, mainly UDP-glucuronosyltransferases (UGTs), by co-administered herbal drugs (e.g., Williams et al. [16]) do exist. This study was therefore performed to investigate the relative bioavailability of ethinyl estradiol (EE) and levonorgestrel (LNG), the active ingredients of the monophasic combination oral contraceptive Microgynon\(^{\text{®}}\), during co-administration of Silexan as compared with the administration of the oral contraceptive alone, as well as any attenuating effect on contraceptive efficacy.

2 Experimental Procedures

2.1 Protocol and Design, Objectives

The investigation was performed as a double-blind, randomised, placebo-controlled, monocentric, crossover study in healthy volunteers whose primary objective was to assess the interacting potential of Silexan with the pharmacokinetic properties of EE and LNG, the active ingredients of the combination oral contraceptive Microgynon\(^{\text{®}}\). An important secondary objective was to assess whether Silexan had an effect on the suppressive efficacy of Microgynon\(^{\text{®}}\) on hormonal and ovarian status.

The protocol was reviewed and approved by an independent ethics committee. All subjects provided written informed consent. The principles of Good Clinical Practice and the Declaration of Helsinki were adhered to.

2.2 Participants

The study participants were healthy, female, premenopausal volunteers aged between 18 and 38 years (smokers had to be ≤30 years of age, and those with a consumption of more than ten cigarettes per day were excluded), who presented with a body mass index between 18 and 30 kg/m\(^2\). For eligibility for randomised treatment, subjects had to take Microgynon\(^{\text{®}}\) for at least 2 cycles before randomisation, with a ‘usual’ 28-day pill cycle and withdrawal bleedings during the 7-day pill-free interval. Eligible subjects were also required to use a non-hormonal method of contraception in addition to Microgynon\(^{\text{®}}\) throughout the study period. Women with delivery, abortion, or lactation within 3 months of the study were excluded.

During study participation, the subjects were not allowed to take any of the following drugs: any drugs known or discussed to potentially induce liver enzymes
(e.g. rifampicin, dexamethasone, barbiturates, anticonvulsants, St. John’s wort); any drugs known to inhibit CYP3A4 (e.g. ketoconazole, verapamil, cimetidine, macrolides); any broad-spectrum antibiotics; and long-acting, injectable or implant hormonal therapy within 26 weeks prior to the screening visit, as well as any oral contraceptives other than the investigational treatment.

2.3 Interventions, Blinding

Silexan was presented in immediate-release soft gelatine capsules containing 160 mg of lavender oil. An identically matched placebo was available in which the smell of the study drug was concealed by flavouring the capsules with 0.08 mg of Lavender oil, i.e., with 1/2000 of the amount of Lavender oil contained in the active capsules. During randomised treatment, the subjects had to take one capsule of Silexan or placebo per day in the morning unchewed. A daily dose of 160 mg corresponds to twice the currently recommended dose of the marketed product. Moreover, Silexan 160 mg/day has been shown to be highly efficacious in patients with GAD [17].

Microgynon® is a combination oral contraceptive containing 0.03 mg of EE and 0.15 mg of LNG in a coated tablet. The drug is widely used in combination oral contraceptive interaction studies [18, 19] and was packed in the original marketed package, which carried an additional label presenting the information required for use in a clinical trial. Starting on the first day of each cycle during study participation, subjects were instructed to take one capsule of Silexan or placebo per day in the morning unchewed. A daily dose of 160 mg corresponds to twice the currently recommended dose of the marketed product. Moreover, Silexan 160 mg/day has been shown to be highly efficacious in patients with GAD [17].

The study participants had to complete at least 2 cycles of oral contraception with Microgynon® in order to achieve steady state conditions. Since LNG highly binds to sex hormone-binding globulin (SHBG) in plasma, and EE induces the SHBG binding capacity, LNG levels were expected to be in a steady state after 2 cycles. Eligible subjects were then randomised to 2 additional cycles with co-administration of Silexan during the first and placebo during the second cycle, or vice versa (Table 1). Double-blind treatment was always started on the first day of the pill cycle following randomisation. Treatment compliance was monitored by recording each drug intake in a diary as well as by counting the unused medication. Furthermore, plasma levels of linalool, the main metabolite of Silexan, were determined at weekly intervals during randomised treatment.

2.4 Study Schedule, Outcomes

During randomised treatment, weekly visits were scheduled on days 7, 14, 21, and 28 of each cycle. For each visit, a deviation of ±1 day was permitted. In addition, blood sampling for determining the pharmacokinetic (PK) parameters of EE and LNG was performed on 4 consecutive days starting on day 18 ± 1 day of each cycle: on each of these days EE and LNG trough levels were determined. Furthermore, blood samples for determining PK profiles were obtained at 0.5, 1, 1.5, 2, 3, 4, 6, 8, and 24 h after the intake of Microgynon® on the third and into the fourth day. The subjects were required to abstain from any food and beverages except for water after 22:00 h on the 2nd day of sampling and received standardised meals on the 3rd day. After the end of the 2nd cycle during randomised treatment, a follow-up examination was scheduled to be performed within 7–14 days. The samples were analysed for the determination of EE, LNG, and linalool levels using a validated high-performance liquid chromatography (HPLC)–tandem mass spectrometry (MS/MS) method and ConCalc for Windows (CCW; Integ GmbH) software version 1.15.

The primary outcome measures for assessing potential pharmacological interactions between Silexan and the combination oral contraceptive were the area under the curve over a dosing interval of τ = 24 h (AUCτ) derived from the 24-h concentration–time curves for EE and LNG at steady state. Secondary outcome measures included the EE and LNG peak concentration (Cmax), time to peak concentration (tmax), and trough levels. Ovarian follicle size measurements were performed through transvaginal ultrasound examinations at screening and at each weekly visit during randomised treatment, and the Hoogland score [20] was determined for each treatment cycle as a measure of ovarian activity. SHBG levels were determined at the end of the 3rd week (i.e., end of Microgynon® intake) during each cycle.

Assessment of safety and tolerability was based on spontaneous reports of adverse events (AEs), physical examinations and safety laboratory measurements.

2.5 Random Sequence Generation, Allocation

Concealment, Implementation

After subjects were determined as eligible for the trial, they were randomised to treatment with Silexan followed by placebo, or vice versa, at a ratio of 1:1. Fixed block randomisation was used; however, the investigator was not informed about the random block size until completion of the trial. A biometrician otherwise not involved in the trial generated the code using a validated computer program. Upon inclusion into randomised treatment, each subject received the lowest available number.

2.6 Statistical Methods, Sample Size

PK parameters were derived by non-compartmental methods using Phoenix WinNonlin® software (Certara, L.P., St
Louis, MO, USA). $C_{\text{max}}$ and $t_{\text{max}}$ were read directly from the data, the AUCs was derived by the linear trapezoidal rule, $t_{\text{1/2}} = (\ln 2)/k_z$, where the terminal rate constant was derived by log-linear regression, apparent clearance $CL/f = \text{AUC}_s/dose$, and apparent volume distribution $V_z/f = k_z9CL/f/k_z$.

PK measures determined from the EE and LNG concentration-time profiles were compared between Silexan and placebo using analysis of variance (ANOVA) models with main effects for sequence, subject within sequence, period, and treatment. Tests for AUCs and $C_{\text{max}}$ were performed after log transformation, and for $t_{\text{max}}$ on the original untransformed scale. Confidence intervals determined on the log scale were back-transformed into the original scale.

In the absence of other guidance, the assessment of the pharmacological interacting potential of Silexan was based on the same approach as when testing for bioequivalence [21]: the absence of a clinically relevant PK interaction between Silexan and EE or LNG was concluded if the 90 % confidence intervals for the AUC ratios (Silexan/placebo) determined from the ANOVA models were included within the standard 80–125 % equivalence range. Since the confidence intervals for the AUC ratios of both hormones were required to fall within the pre-specified equivalence range simultaneously, type I error level adjustment for multiple testing was not required.

Comparable ANOVA models were also applied to $C_{\text{max}}$ and $t_{\text{max}}$; however, these were interpreted descriptively. For SHBG concentration, an ANOVA model was fitted using the same factors after log-transformation of the dependent variable. All other interaction and safety measures were evaluated using methods of descriptive data analysis.

The sample size estimation was based on the assumption of a true within-subject coefficient of variation of 20 % for $\ln(\text{AUC}_\tau)$ of both EE and LNG. When the true AUC$_\tau$ ratio between Silexan and placebo is between 0.95 and 1.05, a sample size of 20 subjects will have at least 80 % power for rejecting the null hypothesis predicting an interaction between Silexan and EE or LNG according to the above criteria. In order to compensate for study participants who did not complete both treatment phases, a total of 24 subjects were to be randomised.

### 3 Results

#### 3.1 Recruitment, Participant Flow, Protocol Deviations

Between November 2009 and May 2010, a total of 32 women were included and 24 were randomised and treated. All randomised subjects completed the study as scheduled. All treated subjects were analysed for safety. Two subjects were excluded from all PK and pharmacodynamic (PD) analyses; one took prohibited concomitant medication and one missed one dose of Microgynon®, which was considered a major protocol deviation since it could not be excluded that the dose might have been missed on the day before obtaining the PK profile. One other subject mistakenly received the investigational treatments in reverse order compared with the randomisation; however, she was included into all analyses and evaluated as treated. All decisions concerning the subjects’ eligibility for the different analyses were made before code breaking.

#### 3.2 Sample Characteristics

The randomised study participants were aged between 21 and 37 years (mean ± standard deviation [SD] 27.3 ± 4.1 years) and had a body mass index ranging between 19.3 and 26.8 kg/m$^2$ (22.2 ± 2.0 kg/m$^2$). All women were fertile and had already used hormonal contraception before inclusion. All participants were of Caucasian race, although study participation was open to all ethnic groups. Nine of the 24 women (37.5 %) were current smokers (up to a maximum of ten cigarettes per day allowed) and seven

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(29.2%) were ex-smokers; 23 (95.8%) drank alcohol occasionally. The most frequent diseases in medical history were myopia (ten subjects, 41.7%) and headache (eight subjects, 33.3%). In the physical examination performed at screening, none of the randomised subjects presented with findings from which an inference with the study procedures or outcomes was to be expected. Of note, none of the women had abnormal findings in the gynaecological examination.

3.3 Treatment Compliance

According to capsule counting, compliance with Silexan or placebo was 100% in 23 of the 24 subjects. One woman took one additional capsule during the first treatment period and left out one during the second. The analysis of linalool plasma concentrations confirmed that all participants had taken Silexan according to the randomisation, whereas the treatment sequence had been mixed up in one woman.

Of the 24 randomised participants, 22 took Microgynon® as prescribed. One missed one dose (and was therefore excluded from all PK and PD analyses), whereas another woman took one additional dose while under co-medication with placebo (by mistake two tablets at once).

3.4 Pharmacokinetic and Pharmacodynamic Evaluation

Figure 1 shows the plasma concentration–time profiles for EE and LNG during co-administration of Silexan or placebo. The numerical results are presented in Table 2.

For EE and LNG, the 90% confidence intervals for the AUCt and Cmax ratios between Silexan and placebo were completely included in the pre-specified equivalence range so that an influence of co-administration of Silexan on these PK measures could be excluded. Whereas the results do not indicate a significant influence of Silexan on tmax of EE, the tmax of LNG was prolonged by an average of about 17 min (point estimate for the mean value difference), albeit without significantly affecting Cmax or the AUCt.

The results for the main PD outcomes and biomarkers are summarised in Table 3. The only case of an active follicle-like structure (>13 mm follicular growth accompanied by estradiol production) was observed during placebo treatment. Average follicle sizes decreased during Microgynon® intake until day 21 but showed an increase during the pill-free 4th week of the cycles. Mean values were similar under co-medication with Silexan and placebo; however, at three of the four post-baseline visits, the larger average for both follicles was observed during placebo treatment. Mean endometrial thickness was also comparable during both treatment phases.

Comparable mean levels for progesterone, estradiol, and SHBG were observed during both treatment phases (Table 3). Whereas progesterone showed only minor changes over time, estradiol decreased markedly until the end of the 2nd week of the cycles and continued to decrease slightly during the 3rd week, whereas a sharp average increase was observed in the pill-free 4th week. During the first 3 weeks of each cycle, while administering Microgynon®, none of the women showed an estradiol level >50 pg/ml, and one woman had a progesterone level >3 nmol/l on day 21 while under co-administration of placebo.

Individual values for the linalool concentrations indicate that the PK evaluation has been performed after a steady state was reached.

3.5 Safety/Tolerability

During randomised treatment, 36 AEs were reported by 17 of the 24 women (70.8%) under co-administration of Table 2 Main pharmacokinetic parameters (n = 22)

|                | Placebo                     | Silexan                     | Ratio or difference 90% CI | AUCt (pg × h/ml)a | Cmax (pg/ml)a | tmax (h)b |
|----------------|-----------------------------|-----------------------------|-----------------------------|------------------|--------------|----------|
| Ethinyl estradiol |                             |                             |                             | 937.4 ± 265.8     | 105.6 ± 33.2  | 1.5 (1.0–3.0) |
| Silexan         |                             |                             |                             | 911.7 ± 263.3     | 106.6 ± 38.3  | 1.5 (0.9–3.0) |
| Ratio or difference 90% CI | 0.97 (0.93–1.02) | 0.99 (0.92–1.07) | 0.11 (0.05–0.53) | 0.94 (0.89–1.00) | 0.96 (0.91–1.01) | 0.29 (0.05–0.53) |
| Levonorgestrel   |                             |                             |                             | 94.3 ± 34.6       | 8.0 ± 2.6     | 1.0 (0.5–1.6) |
| Placebo         |                             |                             |                             | 90.3 ± 37.7       | 7.7 ± 2.8     | 1.1 (1.0–3.0) |
| Silexan         |                             |                             |                             | 90.3 ± 37.7       | 7.7 ± 2.8     | 1.1 (1.0–3.0) |
| Ratio or difference 90% CI | 1.00 (0.99–1.01) | 0.96 (0.91–1.01) | 0.29 (0.05–0.53) | 0.94 (0.89–1.00) | 0.96 (0.91–1.01) | 0.29 (0.05–0.53) |

a Mean ± SD, back-transformed CI for ratio between marginal (geometric) means from ANOVA model, Silexan/placebo
b Median (range), confidence interval for difference between marginal (arithmetic) means from ANOVA model, Silexan—placebo
ANOVA analysis of variation, AUCt area under the concentration–time curve over a dosing interval of t = 24 h, CI confidence interval, Cmax peak concentration, SD standard deviation, tmax time to Cmax

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Microgynon\textsuperscript{®} and Silexan, whereas 23 events were reported by 16 subjects (66.7\%) in the Microgynon\textsuperscript{®} plus placebo phase. All events were mild or moderate in intensity, and all were non-serious. AEs in which a causal relationship to the investigational treatment could not be excluded occurred in ten subjects during Microgynon\textsuperscript{®} and Silexan treatment (gastrointestinal disorders, nine subjects; weight increased, one subject; headache, two subjects; and acne, one subject) as well as in two subjects during Microgynon\textsuperscript{®} plus placebo treatment (headache and fatigue, one subject each). Safety laboratory measures reflected changes that were attributed to frequent blood sampling (e.g., anaemia) but were otherwise uneventful.

4 Discussion

It is important to investigate whether anxiolytic drugs may modify the bioavailability and the clinical efficacy of oral contraceptives with steroid hormones. The study demonstrates that Silexan does not show clinically relevant pharmacological interactions with EE or
Data are presented as mean ± SD, which are the active ingredients of Microgynon® C210. Silexan was well tolerated.

Smoking may alter the PK properties of oral contraceptive steroids. Since about one-third of the women in the study sample were mild smokers (subjects smoking more than ten cigarettes per day were excluded from participation), this might have contributed to large inter-individual variability in plasma concentrations, notably in EE. However, the inter- and intra-individual variability of the EE plasma levels in our trial was comparable with the literature [22] and, moreover, higher variability would have biased the data towards an acceptance of the null hypothesis predicting an interaction between Silexan and EE and/or LNG. Since the null hypothesis could be rejected and an interaction could be excluded in accordance with the prespecified criteria, an introduction of bias due to the inclusion of mild smokers can also be excluded.

Progesterone, estradiol, and SHBG level monitoring also did not indicate an impairment of the contraceptive efficacy of Microgynon® during co-administration of Silexan. Throughout the entire cycle of randomised treatment, the average levels of these biomarkers during co-administration of Silexan were always in the range of those observed under placebo treatment.

Silexan had no appreciable effect on mean endometrial thickness and follicle size. The measured individual follicle sizes were always within the range expected during adequate ovarian suppression [23], and a follicle size exceeding a threshold value of 13 mm was observed only once in a woman receiving placebo.

AEs observed during this trial in healthy volunteers were never more than moderately severe and temporary; abnormal laboratory values were more often related to the study procedures (e.g., to frequent blood sampling) than to study medication intake. Among the events assessed to be potentially attributable to Silexan, the only effect that was observed in more than one subject but also not during placebo treatment was eructation. Mild gastrointestinal disorders are among the known side effects of Silexan, but were always tolerable and did not interfere with treatment compliance.

In conclusion, Silexan, at a dose of 160 mg/day, corresponding to twice the currently recommended dose, did not adversely affect the relative bioavailability and the suppressive efficacy on hormonal and ovarian status of the oral combination contraceptive Microgynon® whose active ingredients are EE and LNG. Silexan was well tolerated.

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