The effect of combined oral contraception on testosterone levels in healthy women: a systematic review and meta-analysis

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BACKGROUND: Combined oral contraceptives (COCs) reduce levels of androgen, especially testosterone (T), by inhibiting ovarian and adrenal androgen synthesis and by increasing levels of sex hormone-binding globulin (SHBG). Although this suppressive effect has been investigated by numerous studies over many years, to our knowledge no systematic review concerning this issue had been performed. This systematic review and meta-analysis was performed to evaluate the effect of COCs on concentrations of total T, free T and SHBG in healthy women and to evaluate differences between the various types of COCs (e.g. estrogen dose, type of progestin) and the assays used to assess total T and free T.

METHODS: A review of the literature was performed using database searches (MEDLINE, EMBASE and the Cochrane Central Register of Clinical Trials) and all publications (from inception date until July 2012) investigating the effect of COCs on androgen levels in healthy women were considered eligible for selection. Three reviewers were involved in study selection, data extraction and critical appraisal. For the meta-analysis, data on total T, free T and SHBG were extracted and combined using random effects analysis. Additional subgroup analyses were performed to evaluate differences between the various types of COCs (e.g. estrogen dose, type of progestin) and the assays used to assess total T or free T.
**RESULTS:** A total of 151 records were identified by systematic review and 42 studies with a total of 1495 healthy young women (age range: 18–40 years) were included in the meta-analysis. All included studies were experimental studies and 21 were non-comparative. Pooling of the results derived from all the included papers showed that total T levels significantly decreased during COC use [mean difference (MD) (95% confidence interval, CI) −0.49 nmol/l (−0.55, −0.42); P < 0.001]. Significantly lower levels of free T were also found [relative change (95% CI) 0.39 (0.35, 0.43); P < 0.001], with a mean decrease of 61%. On the contrary, SHBG concentrations significantly increased during all types of COC use [MD (95% CI) 0.98 nmol/l (86.43, 11.73); P < 0.001]. Subgroup analyses revealed that COCs containing 20–25 μg EE had similar effects on total and free T compared with COCs with 30–35 μg EE. In addition, suppressive effects on T levels were not different when comparing different types of progestins. However, subgroup analyses for the estrogen dose and the progestin type in relation to changes in SHBG levels did show significant differences: COCs containing second generation progestins and/or the lower estrogen doses (20–25 μg EE) were found to have less impact on SHBG concentrations.

**CONCLUSIONS:** The current literature review and meta-analysis demonstrates that COCs decrease circulating levels of total T and free T and increase SHBG concentrations. Due to the SHBG increase, free T levels decrease twice as much as total T. The estrogen dose and progestin type of the COC do not influence the decline of total and free T, but both affect SHBG. The clinical implications of suppressed androgen levels during COC use remain to be elucidated.

**Key words:** combined oral contraception / androgens / testosterone / SHBG / systematic review

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**Introduction**

In normo-ovulatory women, testosterone (T) levels (Burger, 2002; Speroff and Fritz, 2005; Haring et al., 2012; Pesant et al., 2012) are reported to vary between 0.42 and 2.94 nmol/l (Haring et al., 2012; Pesant et al., 2012). T arises from three sources in the female. Approximately 50–60% is derived from the peripheral conversion of the prohormones androstenedione (AD) and dehydroepiandrosterone (DHEA) and its sulphate (DHEA-S), whereas 25% is secreted by the ovary and 25% by the adrenal gland (Longcope, 1986; Bachmann et al., 2002; Burger, 2002). Around 65–70% of circulating T is bound and inactivated by sex-hormone-binding globulin (SHBG). Most of the remaining 30–35% is bound by albumin and only 0.5–3% represents freely circulating T (free T). Since the binding of T to albumin is rather weak, the free and albumin-bound T together are defined as the bioavailable T (Dunn et al., 1981; Speroff and Fritz, 2005).

There is much debate regarding adequate methods for measuring total T as well as free T concentrations (Vesper et al., 2008; Legro et al., 2010; Rosner and Vesper, 2010; Vesper and Botelho, 2010; Haring et al., 2012). The most accurate method to assess total T is liquid chromatography-tandem mass spectrometry (LC-MS/MS) (Rosner et al., 2007; Vesper et al., 2009), although even with LC-MS/MS assays variation in precision exists (Legro et al., 2010). The ‘Princeton Consensus Statement’ (Bachmann et al., 2002) recommends equilibrium dialysis as the ‘gold standard’ method to measure free T concentrations. These methods, however, are very expensive, labour intensive and not available to the great majority of the laboratories. Because of their ease and low costs, immunoassays [radioimmunoassay (RIA) and electrochemiluminescence immunoassays (ECLIA)] are most commonly used in clinical practice; however, they are known for their inaccuracy and variability (Taieb et al., 2003; Miller et al., 2004; Groenestege et al., 2012). In particular, immunoassays are affected by cross-reactivity with other steroids (Middle, 2007; Benton et al., 2011) and by high levels of SHBG (Masters and Hahnel, 1989; Wierman et al., 2006). The accuracy of RIA can be improved by the addition of an extraction step or chromatography (Rosner et al., 2007; Groenestege et al., 2012), but these procedures are time consuming. The very low concentrations of free T in female blood (35–700 pmol/l; Speroff and Fritz, 2005) are an additional difficulty in obtaining accurate measurements (Taieb et al., 2003; Groenestege et al., 2012). Therefore, free T is often calculated based on total T, SHBG and albumin concentrations (Södergård et al., 1982; Vermeulen et al., 1999). This method has been demonstrated to be a reliable measure for free T concentrations (Rinaldi et al., 2002; Miller et al., 2004).

Combined oral contraceptives (COCs) reduce androgen levels (Fern et al., 1978; Aden et al., 1998; Wiegart et al., 2003; Ágren et al., 2011a). In particular, blood levels of T, the most potent circulating androgen in women, decrease by up to 50% (Van der Vange et al., 1990; Coen et al., 1996; Greco et al., 2007). Three possible underlying mechanisms may be held responsible for this effect: (i) Suppression of ovarian androgen synthesis; (ii) increased SHBG levels and (iii) suppression of adrenal androgen synthesis.

To exert their contraceptive effect, COCs suppress the gonadotrophins (Gaspar et al., 1984; Aden et al., 1998). Low levels of LH result in an inhibition of the LH-dependent synthesis of T by the ovarian theca cells (Speroff and Fritz, 2005; Davison and Bell, 2006). As the suppression of T already occurs before the LH levels drops, it is suggested that the contraceptive steroids may have a direct inhibitory effect on ovarian androgen synthesis (Kuhl et al., 1985; Jung-Hoffman et al., 1988a; Aden et al., 1998).

SHBG is a carrier protein synthesized by the liver. Its synthesis is stimulated by estrogens and inhibited by androgens (Granger et al., 1982; Speroff and Fritz, 2005). The estrogenic component of COCs, ethinyl estradiol (EE), induces hepatic SHBG production in a dose-dependent manner, resulting in elevated circulating blood levels of SHBG (Jung-Hoffmann and Kuhl, 1987; Van der Vange et al., 1990; Hammond et al., 2008; Ágren et al., 2011a). Consequently, more T is bound and inactivated, resulting in lower levels of T and the free androgen index (which is a measure of bioavailable T, i.e. free and albumin-bound T) (Van der Vange et al., 1990; Coen et al., 1996; Boyd et al., 2001; Wiegart et al., 2003; Ágren et al., 2011a). This EE-induced increase of SHBG can be counteracted by the progestin-component of COC, as a result of the androgenicity of the progestin. COCs containing a progestin with androgenic activity [e.g. levonorgestrel (LNG)] will...
induce a less pronounced increase in SHBG, whereas progestins with concomitant anti-androgenic activity (e.g. cyproterone acetate (CPA)) will lead to higher SHBG levels (Van Kammen et al., 1975; Bergink et al., 1981; Hammond et al., 1984; Van der Vange et al., 1990; Odling et al., 2002; Ågren et al., 2011a, b).

The third mechanism of action causing lower levels of androgens is the inhibitory effect of COCs on adrenal androgen synthesis (Fern et al., 1978; Madden et al., 1978; Carlström et al., 2002). Several studies have shown that COCs decrease both DHEA and DHEA-S concentrations in blood (Gaspard et al., 1984; Falsetti et al., 1987; Wiegratz et al., 1995; Coenen et al., 1996, Wiegratz et al., 2003; Ågren et al., 2011a). The underlying mechanism is not yet established, but it has been suggested that a reduced release of adrenocorticotropic hormone (ACTH) and increased levels of cortisol during COC use may account for the reduced concentrations of adrenal androgens (Carr et al., 1979; Klove et al., 1984; Carlström et al., 2002). Consistent with this, ACTH is known to stimulate adrenal androgen secretion (Rosenfield et al., 1972; Longcope, 1986).

Although little attention has been paid to the suppressive effect of COCs on androgens along with its potential clinical implications. Awareness of the significance of androgens for women is increasing, however, and T deficiency is thought to be associated with a broad range of undesired effects including diminished well-being and quality of life, mood changes (depression, irritation, moodiness), loss of energy, cognitive disturbances, interference with optimal sexual functioning, declining muscle mass and strength and lowering of bone density (Bachmann et al., 2002; Traish et al., 2007).

The study results [mean and standard deviation (SD)] for total T, free T and SHBG in healthy women were reviewed for additional publications that could be used in this review.

**Selection criteria**

All studies investigating the effect of COCs on androgen levels in healthy women were eligible for selection. At least two of the following three outcome parameters had to be reported: total T, free T and SHBG. Exclusion criteria were studies in men or animals, women suffering from acne, polycystic ovary syndrome, hirsutism or endometriosis, (post)menopausal women, no or other types of hormonal contraception (e.g. progestin-only contraceptives, implants, patches, vaginal rings, intrauterine device), COC with a continuous regimen, emergency contraception, treatment duration of less than one cycle (i.e. one cycle is 28 days), other outcome parameters (e.g. cervical mucus, acne, endometriosis) and bioequivalence/bioavailable studies. Reviews, case reports, letters to the editor and conference papers were also excluded.

The results of the searches were screened to meet the pre-defined eligibility criteria. A first selection was performed based on their titles, followed by a second selection performed by two independent reviewers (Y.Z. and W.W.), who reviewed the abstracts of all remaining records. Any disagreement in the selection of abstracts was resolved by consensus or by a third reviewer (B.F.). Full text articles for review and data processing were obtained for all selected abstracts.

**Data extraction**

Data were extracted from full text articles into a specially designed data extraction form by one reviewer (Y.Z.) in close consultation with another reviewer (M.E.). Any obscurities were discussed and decisions were taken by consensus. During the process of full text review and data extraction, studies were excluded if they did not fulfil the eligibility criteria or if the reported results were in a format that could not be used or were judged to be of insufficient quality. No investigators were contacted in the case of missing or obscure data.

The following data were extracted: authors, year of publication, title, study design, characteristics of the study population (e.g. study criteria, age, body weight/height/BMI, use of previous hormonal contraceptives), number of randomized, early discontinued and completed subjects, details on intervention (e.g. pill composition and regimen, multi- or monophasic, treatment duration and number of treatment groups), study objective, outcome parameters with special attention to the androgens and SHBG, sampling details (e.g. timing in treatment period and cycle day), analytical methods and statistical methods.

The study results [mean and standard deviation (SD)] for total T, free T and SHBG concentrations in International System (SI) units were extracted in a separate standardized form. In a few cases, study results were only presented in figures and data were obtained by measuring directly from these figures that were printed on a large scale. T concentrations reported in conventional units were converted to SI units using the following formula: conversion factor (CF) × conventional unit (e.g. gram) = SI (e.g. mol), where CF for T was 3.467. For the conversion of SHBG concentrations, the molecular weight of SHBG (43 780 g/mol) was used.

The standard error (SE) or SE of the mean (SEM) were converted to SD using the formula: $SD = SEM \times \sqrt{n}$, where $n$ is the sample size. In a few cases, neither the SD nor the SEM was reported. The SD was calculated if the SD (or SEM) was reported for the other time-points in the study using the following formula: $SD_2 = SD_1 \times (\text{mean}_2/\text{mean}_1)$, where $SD_2$ is the SD which is unknown, $SD_1$ is the SD reported for another time point in the study, $\text{mean}_2$ is the mean that belongs to $SD_2$ and $\text{mean}_1$ the mean that belongs to $SD_1$. A 95% confidence interval (CI) was converted to SD using the following formula: $(\text{CI}_{1%} - \text{CI}_{95%})/((2 \times 1.96) \times \sqrt{n})$, where CI is the
upper limit, $C_{\text{up}}$, is the lower limit and $n$ is the sample size. In cases where the change from baseline was reported, the end of treatment value was calculated from the baseline value minus the change from baseline. Finally, when a median plus range of variability was reported instead of a mean plus SD, the mean and SD were calculated by assuming the data had a log-normal distribution.

To prevent extraction errors, a quality control check between the final data used in the meta-analysis and the original publications was performed.

**Critical appraisal**

The risk of bias for the individual studies included in the meta-analysis was assessed based on guidance provided in the Cochrane Handbook for Systematic Reviews of Interventions (Higgins and Green, 2011). All studies were critically appraised for selection bias (sequence generation, allocation concealment), performance bias, detection bias, attrition bias, reporting bias and other bias by two reviewers (Y.Z. and M.E.), in relation to the outcome parameters used in the meta-analysis. Studies were not excluded from the meta-analysis based on a high risk of bias, but their risk of bias was taken into consideration during the interpretation of the results.

**Statistical analysis**

Only a few papers reported data from observational studies comparing women using COC with women not using COC (control group) and no specific type of COC was investigated in these studies. The majority of the selected studies were experimental studies investigating the effects of one or more COCs on different outcome parameters. Because an identical control group in these studies could not be identified, the experimental studies which compared the pretreatment study results (i.e. total T, free T and/or SHBG levels) with the end of treatment results were used in the meta-analysis. By using these studies, differences between the various types of COC could also be evaluated. In this context, an important additional inclusion criterion was used; study participants should not have taken a hormonal contraceptive prior to starting the study medication (including a wash-out period).

For the meta-analysis, the inverse variance method with random effects as analysis model was applied. As effect measures, the mean difference (MD) and associated 95% CIs were calculated based on the means of the pre-treatment and the end of treatment levels of total T, free T and SHBG. Data of each treatment group were entered separately. For the end of treatment time point, the assessment (mean/SD) after a 6-cycle treatment duration was used for reasons of standardization. In cases where this time point was not reported, the three cycle or the nearest available reported time point was used.

In cases where the data showed large variation (e.g. larger SD in studies with larger mean values, plus identification of a significant heterogeneity in the meta-analysis), logarithms of these values were used to reduce the wide range of values. For the pooled concentrations in logarithmic scale, the generic inverse variance method with random effects was used. As effect measure, the relative change and associated 95% CI from pretreatment to end of treatment was calculated. Heterogeneity between the results of the different studies was examined using the $I^2$ value. Heterogeneity was found to be significant when $I^2 > 65%$.

Subgroup analyses were performed for different estrogen doses (20–25 μg EE versus 30–35 μg EE) and for the type of progestin (first generation estranes) versus second generation (gonanes) versus third generation (gonanes) versus fourth or unclassified generation) on all outcome parameters. The classification of the progestins was based on their chemical structure combined with the time of introduction.

As the methods to assess total T and free T concentrations could influence the outcome of the meta-analysis, subgroup analyses for the type of assay (RIA/ECLIA/enzyme-linked immunosorbent assay (ELISA) versus RIA with extraction/chromatography) and for the method to determine free T concentrations (direct immunoassay versus calculation or equilibrium dialysis) were performed. In cases where more than two subgroups were included and a significant difference was identified, it was pre-determined which subgroups would undergo separate statistical testing. For the progestin type, an additional comparison was made between second versus third generation or between second versus third generation, only including monophasic COCs containing 30–35 μg EE.

All analyses were performed with Review Manager (RevMan), version 5.2 (2012).

**Results**

**Study selection**

A flow chart of the included/excluded studies is shown in Fig. 1. Database searches yielded 904 records from MEDLINE, 929 from EMBASE and 538 from the Cochrane Central Register of Clinical Trials. Following the first review based on the title, 193 records remained (122 from MEDLINE, 39 from EMBASE and 32 from the Cochrane Central Register of Clinical Trials). After removal of duplicate records, 151 records remained and the abstracts were reviewed based on the pre-defined eligibility criteria. A total of 66 records were selected for full text review and data processing. During this phase, 27 papers were excluded, 3 studies were included by hand search and 42 studies were finally included in the meta-analysis. Table I shows the reasons for exclusion. Not all studies reported all three outcome parameters, therefore 39 studies were included in the meta-analysis for COC effect on total T and SHBG and 29 studies were included for the effect on free T.

**Characteristics of included studies**

Characteristics of included studies are reported in Table II. All 42 studies were experimental with one or more treatment groups. Only three studies used a double-blind or single-blind design (Wiegratz et al., 2003; Greco et al., 2007; Legro et al., 2008). There were 21 studies which were non-comparative, whereas 16 studies compared two COCs and 5 studies compared 3 or more different types of COCs. Various types of COCs were investigated including both monophasic and multiphasic regimens. Almost all included studies investigated the effects of a COC with a 21:7 regimen, except for two studies that both included a COC with a 24:4 regimen (Duijkers et al., 2010; Ågren et al., 2011a). Most of the studies included COCs containing EE as the estrogenic component, but four studies (Wiegratz et al., 2003; Duijkers et al., 2010; Ågren et al., 2011a; Caruso et al., 2011) also investigated the effects of a COC with estradiol (E2) or E2 valerate. These treatments often did not qualify for the subgroup analyses. The duration of treatment varied from 1 treatment cycle (Kuhnz et al., 1991) up to 12 treatment cycles (Jung-Hoffman et al., 1988a; Åkerlund et al., 1994; Wiegratz et al., 1995; Sänger et al., 2008). Treatment was assessed for 6 cycles (the effect measure used for the meta-analysis) in 16 studies, 3 cycles in 14 studies and for 12 studies another time point had to be taken for the analyses.

The total number of women randomized in all studies was 1495 with an average number of 21 (range 5–60) women per treatment group. The study population consisted of healthy women in the age range of 18–40 years. Two studies (Sieberg et al., 1984; De Leo et al., 1991) also enrolled younger women (minimum age of 15 and 17 years, respectively).
One study enrolled Chinese women (Song et al., 1992). The period without intake of hormonal contraceptives before the start of the study was more than 2 or 3 months in most studies, ranging from 4 to 6 weeks (Wiegratz et al., 2003; Sänger et al., 2008; Ågren et al., 2011a) up to 6 months (Boyd et al., 2001; Battaglia et al., 2012) and 12 months (Siegberg et al., 1984; Moutos et al., 1995). For those studies that did not specify a pre-study period without hormonal contraceptives, a regular menstrual cycle with or without proven ovulation was an inclusion criterion (Granger et al., 1982; De Leo et al., 1991; Janaud et al., 1992; Spona et al., 1996; Rickenlund et al., 2004; Duijkers et al., 2010; Caruso et al., 2011).

With regard to the methods to determine total T and free T concentrations, 30 studies used a direct immunoassay, 9 studies incorporated an extraction/chromatography step before RIA and no studies used the LC-MS/MS method. One study did not specify the type of assay used to measure total T concentrations (Janaud et al., 1992). The free T concentrations were analysed by a direct immunoassay in 18 studies, determined by calculation (based on total T, SHBG and albumin concentrations) in 8 studies and by equilibrium dialysis in 3 studies. Five studies used the formula of Vermeulen et al. (1971, 1999).

**Methodological quality**

Outcome parameters used in the meta-analysis and the type of comparison (a within-subject comparison on pretreatment (no COC) versus end of treatment (COC)) were assessed for risk of bias. An overview of the risk of bias of the studies included in the meta-analysis is provided in Table II and Supplementary data, Table SI.

To assess selection bias, reported procedures on sequence generation and allocation concealment were judged for all studies. Often randomization and concealment was not applicable for studies, because only one treatment group was investigated. Furthermore, selection bias was expected not to have a major impact on the outcome of the analyses as study participants acted as their own control. Overall, three studies (Granger et al., 1982; Gaspard et al., 1984; Hammond et al., 1984) were judged as having inadequate sequence generation or
Table I Reasons for excluding articles.

| First author, year | Reason |
|--------------------|--------|
| Bancroft et al. (1980) | Wrong study population (women with impaired sexual function); baseline comparison problem group versus non-problem group, but only problem group takes part in study phase with prescribed study medication |
| Bergink et al. (1981) | Study results are reported as geometric mean, which could not be converted to mean; no SD or SE reported; unit of Free T (%) could not be converted |
| Spona et al. (1986) | Treatment duration was unclear (i.e. from Day 5 to Day 25 and unclear if 21:7 regimen); not specified if results are reported in mean/SD; not clear if study participants had been using previous steroids prior to starting the study medication; not able to convert the unit of SHBG to SI units |
| Jung-Hoffmann and Kuhl (1987) | Assumption same study as reported by Kuhl et al. (1985); the latter has been used in the meta-analysis, but publication of Jung-Hoffmann and Kuhl (1987) was also used to obtain additional information |
| Jung-Hoffmann et al. (1988b) | The same study was published in an English journal by the same first author (Jung-Hoffmann et al., 1988a). The latter has been used in the meta-analysis, but this publication (Jung-Hoffmann et al., 1988b) was also used to obtain additional information |
| Alexander et al. (1990) | Observational study (COC users versus non-COC users); substudy of Bancroft et al. (1991a, b); various types of COCs were used |
| Van der Vange et al. (1990) | Study results are reported as geometric mean, which could not be converted to mean and SD is not reported. Results are only presented in figures with a low quality for direct measurement |
| Bancroft et al. (1991a) | Observational study (non-COC users versus users); various types of COCs were used; study results reported in two publications Bancroft et al. (1991a, b) |
| Bancroft et al. (1991b) | Observational study (non-COC users versus users); various types of COCs were used; study results reported in two publications Bancroft et al. (1991a, b) |
| Sobrio et al. (1991) | Not able to obtain a copy of the full text |
| Janaud et al. (1993) | Not able to obtain a copy of the full text; assumption same study as reported by Janaud et al. (1992) |
| Pasinetti and Falsetti (1993) | Not able to obtain a copy of the full text |
| Erdmann et al. (1994) | Wrong study population (women with acne, seborrhoea, hirsutism and alopecia) |
| Coenen et al. (1995) | Not able to obtain a copy of the full text; assumption same study as reported by Coenen et al. (1996) |
| Oettel et al. (1997) | Not able to obtain a copy of the full text |
| Sulak et al. (1999) | Wrong study population (a part of the study population is post-partum [given birth <= 60 days of screening] and results are not separately reported); units for the lab parameters are not specified |
| Stanczyk et al. (2000) | Not able to obtain a copy of the full text |
| Aleyeva (2002) | Not able to obtain a copy of the full text |
| Endrikat et al. (2002) | Pretreatment values for total T and free T were measured on Day 1 of treatment Cycle 1, which was directly after a 7-day pill-free period. This pill-free period followed the single dose administration part of the study (one tablet on Day 21 of menstrual cycle). This measurement is not a real pretreatment value. In addition, no SD or SEM reported for total T concentrations at pretreatment |
| Oranratanaphan and Taneapanichskul (2006) | Not able to obtain a copy of the full text |
| Graham et al. (2007) | Publication combines results of two studies; results of one of these studies are also reported by Greco et al. (2007). The latter has been used in the meta-analysis, but publication of Graham et al. (2007) was also used to obtain additional information |
| Winkler and Sudik (2009) | Results were judged to be of insufficient quality; SHBG converted, but after conversion values do not seem to be correct; all measurements for free T values during treatment are 2.08 pmol/l, therefore no SD can be reported (assumed that LOQ of assay was 2.08 pmol/l, therefore free T concentrations could not be measured and the LOQ has been reported. |
| Schaffir et al. (2010) | Observational study and wrong study population (women are using a COC at pretreatment) |
| Heiman et al. (2011) | Observational study; no study medication; comparison between women with a normal sexual function and women with Hypoactive Sexual Desire Disorder of which half of the group was using hormonal contraception |
| Sanam and Ziba (2011) | Results were found to be of insufficient quality (e.g. not reported if results are expressed as mean/SD; seems that reported SD is sometimes the mean and vice versa [see body weight] and wrong numbers reported (e.g. baseline acne). |
| Elaut et al. (2012) | Study results are reported in boxplots from which no mean could be retrieved; no pretreatment assessment for COC-only group reported |
| Haring et al. (2012) | Observational study; no study medication; study was performed to establish age-specific reference ranges for serum sex hormone concentrations in women (without and with hormonal contraception) using mass spectrometry (LC-MS/MS method) |
| First author, year | Study design                  | Study population                                | Characteristics per treatment group | Study treatment (duration, composition and regimen) | End-points used in meta-analysis | Details hormone assessment (type of assay or calculation) | Domains with low risk of bias (n)† |
|-------------------|-------------------------------|------------------------------------------------|-------------------------------------|-----------------------------------------------|---------------------------------|----------------------------------------------------------|----------------------------------|
| Cullberg et al. (1982) | Randomized, comparative, parallel group | Healthy women 18–36 years regular menstrual cycle No hormonal contraceptives: ≥ 3 months | 1 n = 10 Mean age: NR Weight/BMI: NR | 3 cycles 30 EE/150DSG 21:7 | Control cycle (day NR) Tr. cycle 3 (Day 21) | Extraction/ chromatography RIA Intra-assay CV: <6% Inter-assay CV: <10% LLQ: NR | Not applicable 5 |
|                   |                               |                                               | 2 n = 10 Mean age: NR Weight/BMI: NR | 3 cycles 30 EE/150 LNG 21:7 |                                          |                                           |                                   |
| Granger et al. (1982)² | Comparative, parallel group | Healthy women 20–30 years normal menstrual cycle No OC: not specified, but only normally cycling women were included | 1 n = 5 Mean age: NR Weight/BMI: NR | 2 cycles 30 EE/300NG 21:7 | Control cycle (early proliferative phase and late luteal phase) Tr. Cycle 2 (Days 18–21) | Extraction/ chromatography RIA Intra-assay CV: NR Inter-assay CV: NR LLQ: NR | Not applicable 4 |
|                   |                               |                                               | 2 n = 5 Mean age: NR Weight/BMI: NR | 2 cycles 35 EE/400NET 21:7 |                                          |                                           |                                   |
| Vermeulen and Thiery (1982) | Non-comparative | Healthy women 20–23 years Normal menstrual cycle No OC: ≥ 2 months | 1 n = 12 Mean age: NR Weight/BMI: NR | 6 cycles 30-40-30 EE/50-75-125 LNG (6-10), 21:7 | Control cycle (NR) Tr. cycle 6 (Days 18–21) | RIA Intra-assay CV: NR Inter-assay CV: NR LLQ: NR | Equilibrium dialysis 6 (measuring free fraction which is used to calculate the AFTC (Vermeulen et al., 1971)) |
|                   |                               |                                               | 2 n = 13 Mean age: 22.75 years Weight: NR (normal body weight) | 6 cycles 30-40-30 EE/50-75-125 LNG (6-10), 21:7 | Control cycle (7 days before presumed menses) Tr. cycle 6 (Days 19–21) | Extraction/ chromatography RIA Intra-assay CV: NR Inter-assay CV: NR LLQ: NR | AFTC; calculated based on total T, SHBG and albumin (Vermeulen et al., 1971) 5 |
| Gaspard et al. (1983) | Comparative, parallel group | Healthy women No OC: ≥ 8 weeks | 1 n = 13 Mean age: 22.75 years Weight: NR (normal body weight) | 6 cycles 50–50 EE/0-125DSG (7–14), 21:7 |                                         |                                           |                                   |
|                   |                               |                                               | 2 n = 13 Mean age: 22.75 years Weight: NR (normal body weight) | 6 cycles 50–50 EE/0-125DSG (7–14), 21:7 |                                         |                                           |                                   |
| Study                | Design                          | Population                        | OC Duration | Mean Age and Weight | Cycle Details                                                                 | Assay Details                                                                 | Notes                                                                 |
|---------------------|---------------------------------|-----------------------------------|-------------|---------------------|-----------------------------------------------------------------------------|----------------------------------------------------------------------------|-------------------------|
| **Gaspard et al. (1984)** | Comparative, parallel group     | Healthy women                     | No OC: ≥8 weeks | 22.75 years, NR     | 6 cycles 30 EE/150 DSG 21:7                                               | Extraction/chromatography RIA Intra-assay CV: NR Inter-assay CV: NR LLQ: NR | AFTC: calculated based on total T, SHBG and albumin (Vermeulen et al., 1971) |
|                     |                                 |                                   |             |                     |                                                                             |                                                                            |                         |
| **Hammond et al. (1984)** | Comparative, parallel group     | Healthy women                     | 20–36 years, Regular menstrual cycle | 24.4 ± 1.6 years, 57.6 ± 1.6 kg | 9 cycles 50 EE/250 LNG 21:7                                               | Control cycle (7 days before presumed menses) Tr. cycle 6 (Days 19–21) | Direct RIA Intra-assay CV: NR Inter-assay CV: NR LLQ: NR Not applicable |
|                     |                                 |                                   |             |                     |                                                                             |                                                                            |                         |
| **Siegberg et al. (1984)** | Open-label, non-comparative     | Adolescent girls                  | 15–20 years, No hormonal contraceptives: ≥12 months | 17.0 ± 1.5 years, 54.7 ± 4.6 kg | 6 cycles 50 EE/1 mg lynestrenol OR 35 EE/0.8 mg lynestrenol 21:7            | Control cycle (≥Day 12; 1 × / week) Tr. cycle 6 (2 times) | RIA Intra-assay CV: 6% Inter-assay CV: 12% LLQ: NR |
|                     |                                 |                                   |             |                     |                                                                             |                                                                            |                         |
| **Kuhl et al. (1985)** | Randomized, crossover, comparative, parallel group | Healthy women                     | 24–35 years, No OC: ≥3 months | 22 Mean age: NR, 54.7 ± 4.6 kg | 9 cycles 30-40-30 EE/50-75-125 LNG (6-5-10), 21:7                          | Control cycle (Day 21) Tr. cycle 3 (Day 21) | Direct RIA Intra-assay CV: 7.2% Inter-assay CV: 8.1% LLQ: 0.87 nmol/l |
|                     |                                 |                                   |             |                     |                                                                             |                                                                            |                         |

*Continued*
### Table II  Continued

| First author, year | Study design | Study population | Characteristics per treatment group | Study treatment (duration, composition and regimen) | End-points used in meta-analysis | Details hormone assessment (type of assay or calculation) | Domains with low risk of bias (n) |
|--------------------|--------------|------------------|-------------------------------------|---------------------------------------------------|-------------------------------|------------------------------------------------------|---------------------------------|
| Falsetti et al. (1987) | Open-label, non-comparative, non-controlled | Healthy women menstrual cycle 22–38 years No hormonal contraceptives: ≥ 3 months | 1 n = 39 Mean (SD) age: 30.1 ± 4.5 years Mean (SD) weight: 56.4 ± 7.7 kg | 6 cycles 20 EE/150 DSG 21:7 | Control cycle (7 days before presumed menses) Tr. cycle 6 (day NR) | Direct RIA Intra-assay CV: NR Inter-assay CV: NR LLQ: NR | 7 |
| Jung-Hoffmann et al. (1988a) | Randomized, comparative, parallel group | Healthy women 18–35 years Regular menstrual cycle No OC: ≥ 3 months Control cycle: ovulation confirmed | 1 n = 11 Mean age: NR Weight/BMI: NR | 12 cycles 30 EE/75 GSD 21:7 | Control cycle (Day 21) Tr. cycle 6 (Day 21) | Direct RIA Intra-assay CV: NR Inter-assay CV: NR LLQ: NR | 4 |
| Murphy et al. (1990) | Prospective, non-comparative | Healthy women 18–35 years normal menstrual cycle No hormonal contraceptives: ≥ 3 months | 1 n = 33 Mean age: NR Weight/BMI: NR | 9 cycles 35 EE/1000 NET 21:7 | Control cycle (Days 1–5) Tr. cycle 6 (Days 1–5) | Direct RIA Intra-assay CV: 1.5% Inter-assay CV: NR LLQ: 11 ng/dl (0.38 nmol/l)§ | 7 |
| Refn et al. (1990) | Randomized, comparative, parallel group | Healthy women 19–36 years Regular menstrual cycle No hormonal contraceptives: ≥ 3 months | 1 n = 17 Mean age: 24.7 years Weight/BMI: NR | 6 cycles 30–40–30 EE/50–75–125 LNG (6–5–10), 21:7 | Control cycle (luteal phase, Days 24–26) Tr. cycle 6 (Days 18–20) | Direct RIA Intra-assay CV: 6% Inter-assay CV: 10% LLQ: NR | Not applicable 5 |
| De Leo et al. (1991) | Open-label, non-comparative, non-controlled | Healthy women 17–25 years No hormonal contraceptives: not specified, but pretreatment samples were collected during follicular phase, so assumption no OC prior | 1 n = 10 Mean age: NR Weight/BMI: NR | 6 cycles 20 EE/150 DSG 21:7 | Control cycle (Days 8–10) Tr. cycle 6 (Days 16–19) | Not applicable RIA Intra-assay CV: NR Inter-assay CV: NR LLQ: NR | 7 |

**Notes:**
- NR: Not reported.
- §: Conversion factor.
- U: Unknown.
| Study          | Design                | Participants                                      | Menstrual Cycle Overview                                                                 | Mean (SD) Age | Mean (SD) Weight | Control Cycle | Tr. Cycle(s) | Direct RIA        | Intra-assay CV (%) | Inter-assay CV (%) | LLQ (ng/mL) | Equilibrium Dialysis | Intra-assay CV (%) | Inter-assay CV (%) | LLQ (ng/mL) |
|---------------|-----------------------|---------------------------------------------------|----------------------------------------------------------------------------------------|----------------|--------------------|---------------|---------------|-------------------|---------------------|--------------------|-------------|---------------------|---------------------|--------------------|-------------|
| Kuhnz et al. (1991) | Open-label, non-comparative | Healthy women 19–34 years normal menstrual cycle No hormonal contraceptives: ≥ 2 months | 1 cycle 30-40-30 EE/50-70-100 GSD (6-5-10), 21:7                                           | 31.7 ± 6.0 years | 59.9 ± 4.5 kg     | Control cycle (Day 21) | Tr. cycle 1 (Day 21) | Direct RIA | Intra-assay CV: NR | Inter-assay CV: < 10% | LLQ: 0.1 ng/mL | (0.3 nmol/l) | Direct RIA | Intra-assay CV: NR | Inter-assay CV: < 10% | LLQ: 0.2 pg/ml | (0.7 pmol/l) |
| Janaud et al. (1992) | Randomized, comparative, parallel group | Healthy women 18–38 years Regular menstrual cycle No hormonal contraceptives: not specified, but only women with regular menstrual cycle were included | 6 cycles 35 EE/180-215-250 NGM 21:7                                                     | 24.7 ± 4.6 years | 55.6 ± 6.5 kg     | Control cycle (follicular phase) | Tr. cycle 6 (Days 1–6 and Days 14–28) | Not reported | Intra-assay CV: NR | Inter-assay CV: NR | LLQ: NR | Equilibrium dialysis | Intra-assay CV: NR | Inter-assay CV: NR | LLQ: NR |
| Song et al. (1992) | Randomized, comparative, parallel group, crossover | Healthy women Regular menstrual cycle No hormonal contraceptives: ≥ 3 months | 3 cycles 30 EE/150 DSG 21:7                                                            | 32.2 ± 2.4 years | 52.8 ± 4.8 kg     | Control cycle (follicular phase, Days 6–9; luteal phase, Days 21–24) | Tr. cycle 3 (Day 21) | RIA | Intra-assay CV: < 10% | Inter-assay CV: < 10% | LLQ: NR | Not applicable | 4 |
|                |                       |                                                   | 2 cycles 20 EE/150 DSG 21:7                                                            | 32.2 ± 2.4 years | 52.8 ± 4.8 kg     |                                                               |               |               |                   |                     |                   |             |                     |                     |                   |             |
|                |                       |                                                   | 3 cycles 30 EE/150 LNG 21:7                                                            | 32.2 ± 2.4 years | 52.8 ± 4.8 kg     |                                                               |               |               |                   |                     |                   |             |                     |                     |                   |             |

Continued
| First author, year | Study design                      | Study population                          | Characteristics per treatment group | Study treatment (duration, composition and regimen) | End-points used in meta-analysis | Details hormone assessment (type of assay or calculation) | Domains with low risk of bias (n)^1 |
|-------------------|-----------------------------------|-------------------------------------------|--------------------------------------|-----------------------------------------------------|----------------------------------|----------------------------------------------------------|-----------------------------|
| Kuhl et al. (1993) | Non-comparative                   | Healthy women 20–29 years Regular menstrual cycle No hormonal contraceptives: ≥ 3 months Control cycle: ovulation confirmed | 1 n = 19 Mean age: NR Weight/BMI: NR | 6 cycles 40–30 EE/25–125 DSG (7–15), 21.7 | Control cycle (Days 21–24) Tr. cycle 6 (Days 18–22) | Direct RIA Intra-assay CV: 7.6% Inter-assay CV: 8.2% LLQ: NR | 7                          |
| Kuhnz et al. (1993a) | Open-label, non-comparative       | Healthy women 18–32 years No OC: ≥ 2 months | 1 n = 14 Mean (SD) age: 27 ± 5 years Mean (SD) weight: 66 ± 6 kg | 3 cycle 30–40–30 EE/50–70–100 GSD (6–5–10), 21.7 | Control cycle (day NR) Tr. cycle 3 (Day 21) | Direct RIA Intra-assay CV: 8% Inter-assay CV: 8% LLQ: NR | 7                          |
| Kuhnz et al. (1993b) | Open-label, non-comparative       | Healthy women normal menstrual cycle No hormonal contraceptives: ≥ 2 months | 1 n = 15 Mean (SD) age: 26 ± 6 years Mean (SD) weight: 63 ± 7 kg | 3 cycles 35 EE/2 mg CPA 21.7 | Control cycle (day NR) Tr. cycle 3 (Day 21) | Direct RIA Intra-assay CV: NR Inter-assay CV: 8–12% LLQ: NR | 7                          |
| Åkerlund et al. (1994) | Randomized, double-blind, comparative, parallel group | Healthy women 18–40 years No OC: ≥ 2 months | 1 n = 25 Mean age: 25.0 years Weight/BMI: NR | 12 cycles 20 EE/150 DSG 21.7 | Control cycle (Days 18–21) Tr. cycle 6 (Days 18–21) | Direct RIA CV: 6.5–8.9% at 3 nmol/l Intra-assay CV: 5–9% LLQ: NR | 6                          |
| Kuhnz et al. (1994) | Open-label, non-comparative       | Healthy women No OC: ≥ 2 months            | 1 n = 14 Mean (SD) age: 23 ± 3 years Mean (SD) weight: 61 ± 8 kg | 3 cycles 30–40–30 EE/50–75–125 LNG (6–5–10), 21.7 | Control cycle (day NR) Tr. cycle 3 (Day 21) | Direct RIA Intra-assay CV: < 10% Inter-assay CV: 4–8% LLQ: NR | 7                          |
| Study                     | Design                     | Participants                           | Methodology                                                                 | Cycle Duration | Control Cycle | Testosterone Levels | References |
|--------------------------|----------------------------|----------------------------------------|----------------------------------------------------------------------------|----------------|-----------------|---------------------|------------|
| Volpe et al. (1994)²     | Open-label, non-comparative | Healthy women 17–35 years normal menstrual cycle No hormonal contraceptives: ≥ 3 months | 1 n = 30 Mean (SD) age: 23.2 ± 5.21 years Weight/BMI: NR                      | 9 cycles       | Control cycle   | Direct RIA Intra-assay CV: 6.5% Inter-assay CV: 8.4% LLQ: 0.1 ng/mL (0.3 nmol/l) | 7          |
|                          |                            |                                        | 40-25 EE/30-125 DSG (7–15), 21:7                                          |                | (Day 22)        |                     |            |
| Heuner et al. (1995)     | Open-label, non-comparative | Healthy women No OC: ≥ 2 months        | 1 n = 14 Mean (SD) age: 27.9 ± 3.9 years Mean (SD) weight: 65.5 ± 11.8 kg | 3 cycles       | Control cycle   | Direct RIA Intra-assay CV: <10% Inter-assay CV: 4–8% LLQ: NR | 7          |
|                          |                            |                                        | 20 EE/75 GSD 21:7                                                           |                | (Day 21)        |                     |            |
|                          |                            |                                        |                                                                           |                | Tr. cycle 6     | Direct RIA Intra-assay CV: 5% Inter-assay CV: NR LLQ: 4 ng/dl (0.1 nmol/l) | 7          |
|                          |                            |                                        |                                                                           |                | (Day 22)        |                     |            |
|                          |                            |                                        |                                                                           |                | Tr. cycle 6     | Not applicable     |            |
|                          |                            |                                        |                                                                           |                | (Day 21)        |                     |            |
| Moutos et al. (1995)     | Randomized, comparative, parallel group | Healthy women 18–35 years Regular menses (every 25–35 days) No OC: ≥ 12 months | 1 n = 20 Mean age: 25.8 years BMI: 21.7 kg/m²                                 | 6 cycles       | Control cycle   | Direct RIA Intra-assay CV: 5% Inter-assay CV: NR LLQ: 4 ng/dl (0.1 nmol/l) | 7          |
|                          |                            |                                        | 50 EE/1 mg NET 21:7                                                         |                | (Day 21)        |                     |            |
|                          |                            |                                        |                                                                           |                | Tr. cycle 6     | Direct RIA Intra-assay CV: <10% Inter-assay CV: 7–8% LLQ: NR | 7          |
|                          |                            |                                        |                                                                           |                | (Day 21)        |                     |            |
|                          |                            |                                        |                                                                           |                | Tr. cycle 6     | Not applicable     |            |
|                          |                            |                                        |                                                                           |                | (Day 21)        |                     |            |
|                          |                            |                                        |                                                                           |                | Tr. cycle 6     |                     |            |
|                          |                            |                                        |                                                                           |                | (Day 21)        |                     |            |
|                          |                            |                                        |                                                                           |                | Tr. cycle 6     |                     |            |
|                          |                            |                                        |                                                                           |                | (Day 21)        |                     |            |
| Wiegratz et al. (1995)   | Comparative, parallel group | Healthy women 18–36 years Regular menstrual cycle No hormonal contraceptives: ≥ 3 months Control cycle: ovulation confirmed | 1 n = 26 Mean age: NR Weight/BMI: NR                                          | 12 cycles      | Control cycle   | Direct RIA Intra-assay CV: 7.6% Inter-assay CV: 8.2% LLQ: 0.08 ng/mL (0.3 nmol/l) | 5          |
|                          |                            |                                        | 30-40-30 EE/50-70-100 GSD (6-5-10), 21:7                                    |                | (Day 21)        |                     |            |
|                          |                            |                                        |                                                                           |                | Tr. cycle 6     | Direct RIA Intra-assay CV: 3.8% Inter-assay CV: 4.2% LLQ: 0.15 pg/ml (0.5 pmol/l) | 5          |
|                          |                            |                                        |                                                                           |                | (Day 21)        |                     |            |
|                          |                            |                                        |                                                                           |                | Tr. cycle 6     | Direct RIA Intra-assay CV: 3.8% Inter-assay CV: 4.2% LLQ: 0.15 pg/ml (0.5 pmol/l) | 5          |

*Continued*
| First author, year | Study design | Study population | Characteristics per treatment group | Study treatment (duration, composition and regimen) | End-points used in meta-analysis | Details hormone assessment (type of assay or calculation) | Domains with low risk of bias (n) |
|-------------------|--------------|------------------|-----------------------------------|---------------------------------------------|---------------------------------|------------------------------------------------|------------------|
| Coenen et al. (1996) | Randomized, open-label, comparative, parallel group | Healthy women 18–38 years Regular menstrual cycle No hormonal contraceptives: ≥ 2 months | 1 n = 25 Mean (SD) age: 26.9 ± 4.2 years Mean BMI: 22.2 ± 2.3 kg/m² | 6 cycles 35 EE/250 NGM 21:7 | Control cycle (luteal phase, Days 24–27) Tr. cycle 6 (Days 18–21) | Extraction/chromatography RIA Intra-assay CV: 5.6% Inter-assay CV: 6.9% LLQ: NR | 5 |
| | | | 2 n = 25 Mean (SD) age: 26.3 ± 4.9 years Mean (SD) weight: 22.2 ± 2.4 kg | 6 cycles 30 EE/75 GSD 21:7 | | Direct RIA Intra-assay CV: 6.4% Inter-assay CV: 12.1% LLQ: NR | |
| | | | 3 n = 25 Mean (SD) age: 27.1 ± 5.1 years Mean (SD) weight: 22.2 ± 2.4 kg | 6 cycles 30 EE/150 DSG 21:7 | | | |
| | | | 4 n = 25 Mean (SD) age: 27.3 ± 5.3 years Mean (SD) weight: 22.4 ± 2.2 kg | 6 cycles 20 EE/150 DSG 21:7 | | | |
| Study | Design | Participants | Menstrual History | Contraceptives | Sample Size | Age (Mean ± SD) | Weight (Mean ± SD) | Study Design | Cycle Details | Assay Method | Intra-assay CV | Inter-assay CV | LLQ | Comments |
|-------|--------|--------------|------------------|-----------------|-------------|----------------|------------------|--------------|--------------|--------------|----------------|----------------|------|----------|
| Spona et al. (1996) | Non-comparative | Healthy women | 20–34 years | normal menstrual cycle | No hormonal contraceptives: not specified, but women had to have a normal menstrual cycle | 24 | 27.5 ± 4.3 years | NR | Direct RIA | Intra-assay CV: 5–9% | Inter-assay CV: 5–9% | LLQ: NR | 7 |
| Aden et al. (1998) | Randomized, crossover, comparative | Healthy women | 21–32 years | Regular menstrual cycle | No OC: ≥ 2 months | Control cycle: ovulation confirmed | 29 | NR | Direct RIA | Intra-assay CV: 5.1% | Inter-assay CV: 10.4% | LLQ: 0.2 ng/ml (0.7 nmol/l) | 7 |
| Thorneycroft et al. (1999) | Randomized, open-label, comparative, parallel group | Healthy women | 18–28 years | Regular menstrual cycle | No hormonal contraceptives: ≥ 3 months | Control cycle | 27 | 22.9 ± 2.8 years | 74.9 ± 25.0 kg | Extraction/chromatography RIA | Intra-assay CV: 5–10% | Inter-assay CV: 10–15% | LLQ: NR | 5 |
| Boyd et al. (2001) | Non-randomized, open-label, non-comparative | Healthy women | 20–39 years | Regular menstrual cycle | No hormonal contraceptives: ≥ 6 months (implants/OC) or ≥ 12 months (injectables) | 17 | 52.6–141 kg | NR | Extraction/chromatography RIA | Intra-assay CV (%RSD): < 6.86% | Inter-assay CV (%RSD): < 10.9% | LLQ: 0.1% free | 5 |

Continued
| First author, year | Study design | Study population | Characteristics per treatment group | Study treatment (duration, composition and regimen) | End-points used in meta-analysis | Details hormone assessment (type of assay or calculation) | Domains with low risk of bias (n) |
|-------------------|--------------|-----------------|-----------------------------------|-------------------------------------------------|----------------------------------|-----------------------------------------------------|-----------------------------|
| Wiegratz et al. (2003) | Randomized, double-blind, comparative, parallel group | Healthy women 18–35 years Regular menstrual cycle No hormonal contraceptives: ≥ 4 weeks Control cycle: ovulation confirmed | | 1 n = 25 Mean age: NR Weight/BMI: NR 6 cycles 30 EE/2 mg DNG 21:7 | Control cycle (Days 21–26) Tr. cycle 6 (Days 18–21) | Not applicable | Direct RIA Intra-assay CV: 23.7% Inter-assay CV: 24.1% LLQ: 0.15 pg/ml (0.5 pmol/l) | 5 |
| | | | | 2 n = 25 Mean age: NR Weight/BMI: NR 6 cycles 20 EE/2 mg DNG 21:7 | | | | |
| | | | | 3 n = 25 Mean age: NR Weight/BMI: NR 6 cycles 10 EE/E2 V/2 mg DNG, 21:7 | | | | |
| | | | | 4 n = 25 Mean age: NR Weight/BMI: NR 6 cycles 20 EE/100 LNG 21:7 | | | | |
| Rickenlund et al. (2004)² | Open-label, non-comparative | Healthy women 16–35 years Regular menstrual cycle No hormonal contraceptives: not specified, but only women with regular menstrual cycles were included | | 1 n = 12 Mean (SD) age: 20.9 ± 4.2 years BMI: 19.9 ± 1.4 kg/m² 10 cycles 30 EE/150 LNG 21:7 | Control cycle (day NR) Tr. cycle 10 (day NR) | Direct RIA Intra-assay CV: 6% Inter-assay CV: 10% LLQ: 0.1 nmol/l | Calculated based on total T, SHBG and fixed albumin [40 g/l] (Södergård et al., 1982) | 7 |
| White et al. (2005)² | Randomized, comparative, parallel group | Healthy women 18–40 years regular menstrual cycle (25–35 days) No hormonal contraceptives: ≥ 3 months | | 1 n = 9 Mean (SD) age: 25.1 ± 2.8 years BMI: 21.8 ± 3.4 kg/m² 3 cycles 35 EE/250 NGM 21:7 | Control cycle (day NR) Tr. cycle 3 (Days 17–21) | Extraction/chromatography RIA Intra-assay CV: 4–7% Inter-assay CV: 9–12% LLQ: NR | Calculated based on total T, SHBG and albumin (Vermeulen et al., 1999) | 5 |
| Study                  | Design                        | Participants                                                                 | Follow-up | Measurement       | Intra-assay CV (%) | Inter-assay CV (%) | LLQ (nmol/l) | Notes                                                                 |
|-----------------------|-------------------------------|-----------------------------------------------------------------------------|-----------|-------------------|--------------------|--------------------|--------------|----------------------------------------------------------------------|
| Elkind-Hirsch et al.  | Randomized, prospective,    | Healthy women 18–40 years No hormonal contraceptives: ≥ 2 months control    | 1         | 5 cycles          | 10%                | 10%                | NR           | Not applicable                                                      |
|                       | comparative, parallel group   | cycle: ovulation confirmed                                                   |           | Control cycle (Days 2–5) | Tr. cycle 5 (Days 8–21) |                    |              |                                                                    |
| Greco et al. (2007)   | Randomized, single-blind      | Healthy women ≥ 8 years Regular menstrual cycle No OC: ≥ 3 months           | 1         | 3 cycles          | 10%                | 10%                | NR           | Method used is based on centrifugal ultrafiltration dialysis;       |
|                       | comparative, parallel group   |                                                                             |           | Control cycle (2 days around ovulation) | Tr. cycle 2 (Days 12–14) |                    |              | calculated from a normogram constructed using serum SHBG and %FT     |
|                       |                               |                                                                             |           |                   |                    |                    |              | (Hammond et al., 1980); free T can then be determined from T.        |
|                       |                               |                                                                             |           |                   |                    |                    |              |                                                                    |
| Legro et al. (2008)   | Randomized, double-blind      | Healthy women No OC: ≥ 3 months normal menstrual cycle (21–35 days) non-smoking | 2         | 6 cycles          | 10%                | 10%                | NR           | Not applicable                                                      |
|                       | comparative, parallel group   |                                                                             |           | Control cycle (Days 15–21) | Tr. cycle 6 (days 15–21) |                    |              |                                                                    |
|                       |                               |                                                                             |           |                   |                    |                    |              | Method used is based on centrifugal ultrafiltration dialysis;       |
|                       |                               |                                                                             |           |                   |                    |                    |              | calculated from a normogram constructed using serum SHBG and %FT     |
|                       |                               |                                                                             |           |                   |                    |                    |              | (Hammond et al., 1980); free T can then be determined from T.        |
|                       |                               |                                                                             |           |                   |                    |                    |              |                                                                    |
| Sänger et al. (2008)  | Randomized, prospective,     | Healthy women 18–40 years Regular menstrual cycle No hormonal contraceptives: ≥ 4 weeks | 1         | 12 cycles         | 12.3%              | 12.3%              | 0.07 nmol/l  | Direct RIA Inter-assay CV: 12.3% Inter-assay CV: 18.3% LLQ: 0.52 pmol/l |
|                       | comparative, parallel group   |                                                                             |           | Control cycle (Days 21–26) | Tr. cycle 4 (Days 19–21) |                    |              |                                                                    |
|                       |                               |                                                                             |           |                   |                    |                    |              | Direct RIA Intra-assay CV: 4.6% Inter-assay CV: 7.4% LLQ: 0.07 nmol/l |
|                       |                               |                                                                             |           |                   |                    |                    |              |                                                                    |
| Duijkers et al. (2010)| Randomized, open-label,      | Healthy women 18–35 years No hormonal contraceptives: not specified, but women using OC and only after return of spontaneous menses control cycle was started (= at least 2 cycles no OC) Control cycle: ovulation confirmed | 1         | 6 cycles          | 4.6%               | 4.6%               | NR           | Calculated based on total T, SHBG and albumin (Vermeulen et al., 1999) |
|                       | comparative, parallel group   |                                                                             |           | Control cycle (Days 5–7) | Tr. cycle 6 (Days 21) |                    |              |                                                                    |
|                       |                               |                                                                             |           |                   |                    |                    |              | Direct RIA Intra-assay CV: NR Inter-assay CV: 4.6% LLQ: NR          |
|                       |                               |                                                                             |           |                   |                    |                    |              |                                                                    |
| First author, year | Study design | Study population | Characteristics per treatment group | Study treatment (duration, composition and regimen) | End-points used in meta-analysis | Details hormone assessment (type of assay or calculation) | Domains with low risk of bias (n) |
|-------------------|--------------|------------------|-----------------------------------|---------------------------------------------------|---------------------------------|--------------------------------------------------------|--------------------------|
| Sorufaldi et al. (2010) | Randomized, prospective, open-label, comparative, parallel group | Healthy women, 18–40 years, No hormonal contraceptives: ≥3 months | n = 16  
Mean (SD) age: 22.9 ± 4.3 years  
Mean (SD) weight: 66.3 ± 10.4 kg | 6 cycles  
30 EE/3 DRSP 21:7 |  | CMIA  
Total precision: 3.1–8.0%  
LLQ: 0.28 nmol/l | 7 |
| Ågren et al. (2011) | Randomized, open-label, comparative, parallel group | Healthy women, 18–50 years, No OC: 6 weeks | n = 51  
Mean (SD) age: 28.7 ± 6.83 years  
Mean (SD) weight: 60.6 ± 3.0 kg | 6 cycles  
30 EE/150 LNG 21:7 | Control cycle  
(Days 2–4)  
Tr. cycle 6  
(Days 2–4) | CMIA  
Total precision: 3.1–8.0%  
LLQ: 0.28 nmol/l | 7 |
| | | | n = 50  
Mean (SD) age: 26.8 ± 5.89 years  
Mean (SD) weight: 62.5 ± 12.6 kg | 6 cycles  
20 EE/100 LNG 21:7 |  |  |  |
| | | | n = 60  
Mean (SD) age: 28.2 ± 8.2 years  
Mean (SD) weight: 62.8 ± 9.7 kg | 6 cycles  
1.5 mg E2/2.5 mg NOMAC, 24:4 | Control cycle  
(follicular phase, nd half)  
Tr. cycle 6  
(Days 15–21) | ECLIA  
Intra-assay CV: NR  
Inter-assay CV: NR  
LLQ: NR | Calculated based on total T, SHBG and albumin (Vermeulen et al., 1999) |
| | | | n = 58  
Mean (SD) age: 29.1 ± 7.8 years  
Mean (SD) weight: 61.7 ± 9.0 kg | 6 cycles  
30 EE/150 LNG 21:7 |  |  |  |
Testosterone levels during pill use

ELISA

Control cycle

Testosterone levels during pill use

Prospective, Healthy women

Caruso et al. (2011)

Control cycle

Prospective, open-label, non-comparative

Song et al. (2012)

Pilot, non-comparative

Boyd et al. (2005)

AC-FTC, apparent free testosterone concentration; BMI, body mass index; CPA, cryopreserved aliquots; CV, coefficients of variation; DNG, dienogest; DRSP, drospirenone; DSG, desogestrel; E2, estradiol; E2 V, estradiol valerate; (E)CLIA, (electro)chemiluminescence immunoassay; EE, ethinyl estradiol; ELISA, enzyme-linked immunosorbent assay; GSD, gestodene; LLQ, lower limit of quantification; (L)NG, (levo)norgestrel; OC, oral contraceptive; N, number; N/A, not applicable; T, testosterone; Tr., treatment.

allocation concealment (high risk of bias). In many studies the procedures were not clearly described and no judgement on risk of bias could be made (Cullberg et al., 1982; Gaspard et al., 1983; Kuhl et al., 1985; Jung-Hoffmann et al., 1988a, b; Refin et al., 1990; Janaeu et al., 1992; Song et al., 1992; Wiegartz et al., 1995; Coenen et al., 1996; Thorneycroft et al., 1999; Wiegartz et al., 2003; White et al., 2005). With regard to performance and detection bias, it was assumed that lack of blinding of participants and personnel or lack of blinding of outcome assessment were not likely to have had an influence on the measurement of the biochemical outcome parameters. Therefore, risk of performance and detection bias was judged as not relevant and was considered to be low for all studies. On the other hand, incomplete outcome data (attrition bias) was found to be important, as missing data could potentially have an impact on the observed effect size. Reasons for early discontinuation could be due to being lost to follow-up, or could be related to side effects associated with low androgen levels (e.g. emotionally labile, lack of sexual desire). For the judgement of attrition bias, it was assumed that a completion rate of ≥ 80% per treatment group had not affected the outcome. Ten studies were considered to have a high risk of attrition bias (Åkerlund et al., 1994; Thorneycroft et al., 1999; White et al., 2005; Elkind-Hirsch et al., 2007; Greco et al., 2007) or an uncertain risk (Granger et al., 1982; Vermeulen and Thiery, 1982; Sieberg et al., 1984; Jung-Hoffmann et al., 1988a, b; Boyd et al., 2001). The reporting bias (selective outcome reporting) was difficult to assess due to the lack of study protocols. For all studies it was verified whether the reported laboratory parameters were in adherence to the objectives of the study as specified in the paper. This was also done for the reported statistical significances. One study (Boyd et al., 2001) was judged as having a high risk of reporting bias, because total T results were not reported. Finally, all studies were also judged for other possible sources of bias. The only additional source for bias was the inadequate statistics used in a crossover study pooling the data for both treatment groups (Song et al., 1992).

Several other issues were identified that were not considered as a source for bias, but could still have affected the outcome of the analyses. In one study, only Chinese women were enrolled (Song et al., 1992). Some of the procedures performed during data extraction (e.g. obtaining values from figures, converting SEM to SD, converting to SI units) were not clearly described and no judgement on risk of bias could be made (Granger et al., 1982; Murphy et al., 1990; Kuhnz et al., 1994; Moutos et al., 1995; Legro et al., 2008). Furthermore, for some studies the reported units or measures for variation (SEM) did not seem correct in view of the expected ranges and assumptions were made for correction (Granger et al., 1982; Falsetti et al., 1987; White et al., 2005). Most of the studies did perform their sampling at the end of the pill cycle just prior to the pill-free period (e.g. on Days 19–21); however, in a few studies samples were collected during the first week (Murphy et al., 1990; Strufaldi et al., 2010; Caruso et al., 2011; Battaglia et al., 2012). One study (Moutos et al., 1995) retrospectively collected samples from a larger study. Although steroids are known to be stable when stored for a longer period, it could be possible that there is some in vitro effect during storage, which could result in small deviations of the results. Finally, in one study (Sieberg et al., 1984) two different COCs were used in one treatment group (50 μg EE/1 mg lynestrenol and 35 μg EE/0.8 mg lynestrenol), and this study was...
Excluded from the subgroup analyses between the different EE dosages. These issues are not considered as bias, but could be a source for heterogeneity.

**Effect of COC treatment on total testosterone, SHBG and free testosterone concentrations**

The results of the analyses including the different subgroup analyses can be found in Table III and Supplementary data, Table SII.

**Effect of COC treatment on total testosterone concentrations**

A total of 39 studies and 64 treatment groups were included in the meta-analysis for the effect of COC treatment on total T levels with data from 1405 women (no COC) versus 1336 women (COC). The pooled total T concentration at the end of treatment was significantly (P < 0.001) lower than the pooled total T concentration at pretreatment. No significant between-study heterogeneity was found (Table III and Fig. 2).

The suppressive effect of COCs on total T did not reveal a significant difference (P = 0.54) between the COCs containing 20–25 μg EE dose and the COCs containing 30–35 μg EE (Supplementary data, Table SII and Supplementary data, Fig. S1). The subgroup analysis between the different types of progestin (first generation versus second generation versus third generation versus fourth or unclassified generation) also revealed no significant difference (P = 0.20) between the groups for the effect on total T (Table III and Fig. 3). When a more detailed comparison was made combining the EE dose and the type of progestin, a significant difference was found between the subgroups (P = 0.02). In particular, the first and second generation COCs containing 20–25 μg EE showed less suppressive effects on total T than did the other subgroups (Supplementary data, Table SII and Supplementary data, Fig. S2). A separate analysis comparing the second and third generation monophasic COCs containing 30–35 μg EE revealed no differences between progestins on total T levels (Fig. 3 Supplementary data, Table SII, Figs S1, S2 and S3). For all these subgroup analyses, no significant heterogeneity was found (Table III and Supplementary data, Table SII).

For the comparison between the methods used to measure total T, lower differences in total T concentrations were reported with the RIA incorporating extraction or chromatography compared with the assays without this extra step. This difference was statistically significant (P = 0.003); however, a significant heterogeneity was also identified (Supplementary data, Table SII and Supplementary data, Fig. S4).

**Effect of COC treatment on SHBG concentrations**

The 29 studies and 47 treatment groups used for the meta-analysis for the COC effect on free T concentrations included 1046 women at pretreatment and 977 women at end of treatment. The levels of free T showed a large variation. After inspection of the data, it was decided to use a logarithmic scale for the free T concentrations with the relative change as the effect measure. At the end of COC use, the free T levels were significantly (P < 0.001) decreased. The average relative change was 0.39, which means that the free T levels were 61% lower compared with the levels before starting the COC (Table III and Fig. 6). No significant difference in treatment effect was found after performing subgroup analyses for comparison between the EE dosages and between the type of progestins or when combining the EE dosage with the type of progestin (Table III, Figs 6, 7, Supplementary data, Table SII, Figs S8 and S9). Significant heterogeneity was found for the main analysis, but not for the subgroup analyses (Table III and Supplementary data, Table SII).

To judge whether the methods to assess the free T levels had any effect on the size of the observed effect, a subgroup analysis between the different methods was performed. No significant difference (P = 0.27) in pooled free T concentration was found between the results obtained by direct RIA and with the methods reported to be more accurate (calculation or equilibrium dialysis). No significant between-study heterogeneity was found (Supplementary data, Table SII and Supplementary data, Fig. S10).

**Discussion**

This systematic review and meta-analysis was performed to evaluate the effect of COCs on concentrations of total T, free T and SHBG in healthy women. A total of 42 studies involving 1495 women were included in the current analyses. Both pooled total T and free T concentrations decreased significantly following COC use. All studies reported such an effect, except one (Janaud et al., 1992). The suppression of free T was twice as high compared with total T. The decline in total T and free T levels were both independent from the estrogen dose or progesterin type. SHBG levels increased significantly during COC use, as reported by all included studies. The lower dosage of EE had less of an effect on SHBG than the higher dosage, and the greatest effect was with the COCs containing a third or fourth generation progesterin. The COCs with a second...
Table III  Summary of key findings of the meta-analysis on the effect of COCs on total T, free T and SHBG concentrations with the comparison no COC (pretreatment) versus COC (end of treatment).

| Type of analysis                  | Number of studies | Number of treatment groups | Number of subjects Total (no COC versus COC) | MD [95% CI]                      | P value | Heterogeneity (I²) (%) |
|-----------------------------------|-------------------|----------------------------|----------------------------------------------|----------------------------------|---------|------------------------|
| COC effect on total T [nmol/l]    | 39                | 64                        | 2741 (1405 versus 1336)                      | 0.49 [−0.55, −0.43]              | <0.001  | 55                     |
| Subgroup: type of progestin      | 39                | 61                        | 2452 (1256 versus 1196)                      | 0.50 [−0.56, −0.43]              | 0.20    | 34.7                   |
|  First generation (estranes)     | 8                 | 31                        | 313 (159 versus 154)                         | −0.33 [−0.50, −0.17]             |         |                        |
|  Second generation (gonanes)     | 21                | 93                        | 934 (474 versus 460)                         | −0.54 [−0.63, −0.45]             |         |                        |
|  Third generation (gonanes)      | 28                | 104                       | 1044 (542 versus 502)                        | −0.51 [−0.62, −0.40]             |         |                        |
|  Fourth or unclassified generation | 4              | 16                        | 161 (81 versus 80)                           | −0.52 [−0.77, −0.27]             |         |                        |
| COC effect on SHBG [nmol/l]      | 39                | 67                        | 2917 (1487 versus 1430)                      | 99.10 [86.44, 111.75]            | <0.001  | 96                     |
| Subgroup: type of progestin      | 38                | 64                        | 2628 (1338 versus 1290)                      | 103.09 [89.51, 116.66]           | <0.001  | 97.8                   |
|  First generation (estranes)     | 9                 | 34                        | 347 (176 versus 171)                         | 89.06 [65.94, 112.19]            |         |                        |
|  Second generation (gonanes)     | 20                | 93                        | 934 (470 versus 464)                         | 35.84 [26.86, 44.82]             |         |                        |
|  Third generation (gonanes)      | 28                | 1036                      | 1036 (536 versus 500)                        | 136.76 [120.66, 152.86]          |         |                        |
|  Fourth or unclassified generation | 7              | 31                        | 311 (156 versus 155)                         | 157.39 [111.62, 203.16]          |         |                        |
| COC effect on free T (log scale; relative change) | 29                | 47                        | 2043 (997 versus 1046)                       | 0.39 [0.35, 0.43]                | <0.001  | 79                     |
| Subgroup: type of progestin      | 29                | 45                        | 1865 (954 versus 911)                        | 0.38 [0.35, 0.42]                | 0.32    | 12.0                   |
|  First generation (estranes)     | 2                 | 54                        | 54 (27 versus 27)                            | 0.38 [0.31, 0.48]                |         |                        |
|  Second generation (gonanes)     | 13                | 596                       | 596 (302 versus 294)                         | 0.42 [0.36, 0.50]                |         |                        |
|  Third generation (gonanes)      | 24                | 946                       | 946 (490 versus 456)                         | 0.37 [0.32, 0.43]                |         |                        |
|  Fourth or unclassified generation | 6              | 269                       | 269 (135 versus 134)                         | 0.34 [0.25, 0.46]                |         |                        |

CI, confidence interval; COC, combined oral contraceptive; MD, mean difference; SHBG, sex hormone-binding globulin; T, testosterone.
generation progestin caused the smallest increase in SHBG. The between-study heterogeneity was acceptable for all comparisons with total T and the subgroup comparisons with free T. However, a significant heterogeneity was found for the analyses with SHBG and for the main analysis with free T.

The between-study heterogeneity can be explained by the different types of COCs that were investigated in the included studies. Additionally, the duration without the use of a hormonal contraceptive prior to starting the COC varied from 1 month to 12 months. Another explanation may be the difference in handling of blood samples and assessment methods. Timing of blood sampling was not always consistent among studies. Most of the studies performed their sampling just prior to the start of the pill-free period, whereas a few studies (Murphy et al., 1990; Strufaldi et al., 2010; Caruso et al., 2011; Battaglia et al., 2012) collected samples during the first week of the cycle. Some studies did not provide information on the sampling time points, so it could be that no specific time window was followed. The sample timing could be an important factor when evaluating the effect of COC use on hormonal parameters. The fact that some studies measured the hormone concentrations in plasma and some in serum could also be a confounding factor, although this is mainly overcome by the within-subject comparison used in the current meta-analysis. Furthermore, there is no consensus on adequate methods for measuring total T as well as free T concentrations and its standardization (Vesper et al., 2008; Legro et al., 2010; Rosner

Figure 2 Meta-analysis of 39 studies on the effect of COCs on total T concentrations. COC, combined oral contraceptive; T, testosterone.
Figure 3 Subgroup analysis for the effect of type of progestin on total T concentrations in the meta-analysis. COC, combined oral contraceptive; T, testosterone.
and Vesper, 2010; Vesper and Botelho, 2010; Haring et al., 2012), which is apparent from the studies using different types of analytical methods. The subgroup analysis between the studies which used a direct immunoassay with the studies using RIA with extraction and chromatography revealed a significant difference. When an extraction and chromatography step was included, significantly lower differences in concentrations of total T were measured. Other studies have also reported the overestimation of concentrations by direct immunoassays (Rosner et al., 2007; Groenestege et al., 2012). As data were grouped per assay method, the between-study heterogeneity in the subgroups was lower compared with the main analysis. The subgroup analysis performed for the free T assessment did not identify a significant difference between concentrations obtained using direct immunoassay versus the calculation method or equilibrium dialysis. The latter two methods are regarded to be a more reliable measure of free T concentrations (Bachmann et al., 2002; Rinaldi et al., 2002; Miller et al., 2004), so this is an unexpected finding. This may be attributable to procedural differences, which, in the case of the calculation method or equilibrium

![Figure 4](meta-analysis.png)

Meta-analysis of 39 studies on the effect of COCs on SHBG concentrations. COC, combined oral contraceptive; SHBG, sex hormone-binding globulin.
Figure 5 Subgroup analysis for the effect of type of progestin on SHBG concentrations in the meta-analysis. COC, combined oral contraceptive; SHBG, sex hormone-binding globulin.
dialysis apparently differ more widely between laboratories than the RIA method. Other sources for the variation can be explained by the variations in the study population, e.g. different age and body mass index (BMI) ranges. No adjustment could be made for these confounding factors.

This meta-analysis has limitations which should be considered. The included studies were not strong in design with regard to preventing selection bias and only a few studies used a double-blind or single-blind design. However, due to the within-subject comparison, this bias seems to be of little relevance. Incomplete outcome data (attrition bias) seemed to be one of the most important factors affecting the analyses as some reasons for early discontinuation could be related to low androgen levels. The search was performed using terms (MeSH) in the title or abstracts, and therefore some publications could have been missed. However, this problem was partly covered by performing a hand search on the reference lists of all selected papers. For consistency

Figure 6 Meta-analysis of 29 studies on the effect of COCs on free T concentrations (log scale; relative change). COC, combined oral contraceptive; T, testosterone.
Figure 7 Subgroup analysis for the effect of type of progestin on free T concentrations in the meta-analysis on free T (log scale; relative change). T, testosterone.
reasons, only studies investigating COCs with a 21:7 or 24:4 regimen were eligible for inclusion. Only two of the included studies evaluated the effects of a COC with a 24:4 regimen (Duijkers et al., 2010; Ågren et al., 2011a, b). This COC containing E2 and nomegestrol acetate (NOMAC) was not included in the subgroup analyses, because it did not qualify for the subgroups. The effects of COCs with a continuous regimen were also not evaluated in the current meta-analysis. However, studies comparing COCs using a continuous regimen with COCs using the 21:7 regimen report that T and SHBG levels are affected to the same extent (Legro et al., 2008; Sänger et al., 2008). This shows that the maximal effect of a COC on T and SHBG is already reached within 3 weeks of use.

Based on the outcome of this review, it seems reasonable to conclude that the suppression of T is caused by all three mechanisms of action mentioned in the introduction of this paper, i.e. suppression of both ovarian and adrenal androgen synthesis, along with increased SHBG levels. The average decrease of total T was 31% and for free T it was 61%, which cannot solely be explained by direct inhibition of ovarian androgen synthesis. Under normal conditions, ~25% of T is produced by the ovaries, whereas the adrenals synthesize another 25% and the remaining 50% is derived from peripheral conversion (Longcope, 1986; Bachmann et al., 2002; Burger, 2002). Thus, the other two mechanisms of action, the increase in SHBG increase and the inhibition of adrenal androgen synthesis, seem to be relevant too. Possibly the peripheral conversion of T from (pre)androgens is diminished due to lower levels of (pre)androgens. This contention is supported by the fact that many studies have also found decreased levels of dihydrotestosterone, AD, DHEA, DHEA-S and androstenediol glucuronide after COC use (Gaspard et al., 1984; Wiegratz et al., 2003; Ågren et al., 2011a, b). AD is the major source of peripheral T production (Longcope, 1986; Burger, 2002). The lower levels of AD, DHEA and DHEA-S are possibly the result of suppression of the adrenal androgen synthesis (Carr et al., 1979; Klove et al., 1984; Coenen et al., 1996).

This review also confirms that SHBG significantly increases during COC use, an effect which varied depending on the dose of the EE component. COCs with the second generation progesterin combined with the lowest EE dose (20–25 μg) had the least effect on SHBG. A separate analysis between the monophasic COCs with a second or third generation progesterin, both containing 30–35 μg EE, revealed that the third generation progesterin had a stronger effect on SHBG. All data confirm that COCs containing a second generation progesterin increase SHBG considerably less compared with all other progestins (~50% versus 150–250% elevation). Since SHBG is the main carrier protein of T one would expect more T to be bound when the SHBG increases are higher (Anderson, 1974; Van der Vange et al., 1990; Wiegratz et al., 2003; Ågren et al., 2011a). However, all progesterin types reduce total and free T to the same extent and no significant difference in lowering T levels was found when comparing the 20–25 μg with the 30–35 μg EE dose except for a small but significant difference (P = 0.02) when the lower EE dose was combined with a first or second generation progesterin. Similar results have been found in other studies (Jung-Hoffmann and Kuhl, 1987; Heuner et al., 1995; Boyd et al., 2001; Sänger et al., 2008). Apparently, the moderate SHBG increase caused by second generation progesterins already exerts the maximal effect on additional binding of T and further lowering of free T levels.

Apart from estrogens and androgens, progestins can also bind to SHBG and albumin (Schindler et al., 2003). Competition with T or E2 for SHBG might decrease the circulating concentration of the respective progestin in the circulation and, concomitantly, increase the free T and E2 levels, thereby indirectly influencing the SHBG synthesis, be it on an as yet undetermined extent.

As mentioned above, T deficiency is thought to be associated with a broad range of undesired effects including diminished well-being and quality of life, mood changes (depression, irritation, moodiness), loss of energy, cognitive disturbances, interference with optimal sexual function, declining muscle mass and strength and lowering of bone mass and bone density (Bachmann et al., 2002; Traish et al., 2007). The clinical consequences of the suppressed T levels during COC use have so far gained little attention. Only a few studies have combined the investigation of the effects of COCs on androgens with the occurrence of side effects which could be attributed to suppressed androgen levels. These studies report conflicting results (Bancroft et al., 1991b; Schaffir, 2006; Graham et al., 2007; Greco et al., 2007; Strufaldi et al., 2010; Caruso et al., 2011; Battaglia et al., 2012; Elaut et al., 2012; Pastor et al., 2013). Although estrogens have been clearly identified as key hormones for maintaining bone mass (Ott et al., 2001; Riggs et al., 2002; Ott, 2008), androgens are also important regulators of skeletal growth and maturation (Martel et al., 1998; Riggs et al., 2002; Hartard et al., 2006). Data on the effect of androgens in women are scarce but T may exert an effect on protein synthesis and increase muscle mass (Gower and Nyman, 2000; Ružič et al., 2003). Therefore, the negative effect of COCs on T levels may have some impact on bone mass and also on muscle strength.

In conclusion, the current systematic literature review and meta-analysis demonstrates that COCs decrease circulating levels of total T and free T and increase SHBG concentrations. Due to the SHBG increase, free T levels decrease twice as much as total T. The estrogen dose and progesterin type of the COC do not influence the decline of total or free T, but both affect the magnitude of the effect on SHBG. The clinical implications of suppressed androgen levels during COC use remain to be elucidated. Further research investigating the COC’s effects on androgens combined with their effects on other clinical outcomes is warranted.

**Supplementary data**

Supplementary data are available at http://humupd.oxfordjournals.org/.

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**Authors’ roles**

Y.Z. made substantial contributions to the conception and design of the study, the acquisition of data, the analysis and interpretation of the data, the drafting of the article and the adjustments after the review by other authors and gave final approval of the submitted version of the manuscript. M.J.C.E. made substantial contributions to the conception and design of the study, the analysis and interpretation of the data, the critical review of the article and gave final approval of the submitted version of the manuscript. M.A.B. made substantial contributions to the
interpretation of the data, the critical review of the article and gave final approval of the submitted version of the manuscript. H.J.T.C.B. and B.C.J.M.F. made substantial contributions to the conception and design of the study, the interpretation of the data, the critical review of the article and gave final approval of the submitted version of the manuscript.

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Conflict of interest

Y.Z. is employed by PRB. H.J.T.C.B. is CEO of PRB and has equity interests in the company. M.J.C.E. and M.A.B. have nothing to declare. B.C.J.M.F. has received fees and grant support from the following companies (in alphabetic order): Andromed, Ardana, Auxogyn, Ferring, Genovum, Merck (MSD), Merck Serono, Organon, Pantheari Bioscience, Preglem, Schering, Schering Plough, Serono, Uteron Pharma, Watson Pharmaceuticals and Wyeth.

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