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Progestin-only contraceptives: effects on weight

Laureen M Lopez¹, Shanthi Ramesh², Mario Chen³, Alison Edelman⁴, Conrad Otterness⁵, James Trussell⁶, and Frans M Helmerhorst⁷

¹Clinical and Epidemiological Sciences, FHI 360, Durham, North Carolina, USA ²Department of Obstetrics and Gynaecology, University of North Carolina, Chapel Hill, NC - North Carolina, USA ³Biostatistics, FHI 360, Durham, North Carolina, USA ⁴Dept. of Obstetrics and Gynecology, Oregon Health & Science University, Portland, Oregon, USA ⁵Global Health Access Program, Mae Sot, Thailand ⁶Office of Population research, Princeton University, Princeton, New Jersey, USA ⁷Dept. of Clinical Epidemiology, Leiden University Medical Center, Leiden, Netherlands

Abstract

Background—Progestin-only contraceptives (POCs) are appropriate for many women who cannot or should not take estrogen. POCs include injectables, intrauterine contraception, implants, and oral contraceptives. Many POCs are long-acting, cost-effective methods of preventing pregnancy. However, concern about weight gain can deter the initiation of contraceptives and cause early discontinuation among users.

Objectives—The primary objective was to evaluate the association between progestin-only contraceptive use and changes in body weight.

Search methods—Until 4 August 2016, we searched MEDLINE, CENTRAL, POPLINE, LILACS, ClinicalTrials.gov, and ICTRP. For the initial review, we contacted investigators to identify other trials.

Selection criteria—We considered comparative studies that examined a POC versus another contraceptive method or no contraceptive. The primary outcome was mean change in body weight

DECLARATIONS OF INTEREST

Contact address: Laureen M Lopez, Clinical and Epidemiological Sciences, FHI 360, 359 Blackwell St, Suite 200, Durham, North Carolina, 27701, USA. llopez@fhi360.org.

CONTRIBUTIONS OF AUTHORS

^{2010:} L Lopez reviewed the search results, extracted and entered the data, and drafted the initial review. A Edelman and F Helmerhorst did part of the second data extraction. M Chen provided guidance on study design, data analysis, and interpretation of results. All authors reviewed and commented on the document.

^{2013:} C Otterness reviewed the search results, did the primary data extraction and entering, and incorporated the new study. L Lopez did the second data extraction and updated the text. M Chen provided guidance on study design, data analysis, and interpretation of results. All authors reviewed and commented on the document.

^{2016:} L Lopez ran the searches, revised the quality assessment for NRS, revised the text and tables to incorporate the new studies, and added the 'Summary of findings tables.' S Ramesh reviewed the search results, extracted and entered study characteristics, and helped update the text. L Lopez and S Ramesh entered and checked the outcome data for new studies. M Chen reviewed and contributed to the revised criteria for quality of evidence for all studies, and consulted on the presentation and interpretation of the outcome data. All authors reviewed and commented on the document.

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or mean change in body composition. We also considered the dichotomous outcome of loss or gain of a specified amount of weight.

Data collection and analysis—Two authors extracted the data. Non-randomized studies (NRS) need to control for confounding factors. We used adjusted measures for the primary effects in NRS or the results of matched analysis from paired samples. If the report did not provide adjusted measures for the primary analysis, we used unadjusted outcomes. For RCTs and NRS without adjusted measures, we computed the mean difference (MD) with 95% confidence interval (CI) for continuous variables. For dichotomous outcomes, we calculated the Mantel-Haenszel odds ratio (OR) with 95% CI.

Main results—We found 22 eligible studies that included a total of 11,450 women. With 6 NRS added to this update, the review includes 17 NRS and 5 RCTs. By contraceptive method, the review has 16 studies of depot medroxyprogesterone acetate (DMPA), 4 of levonorgestrel-releasing intrauterine contraception (LNG-IUC), 5 for implants, and 2 for progestin-only pills.

Comparison groups did not differ significantly for weight change or other body composition measure in 15 studies. Five studies with moderate or low quality evidence showed differences between study arms. Two studies of a six-rod implant also indicated some differences, but the evidence was low quality.

Three studies showed differences for DMPA users compared with women not using a hormonal method. In a retrospective study, weight gain (kg) was greater for DMPA versus copper (Cu) IUC in years one (MD 2.28, 95% CI 1.79 to 2.77), two (MD 2.71, 95% CI 2.12 to 3.30), and three (MD 3.17, 95% CI 2.51 to 3.83). A prospective study showed adolescents using DMPA had a greater increase in body fat (%) compared with a group not using a hormonal method (MD 11.00, 95% CI 2.64 to 19.36). The DMPA group also had a greater decrease in lean body mass (%) (MD –4.00, 95% CI –6.93 to –1.07). A more recent retrospective study reported greater mean increases with use of DMPA versus Cu IUC for weight (kg) at years 1 (1.3 vs 0.2), 4 (3.5 vs 1.9), and 10 (6.6 vs 4.9).

Two studies reported a greater mean increase in body fat mass (%) for POC users versus women not using a hormonal method. The method was LNG-IUC in two studies (reported means 2.5 versus -1.3; P = 0.029); (MD 1.60, 95% CI 0.45 to 2.75). One also studied a desogestrel-containing pill (MD 3.30, 95% CI 2.08 to 4.52). Both studies showed a greater decrease in lean body mass among POC users.

Authors' conclusions—We considered the overall quality of evidence to be low; more than half of the studies had low quality evidence. The main reasons for downgrading were lack of randomizations (NRS) and high loss to follow-up or early discontinuation.

These 22 studies showed limited evidence of change in weight or body composition with use of POCs. Mean weight gain at 6 or 12 months was less than 2 kg (4.4 lb) for most studies. Those with multiyear data showed mean weight change was approximately twice as much at two to four years than at one year, but generally the study groups did not differ significantly. Appropriate counseling about typical weight gain may help reduce discontinuation of contraceptives due to perceptions of weight gain.

Body Composition [drug effects]; Body Weight [*drug effects]; Contraceptives, Oral, Hormonal [*pharmacology]; Levonorgestrel [*pharmacology]; Medroxyprogesterone Acetate [*pharmacology]; Progestins [*pharmacology]; Weight Gain [drug effects]

MeSH check words

Female; Humans

PLAIN LANGUAGE SUMMARY

Effects of progestin-only birth control on weight

Progestin-only contraceptives (POCs) can be used by women who cannot or should not take the hormone estrogen. Many POCs are long acting, cost less than some other methods, and work well to prevent pregnancy. Some people worry that weight gain is a side effect of these birth control methods. Concern about weight gain can keep women from using these methods. Further, some women may stop using birth control early, which can lead to unplanned pregnancy. We looked at studies of POCs and changes in body weight.

Until 4 August 2016, we did computer searches for studies of a POC compared with another birth control method or no contraceptive. For the initial review, we wrote to investigators to find other trials. The focus was on change in body weight or other body measure of lean or fat mass.

With six new studies in this update, we have 22 studies that included 11,450 women. The groups compared did not differ much for weight change or other body measures in 15 studies. Five studies with moderate or low quality results showed a difference between study groups. Three studies showed differences for users of the injectable 'depo' versus no hormonal method. Depo users had a greater weight gain in two studies. In the third study, adolescents had a greater increase in body fat (%) and decrease in lean body mass (%). Two studies showed a greater increase in body fat (%) for users of hormonal intrauterine contraception versus women not using a hormonal method. One also showed a similar difference with a progestin-only pill. Both studies showed a greater decrease in lean body mass with POC use.

We found little evidence of weight gain when using POCs. Mean weight gain at 6 or 12 months was less than 2 kg (4.4 lb) for most studies. The groups using other birth control methods had about the same weight gain. Good counseling about typical weight gain may help women continue using birth control.

BACKGROUND

Description of the condition

Many women consider weight gain a side effect of using hormonal contraceptives (Bartz 2011; Raymond 2011). This perception may be based on self-report of side effects rather

than actual weight changes (Paul 1997; Berenson 2008; Nault 2013). Many clinicians and women believe that progestin-only contraceptives cause weight gain (WebMD 2010; Albright 2015).

Concern about weight gain can deter the initiation of contraceptives and cause early discontinuation among users. In a United States (US) study of bone mineral density, weight gain was reported more often by women using depot medroxyprogesterone acetate (DMPA) than those using a low-dose oral contraceptive (Berenson 2008). Weight gain was the most common side effect reported with DMPA use, after menstrual disturbances, in a New Zealand study (Paul 1997), and was the most common self-reported side effect in a study from Iran (Veisi 2013). Reported weight gain has been a major reason for discontinuing DMPA use in the US (Bonny 2004). Some evidence suggests that DMPA is a concern for adolescents who are already obese (Curtis 2009). From a survey of Latin American women across four countries, more women believed levonorgestrel-releasing intrauterine contraception (LNG-IUC) led to weight gain, mood swings, and infertility compared with copper IUC (Silva-Filho 2016). In a US study, more women reported weight gain as a side effect for the etonogestrel implant than for LNG-IUC (Dickerson 2013). Weight gain was also reported for levonorgestrel implants (Sivin 2003). The gain may have been greater among women in the US than among those in China, and may be partly attributable to differences in dietary habits.

Description of the intervention

Progestin-only contraceptives (POCs) include injectables, implants, hormonal intrauterine contraception (IUC), and pills. Except for the pills, POCs are longer-acting and help free women from daily action to prevent unintended pregnancy. Such methods are among the most cost-effective contraceptives in many areas. Studies of long-acting methods are often of longer duration than those for pills, making study of weight change over time more feasible.

POCs do not contain estrogen, unlike combined hormonal contraceptives that have both progestin and estrogen. Therefore, POCs are appropriate for women who cannot or should not take estrogen (ACOG 2006). In Medical Eligibility Criteria, POCs are category 1 for women who are obese (body mass index (BMI) 30 kg/m²) (CDC 2012b; WHO 2015a; WHO 2015b). Category 1 is a condition with no restriction for use of the contraceptive method. For obese adolescents, DMPA is category 2 due to possible effects on bone mineral density. For category 2, method advantages generally outweigh the theoretical or proven risks. POCs are also category 1 for breastfeeding women who are at least six weeks postpartum. Combined hormonal contraceptives are category 3 for such women until six months postpartum (WHO 2015b). In the US, CHCs are considered category 2 by one month postpartum (CDC 2012b).

Worldwide, intrauterine contraception (IUC) is the most commonly used reversible method among women married or in union (UN 2015). In the US in 2012, IUC use was low compared with oral contraceptive use (Guttmacher 2015). However, use of long-acting reversible methods (LARC) increased to 11.6% from 8.5% in 2009 (Kavanaugh 2015). Nearly three-fourths of IUC users were using hormonal IUC. Worldwide, the method used most frequently after IUC is oral contraceptives, which include combined oral contraceptives

(COCs) and progestin-only pills (POPs). In the least developed countries, injectables are most commonly used, followed by oral contraceptives (UN 2015). Some injectable contraceptives contain both estrogen and progestin, while others like DMPA are progestin-only.

How the intervention might work

In general, weight gain is due to an increase in fluid retention, muscle mass, or fat deposition. Research on mechanisms for weight change include investigations related to contraceptive use. Two uncontrolled studies included DMPA initiators, 12 to 21 years old. In a six-month study with 43 DMPA users, weight increased 1.2 kg among African Americans, as did BMI and total body fat. However, appetite score decreased while on DMPA for both African American and white participants (Bonny 2004). A 12-month study with 45 young women looked for associations of reported dietary intake with body composition change (Lange 2015). BMI increased significantly over 12 months, i.e. 1.6 kg/m², but was not associated with total energy intake or macronutrient composition of the diet (carbohydrates, fat, or protein). The study lost 31% of participants.

Other experimental studies included adult women. A nine-week study of resting metabolic rate (RMR) with DMPA initiation included 13 women with BMI 20 to 35 kg/m². RMR increased significantly during the first three weeks compared with the next six, especially for those who initiated during the luteal cycle (Steward 2016). An increase in non-shivering thermogenesis was consistent with the RMR change. A six-month metabolic study of P-O methods focused on 25 obese women (BMI 30) (Bender 2013). Participants chose the levonorgestrel-releasing intrauterine system (LNG-IUS), the etonogestrel-releasing (ETG) implant, or a non-hormonal method. Fasting glucose increased and insulin sensitivity decreased more with the ETG implant than with the LNG-IUS when compared to a nonhormonal method. An eight-week study examined DMPA effects on food motivation centers in the brain (Basu 2016). Eight of 14 women completed the protocol with data for analysis. All had BMI < 30 and most were Latina. The investigators used functional magnetic resonance imaging (MRI) to assess response to food cues. The blood oxygen level dependent signal was greater after eight weeks of DMPA compared with baseline. Some brain regions had significant activation after DMPA with food versus nonfood images and with high-calorie versus low-calorie food cues. Circulating leptin and ghrelin, hormones known to regulate eating behavior, did not change significantly. Such work may help elucidate mechanisms when conducted with larger sample sizes.

During adolescence, some weight gain is developmentally normal and appropriate. Also, people tend to gain weight over time (Flegal 2000). In the US, the prevalence of overweight or obesity is higher for men and women 40 to 59 years of age compared with those aged 20 to 39 years (Ogden 2014). In contraceptive studies, weight change is rarely a primary outcome in contraceptive studies. No consensus exists regarding what is excessive weight gain. Examining contraceptive use and weight gain can be complicated by the initial weight of the users. Recent interest in the effectiveness of hormonal contraceptives among obese women has led to more research with such women, who had been historically excluded from such studies (Bender 2013; Lopez 2013; Edelman 2014).

Concern about contraceptive effectiveness among overweight women (Robinson 2013; Merki-Feld 2015) has led to questions about contraceptive usage by overweight or obese women. An analysis of medical records from 231 health centers examined contraceptive use among 147,336 US women, age 15 to 44 years (Kohn 2015). The obese women (BMI 30) were more likely to use LARC than women with a BMI < 30 (13% versus 9%, respectively). Obese women were less likely than women with a lower BMI to use OCs, the injectable, the vaginal ring, or the subdermal patch (76% versus 82%, respectively). An analysis of US survey data examined contraceptive use in the past month (Callegari 2014). The women were sexually active, obese (BMI > 30), and age 20 to 44 years. LARC use among these obese women was nearly 10%, but only about 38% reported using OCs, the patch, the ring, or injectable contraception. While 21% reported not using a contraceptive method in the past month, 31% used a nonprescription method, i.e. condoms or another barrier method, withdrawal, or fertility awareness methods. Those two groups were more likely to be the youngest and oldest (aged 20 to 24 or 40 to 44). Women who used nonprescription methods were less likely to report having discussed contraception with a healthcare provider in the past year.

Why it is important to do this review

Prior to the initial review, no comprehensive systematic review existed on progestin-only contraceptives and weight change. Concern about weight gain might deter women from using these effective contraceptives and health care providers from recommending them. We did not examine effectiveness nor focus on overweight women. Many reviews have examined effectiveness of specific progestin-only contraceptives, such as progestin-only pills (Grimes 2013) and IUC (Grimes 2007). Further, a Cochrane review examined effectiveness of hormonal contraceptives for overweight women versus women who were not overweight (Lopez 2013).

Progestin-only contraceptives are an attractive option for many women. The longer-acting POCs, especially IUC and implants, are among the more effective methods with typical use (Trussell 2011). The cost for POCs can be less than that of COCs in some areas, and many postpartum women can use them. Further, POCs are appropriate for women at increased risk for venous thromboembolism such as those who are obese (Merki-Feld 2015), which is important given the worldwide epidemic of obesity (Prentice 2006; Flegal 2012; Ogden 2014). Being overweight or obese increases also risk for Type 2 diabetes and other diseases and disorders.

OBJECTIVES

The primary objective was to evaluate the potential association between progestin-only contraceptive use and changes in body weight.

METHODS

Criteria for considering studies for this review

Types of studies—We considered studies that examined progestin-only contraceptives used for contraception and their associations with weight change. Reports had to contain

information on the specific contraceptive method(s) examined. We searched for studies with comparative data on a progestin-only contraceptive versus another contraceptive (differing in formulation, dose, regimen, or initiation time) or no hormonal contraceptive. Potential studies included comparisons of a POC with a combination contraceptive as well as comparisons of two different types of progestin-only contraceptives.

Types of participants—Participants were the women in the studies who used the progestin only contraceptive for contraception or who had the comparison intervention or placebo. We did not consider studies focused on women with specific health problems, such as diabetes or HIV.

Types of interventions—We considered any progestin-only contraceptive, such as an oral contraceptive, an injectable, an implant, or hormonal intrauterine contraception (IUC). Treatment duration must have been at least three cycles or three months.

The progestin method of interest must have been specified and not combined in a group with another method, e.g. a group that used either DMPA or norethisterone enanthate (NET-EN). The comparison could have been another progestin-only contraceptive or a group of contraceptives, such as COCs. We did not include comparison groups identified only as oral contraceptive (OC) users, since the oral contraceptives could have been progestin-only pills or combined oral contraceptives.

The progestin-only method had to be intended for contraception. We did not consider studies of contraceptives used for treatment for specific disorders, e.g. acne, hirsutism, or polycystic ovary syndrome.

Types of outcome measures—The primary outcome was the continuous outcome of mean change in body weight, BMI, or body composition (e.g. percent body fat) over time with the use of progestin-only contraceptives. If mean change in body weight or BMI was not available per study arm, we examined the dichotomous outcome of loss or gain of a specified amount of weight in each study arm.

For high quality evidence, the study had to include mean change in body weight, BMI, or body composition. The time frame had to be 12 months.

We used measured weight and not self-reported weight. We excluded studies that did not report change data but only reported mean weight or BMI at pre- and post-treatment.

Search methods for identification of studies

Electronic searches—Until 4 August 2016, we searched MEDLINE, Cochrane Central Register of Controlled Trials (CENTRAL), POPLINE, Web of Science, and LILACS. We also searched for trials via ClinicalTrials.gov and the search portal of the International Clinical Trials Registry Platform (ICTRP). Appendix 1 shows the 2016 strategies. We listed the previous search strategies in Appendix 2.

Searching other resources—We examined reference lists of relevant articles. For the initial review, we contacted investigators in the field to seek additional unpublished trials or published trials that we may have missed in our search.

Data collection and analysis

Selection of studies—We assessed for inclusion all titles and abstracts identified during the literature searches. One author reviewed the search results and identified reports for inclusion or exclusion. A second author also examined the reports identified for appropriate categorization according to the eligibility criteria above.

We considered all comparative study designs. For example, studies could have been randomized controlled trials (RCTs), other prospective studies (provided intervention; assignment not random), observational studies of users, case-control studies, or retrospective chart reviews. We also considered post hoc analysis from any of these types of studies. However, the studies had to meet the Criteria for considering studies for this review.

Data extraction and management—Two authors conducted the data extraction. One author entered the data into Review Manager (RevMan 2014), and a second author checked accuracy. These data include the study characteristics, risk of bias, and outcomes. We focused on the primary and secondary outcomes for this review, which do not include all outcomes from each study. The authors resolved discrepancies through discussion.

Assessment of risk of bias in included studies—We examined the RCTs for methodological quality in accordance with recommended principles (Higgins 2011), and entered the information into the Risk of bias tables. Factors considered are randomization method, allocation concealment, blinding, and losses to follow-up and early discontinuation.

For the NRS, we used the Newcastle-Ottawa Quality Assessment Scale (NOS) (Higgins 2011; Wells 2014). Of the two NOS versions, i.e. for case-control and cohort studies, the latter was more pertinent here (Appendix 3). The NOS investigators are examining the criterion validity and construct validity of this scale as well as the inter-rater reliability and intra-rater reliability. The scale does not yet have an overall scoring or threshold for a 'good' or 'poor' quality study. The NOS has eight items within three domains: selection (representativeness), comparability (due to design or analysis), and outcomes (assessment and follow-up). A study can receive one star (*) for meeting each criterion. The exception is comparability (design or analysis), for which a study can receive two stars (for design and analysis). We adapted the NOS items for this project as suggested by the developers (Wells 2014).

Measures of treatment effect—Outcomes listed in Characteristics of included studies focus on those relevant to this review. We examined weight change in relation to initial body weight or body mass index (BMI) [weight (kg)/height (m)²] when we had the necessary data. Weight change may differ for women who were initially overweight or obese versus those who were not. We preferred BMI over weight alone, as BMI is a better reflection of body fat (CDC 2012a). The measures and cutoffs depended on those used in the included

studies. Frequently used BMI categories are 25 to 29.9 (kg/m^2) for overweight and 30 or higher for obesity (CDC 2012a).

We examined results by the contraceptive method studied, e.g. injectable or implant, as well as by formulation, dose, or regimen as appropriate. The main comparisons for this review were between users of progestin-only contraceptives and users of another contraceptive (differing in formulation, dose, or regimen) or no hormonal contraceptive.

For weight change measure with follow-up of less than one year, we selected the six-month assessment (if available) and the latest date. If multiple time points were reported up to one year, we used the 6- and 12-month data. If data were available for more than three years, we used one-year data, the midpoint, and the last measure.

Randomized trials: For continuous variables, we computed the mean difference (MD) with 95% confidence interval (CI). Review Manager uses the inverse variance approach. For the dichotomous outcomes, we calculated the Mantel-Haenszel odds ratio (OR) with 95% CI. An example is the proportion of women who gained or lost more than 2 kg. Fixed and random effects give the same result if no heterogeneity exists, as when a comparison includes only one study.

Non-randomized studies: Given the need to control for confounding factors in NRS, we used adjusted measures for the primary effect measures when available or the results of matched analysis from paired samples. Investigators may have used a variety of adjustment strategies. When presenting results, we note the confounding factors considered in the design or analysis. If the report did not provide adjusted measures for the primary analysis, we used unadjusted outcomes with the methods described above for use with RCTs.

Dealing with missing data—We excluded studies with insufficient data on weight or BMI for analysis in this review. Reports sometimes provided results in figures without specific numbers; others presented means without any variance estimate. We contacted investigators for other missing data and for clarifications if the studies were less than 10 years old or had a report within the past five years. Investigators are unlikely to have access to data from older studies. Many studies in the initial review were more than 10 years old.

Assessment of heterogeneity—We expected study populations, designs, and interventions to be heterogeneous. We described the clinical and methodological diversity (or heterogeneity) of the studies. We did not pool data from studies that had different contraceptive methods (e.g. DMPA or implants), different doses of the same method, or different criteria for reporting weight change. Therefore, we were not able to conduct meta-analysis due to the range of contraceptive methods examined and different reporting for weight change. Heterogeneity is not an issue when a comparison has a single study.

Data synthesis—To assess the quality of evidence and address confidence in the effect estimates, we applied principles from GRADE (Grades of Recommendation, Assessment, Development and Evaluation) (Higgins 2011; GRADE 2013). If meta-analysis is not viable because of varied interventions or outcome measures, a typical 'Summary of findings' table

is not feasible. Also, the criteria for quality assessment differ for NRS and RCTs. We provide 'Summary of findings' tables for the main results, although we did not conduct a formal GRADE assessment for all outcomes (GRADE 2013).

We based our assessment of the body of evidence on the quality of evidence from the studies. In 2016, we revised the Risk of bias tables to accommodate RCTs and NRS. For the NRS, we used the Newcastle-Ottawa Quality Assessment Scale as noted earlier (Appendix 3). Evidence quality included the design, implementation, and reporting of the study. We list the criteria for downgrading below.

- 1. Inadequate randomization sequence generation or allocation concealment, or no information provided for either one (RCT), or study was not randomized (NRS)
- 2. NRS: high risk of bias in selection (NOS) or retrospective study of selected cases
- **3.** NRS: no stars for comparability (NOS), i.e. not controlling for relevant confounding
- 4. Follow-up less than 12 months for change in weight or BMI
- 5. Losses (by one year or primary endpoint if more than one year): loss to follow-up greater than 20%, combined loss to follow-up and discontinuation greater than 50%, or differential losses between groups (greater than 50% difference)

For the initial review in 2010 and the minor update in 2013, we used a basic process to assess evidence quality. For those versions, the initial grade was based on study design: RCTs were considered to provide high quality evidence; prospective non-randomized studies, moderate quality; and retrospective studies, low quality. Those ratings were then downgraded for high loss to follow-up and inappropriate exclusions after randomization.

Sensitivity analysis—We examined separately the studies that provided evidence of moderate or high quality.

RESULTS

Description of studies

Results of the search—The 2013 search produced 189 citations: 123 references from the database searches, 63 trials from the clinical trials sites, and 3 references from other sources. After reviewing the full text, we included one new study for a total of 16 studies. We excluded five studies and two secondary articles related to previously excluded studies. The remaining references were discarded after reviewing the titles and abstracts. From the clinical trials sites, we added three new trials to Ongoing studies.

The 2016 search produced 138 unduplicated references from the database searches (Figure 1). With one item from another source, the total was 139. We discarded 122 citations based on title or abstract. After reviewing the full text of 17 articles, we excluded 9 reports (8

primary articles plus a secondary article). We included six new studies that involved six primary reports plus two secondary articles. Searches of recent clinical trials yielded 61 unduplicated listings. Two studies are completed but have not yet produced full reports (Studies awaiting classification). We will assess them for inclusion when full reports are available. Two other trials are Ongoing studies.

Included studies—With 6 new studies in this update, 22 studies now met our inclusion criteria. Fifteen were prospective and seven were retrospective.

- 15 prospective studies
 - 5 randomized controlled trials (WHO 1983; Salem 1988; Ball 1991; Sivin 1998; Westhoff 2007)
 - 10 non-randomized studies (NRS) (Tankeyoon 1976; Castle 1978; Salem 1984; Bonny 2009; Dal'Ava 2012; Nyirati 2013; Vickery 2013; Dal'Ava 2014; Dos Santos 2014; Napolitano 2015)
- 7 retrospective studies: NRS (Moore 1995; Taneepanichskul 1998; Espey 2000; Sule 2005; Tuchman 2005; Pantoja 2010; Modesto 2015)

The studies examined four categories of progestin-only contraceptives (Table 2).

- Oral contraceptives (OCs) containing norethisterone 350 μg, levonorgestrel 30 μg, or desogestrel 75 μg
 - Injectables
 - depot medroxyprogesterone acetate (DMPA): 150 mg/mL versus 450 mg/mL (intramuscular) or versus 104 mg/0.65mL (subcutaneous)
 - norethisterone enanthate (NET-EN) 200 mg
- Implants: levonorgestrel 6 capsules or 2 rods; etonogestrel 1 rod
- Levonorgestrel-releasing intrauterine contraception (LNG-IUC)

Comparison groups included no hormonal method or a non-hormonal contraceptive; a different formulation, regimen, or initiation time of the same POC; another POC; and a combined contraceptive or a supplement containing estrogen.

Studies were conducted in the USA, South America, Europe, Africa, and Asia; some were conducted on multiple continents. Publication dates covered nearly 50 years: five studies from 1976 to 1988; four from 1991 to 1998; five from 2000 to 2009; and eight from 2010 to 2015. Duration of prospective follow-up or retrospective data collection was six months to two years for 16 studies, while four studies gathered data for three to five years of use, and two studies collected retrospective data for 10 years of use.

The studies included a total of 11,450 women with an average of 520 and a median about 160. Six studies had fewer than 100 participants, seven had 100 to 222 participants, six

ranged from 400 to 1000 women, and three had more than 1000 women. A few had comparison groups not used in this review because they did not meet our inclusion criteria.

We were not able to examine weight change in relation to age. Earlier, we identified three studies focused on adolescents and young women. Bonny 2009 analyzed data from a larger study of hormonal contraceptives and bone mineral density. Moore 1995 and Tuchman 2005 were retrospective chart reviews. A certain amount of weight gain is part of normal development for adolescents. None of the newer studies focused on young women. Studies that included both adolescents and adult women did not provide outcome data for age subgroups.

Risk of bias in included studies

Figure 2 summarizes our assessments for the overall review. Table 1 shows how we rated each study, and Figure 3 illustrates our assessment for each study. Because we adapted the Risk of bias tables to accommodate criteria for NRS, some categories are not relevant to an RCT or an NRS. In those cases, we left the cell empty rather than state 'not applicable' to distinguish between 'unclear' and no assessment.

Allocation—Of the five RCTs, two reported the method of randomization and allocation concealment (Salem 1988; Sivin 1998). The other three had no information on randomization method or allocation concealment (WHO 1983; Ball 1991; Westhoff 2007). Of the 17 NRS, one did not meet the NOS selection criteria (Taneepanichskul 1998).

Blinding—Two of the five RCTs had information on blinding. Ball 1991 was reportedly "single-blind" without any specifics. For one trial used in Westhoff 2007, the evaluators were blinded. For most studies, blinding was not feasible due to differences in the contraceptive methods or to women having chosen their contraceptive method in the NRS.

Incomplete outcome data—Of 22 studies, 18 had high risk of bias related to incomplete outcome data. Nine had loss to follow-up or discontinuation greater than 50% (Castle 1978; WHO 1983; Salem 1988; Ball 1991; Sivin 1998; Westhoff 2007; Nyirati 2013; Dal'Ava 2014; Dos Santos 2014). In Tankeyoon 1976, loss differed substantially between groups. Seven retrospective studies may have selected charts for those with complete data and not accounted for losses (Moore 1995; Taneepanichskul 1998; Espey 2000; Sule 2005; Tuchman 2005; Pantoja 2010; Modesto 2015). Modesto 2015 also had differential losses across group that varied over time. Vickery 2013 recruited women who completed at least 11 months of use.

Selective reporting—After randomization, Taneepanichskul 1998 excluded women who developed a chronic disease or disorder during method use. This exclusion may have biased the results. Weight gain is associated with development of some diseases and disorders.

Other potential sources of bias

<u>NRS: comparability (NOS)</u>: Of 17 non-randomized studies, eight addressed potential confounding factors. Four considered confounding in the design by matching on age and

baseline BMI (Pantoja 2010; Dal'Ava 2012; Dos Santos 2014) or age and weight (Dal'Ava 2014). Four studies conducted analysis that adjusted for potential confounders (Moore 1995; Bonny 2009;Modesto 2015; Vickery 2013). In some cases, a comparison group did not meet our inclusion criteria, so we did not include that group in this review. Details are in Characteristics of included studies.

Effects of interventions

See: Summary of findings for the main comparison; Summary of findings 2; Summary of findings 3

We grouped results according to four types of progestin-only contraceptives studied, though some studies included more than one progestin-only (P-O) method. Table 2 summarizes the study interventions and outcomes, along with the mean changes in weight or other body composition measure. Two studies examined progestin-only pills, 15 addressed DMPA, 5 investigated implants, and four evaluated LNG-IUC. We subdivided the DMPA studies into those comparing DMPA with a combination contraceptive, another progestin-only injectable formulation or regimen, or no hormonal contraception.

Progestin-only oral contraceptives—Two studies examined P-O oral contraceptives. In the RCT of Ball 1991, weight change at six months did not differ significantly between the norethisterone 350 μ g and the levonorgestrel 30 μ g groups (Analysis 1.1). Mean changes were small. The NRS of Napolitano 2015 compared body composition changes at 12 months for perimenopausal women receiving desogestrel 75 μ g versus a control group with no hormonal treatment. The study also examined LNG-IUS. Changes in mean weight and BMI did not differ significantly between the OC group and the control group at 12 months (Analysis 2.1; Analysis 2.2). However, the desogestrel group had a greater increase in fat mass (%) (MD3.30, 95% CI 2.08 to 4.52) (Analysis 2.3).

Injectables

<u>DMPA versus a combination contraceptive:</u> Three NRS examined DMPA 150 mg/mL versus a contraceptive or supplement that also contained estrogen.

- In the small study of Tankeyoon 1976, the DMPA and COC groups were not significantly different in the proportions that gained (or lost) at least 1 kg by months 6 and 12 (Analysis 3.1 to Analysis 3.4).
- The retrospective study of Tuchman 2005 focused on adolescents and young women, age 12 to 21 years. At 6 and 12 months, weight changes were not significantly different between the DMPA group and the COC users (Analysis 3.5 to Analysis 3.8) or the group using medroxyprogesterone acetate (MPA) plus E₂C (Analysis 4.1 to Analysis 4.4).
- Bonny 2009 compared mean changes in total body fat (%) and lean body mass (%) at six months for DMPA 150 + placebo injection versus DMPA + estradiol cypionate 5 mg (E_2C). The study targeted adolescents, age 12 to 18 years. The DMPA group was not significantly different from the

DMPA plus E2C group for mean change in percent body fat or percent lean body mass (Analysis 5.1; Analysis 5.2).

DMPA versus another P-O injectable formulation or regimen: Of five studies in this group, three compared DMPA 150 mg/mL with other DMPA formulations or regimens. In Castle 1978, the mean changes in weight at six months were small and did not differ significantly between the DMPA 150 and DMPA 450 groups (Analysis 6.1). The retrospective study of Espey 2000 did not show a significant difference in weight gain at one or two years for those who initiated DMPA at 20 weeks or more after pregnancy (interval group) compared with those who initiated at 5 to 8 weeks (postpartum group) (Analysis 7.1; Analysis 7.2). In the RCT analyzed in Westhoff 2007, weight change was comparable for the group with intramuscular DMPA 150 and the group with subcutaneous DMPA 104 (Analysis 8.1).

Two RCTs examined DMPA 150 mg/mL versus NET-EN 200 mg. In WHO 1983, mean weight changes at 12 and 24 months did not differ significantly between the DMPA group and with the group administered NET-EN at 60-day intervals (Analysis 9.1; Analysis 9.2). Salem 1988 did not report the units for weight (lb or kg). However, the DMPA and the NET-EN groups did not differ significantly for mean changes in weight at one year (Analysis 9.3). Also in WHO 1983, two NET-EN regimens of 60 day-intervals versus 84-day intervals did not cause a significant difference in weight change (Analysis 10.1; Analysis 10.2).

DMPA versus no hormonal contraceptive: Eight NRS compared DMPA 150 mg/mL versus no hormonal method. Six had copper (Cu) IUC users as the comparison group; three studies were retrospective (Taneepanichskul 1998; Pantoja 2010; Modesto 2015) and three were prospective (Vickery 2013; Dal'Ava 2014; Dos Santos 2014). The remaining two prospective studies had other comparison groups (Bonny 2009;Nyirati 2013).

DMPA versus Cu IUC

Retrospective studies

- Taneepanichskul 1998 did not show a significant difference in weight change between the DMPA and Cu IUC groups at 10 years (Analysis 11.1). Unlike most studies in this review, the participants did not include younger women. All were 37 to 50 years old.
 - For Pantoja 2010, mean weight gain (kg) was greater for the DMPA group versus the Cu IUC group at years one (MD 2.28, 95% CI 1.79 to 2.77) (Analysis 11.2), two (MD 2.71, 95% CI 2.12 to 3.30) (Analysis 11.3), and three (MD 3.17, 95% CI 2.51 to 3.83) (Analysis 11.4). Per year, the mean weight changes for the DMPA group ranged from 1.76 kg to 3.9 kg, while changes within the IUC group were less than 1 kg (Analysis 11.5). For each year, the difference between contraceptive groups was notable within the normal to lower weight group (BMI < 25) and within the overweight group (BMI 25 to 29.9), but not within the obese group (BMI 30).

Modesto 2015 examined cumulative weight changes over 10 years of uninterrupted use of DMPA versus the Cu IUC. Using a generalized linear mixed model, the investigators adjusted for years of school and number of children. The adjusted analysis indicated mean weight change was significantly greater for the DMPA group versus the Cu IUC group for the three time points we used. The reported adjusted means were: at 1 year, 1.3 versus 0.2 (P < 0.0001; Analysis 12.1); at 4 years, 3.5 versus 1.9 (P < 0.0001; Analysis 12.2); at 10 years, 6.6 versus 4.9 (P < 0.0350; Analysis 12.3). By four years, the DMPA and LNG-IUC group lost more than 20%; by 10 years, overall loss was 84%. The groups had differential losses at all three time points.

Prospective studies:

Vickery 2013 was a substudy within CHOICE, a prospective study of 9256 women who received contraceptives at no cost. The investigators of the substudy examined weight change among women who had been continuous users of DMPA or the copper IUC for 11 months or longer. With a linear regression model, the investigators adjusted for the potential confounders of age and race. Weight change was not significantly different for use of DMPA compared with the Cu IUC (Analysis 12.4).

Study designs were similar in Dal'Ava 2014 and Dos Santos 2014. While Dal'Ava 2014 paired participants in the DMPA and Cu IUC groups by age (± 2 years) and weight (± 2 kg), Dos Santos 2014 matched by age (± 1 year) and BMI (± 1 kg/m²). The regression model in Dal'Ava 2014 included the potential confounders of physical activity, consumption of coffee and alcohol, and smoking in regression. At 12 months, the study arms did not differ significantly for changes in total body mass (weight), fat mass, or lean mass in either study (Analysis 11.6). A secondary report from Dos Santos 2014 (Modesto 2014) included 29 women using DMPA and 25 using the Cu IUC. Multiple linear regression adjusted for potential confounders such as age, schooling, and pregnancies. DMPA use was significantly associated with change in total fat mass compared with Cu IUC use by 12 months (reported beta 2.09 \pm SE 0.58; P < 0.002) but was not associated with change in percent body fat.

DMPA versus no hormonal method—Two studies compared women using DMPA to another group using no hormonal method.

Bonny 2009, mentioned above, also compared adolescents using DMPA 150 versus those using no hormonal method. By six months, the DMPA group had a greater increase in percent body fat (MD 11.00, 95% CI 2.64 to 19.36) (Analysis 13.1) and a greater decrease in percent lean body mass (MD -4.00, 95% CI -6.93 to -1.07) (Analysis 13.2).

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Nyirati 2013 compared DMPA at six weeks versus surgical sterilization among postpartum women, age 18 or older. By one year postpartum, the study arms did not differ significantly for change in weight, BMI, or percent body fat (Analysis 13.3 to Analysis 13.5). The sample size for the sterilization group was much smaller than that of the DMPA group.

Implants

Five studies examined implants, one RCT and four NRS. A 2013 report compared the single-rod etonogestrel (ENG) implant versus the copper IUC. Four older studies compared Norplant (six capsules) versus a non-hormonal IUC or another progestin-only contraceptive.

Vickery 2013, mentioned above with DMPA, also examined weight change over 12 months for the single-rod etonogestrel (ENG) implant versus the copper IUC. In the regression model adjusted for age and race, the ENG implant was not significantly associated with weight change compared with the Cu IUC (Analysis 12.4).

- Two studies utilized a non-hormonal IUC as the comparison.
 - In a study with lactating women, Salem 1984 showed a greater weight gain (kg) at six months for the Norplant group versus the Cu IUC group (MD 0.47, 95% CI 0.29 to 0.65) (Analysis 14.1). The Norplant group also had a greater weight gain (kg) than the group that used barrier, 'local,' or no contraceptive method (MD 0.74, 95% CI 0.52 to 0.96) (Analysis 15.1).
 - For the retrospective study of Sule 2005, the Norplant group had a significantly greater weight increase (kg) than the group with a non-hormonal IUC at one year (MD 1.10, 95% CI 0.36 to 1.84) (Analysis 14.2) but not at three years (Analysis 14.3). The same study compared the Norplant group versus a group using COCs. Weight change did not differ significantly between the groups at one year (Analysis 18.1). At three years, the COC group had only two participants.
- The retrospective study of Moore 1995 targeted adolescents and young women, ages 15 to 30 years. The Norplant and DMPA groups were not significantly different in mean weight change at one year (Analysis 16.1).
- In the RCT of Sivin 1998, mean weight change was not significantly different for the Norplant group versus the two-rod implant group at one, three, or five years (Analysis 17.1; Analysis 17.2; Analysis 17.3).

Levonorgestrel-releasing intrauterine contraception (LNG-IUC)

Four NRS examined the LNG-IUC versus the non-hormonal Cu IUC or no treatment.

Dal'Ava 2012 compared body composition changes with LNG-IUC versus with the Cu IUC. The two groups were paired by age (± 2 years) and BMI (± 2 kg/m²). At 12 months, the LNG-IUC group differed in percent change

in body fat mass compared with Cu IUC group (reported means 2.5% versus -1.3%; P = 0.029) (Analysis 19.1). The LNG-IUC group also differed from the Cu IUC users in percent change in lean body mass (reported means 1.4% versus 1.0%; P = 0.027).

In addition to DMPA and the ENG implant, Vickery 2013 also examined LNG-IUC versus the Cu IUC. The linear regression model indicated weight change with the LNG-IUC was not significantly different at 12 months from that with the non-hormonal IUC (Analysis 12.4).

In addition to examining weight change with DMPA use (above), Modesto 2015 compared weight change with LNG-IUC versus that with the Cu IUC. The study groups did not differ significantly for mean weight gain (kg) at 1 and 10 years after adjusting for years of school and number of children (Analysis 12.1; Analysis 12.3). As noted above for DMPA, losses were high and differential across groups.

Besides an OC containing desogestrel (above), Napolitano 2015 also compared body composition changes at 12 months for perimenopausal women using LNG-IUC versus a control group receiving no hormonal treatment. In unadjusted analysis, change in mean weight or BMI did not differ significantly between the LNG-IUC group and the control group at 12 months (Analysis 19.2; Analysis 19.3). However, compared with the control group, the LNG-IUC group had a greater mean increase in percent fat mass (MD 1.60, 95% CI 0.45 to 2.75) (Analysis 19.4) and therefore a greater mean decrease in fat free mass (MD -1.60, 95% CI -2.75 to -0.45) (Analysis 19.5).

DISCUSSION

Summary of main results

Actual mean weight gain at 6 or 12 months was limited, i.e. less than 2 kg for most studies up to one year (Table 2). The six studies with multiyear data showed that mean weight change was approximately twice as much at two to four years compared with one year, but generally the study groups did not differ significantly. These studies and their years of data included two RCTs (WHO 1983 [2 years]; Sivin 1998 [5 years]) and four retrospective studies (Espey 2000 [2 years]; Sule 2005 [3 years]; Pantoja 2010 [3 years]; Modesto 2015 [10 years]). Another RCT (Westhoff 2007) and a retrospective study (Taneepanichskul 1998) had data from 3 and 10 years of use, respectively, but not multiyear data.

In Table 3, we synthesized the results for each contraceptive category. Overall, 7 of the 22 included studies indicated a significant difference between groups for change in weight, body fat, or fat free mass (Salem 1984; Sule 2005; Bonny 2009; Pantoja 2010; Dal'Ava 2012; Modesto 2015; Napolitano 2015). Three examined DMPA, two studied the LNG-IUC, two focused on Norplant (six capsules), and one examined a desogestrel OC. The comparisons were groups using no hormonal method.

Summary of findings tables

This section focuses on the five studies with evidence of moderate or low quality that showed a significant difference between study arms (Table 3). Of 16 studies that examined DMPA (aside from one that also studied Norplant), three indicated an association between DMPA use and greater change in weight or other body composition measure (Summary of findings for the main comparison). Bonny 2009 was a small study of adolescents from a larger trial. Compared with a group using no hormonal contraceptive, the DMPA group had a greater increase in body fat percentage and a greater decrease in lean body mass. In Pantoja 2010, a retrospective study, mean weight gain was greater for the DMPA group versus the copper IUC group at one, two, and three years. The differences were notable within the normal to lower weight group and the overweight group but not within the obese group. For Modesto 2015, also retrospective, mean weight change was greater for the DMPA group compared with the copper IUC group at 1, 4, and 10 years, the three time points we examined. Four studies compared the LNG-IUC with a group not using any hormonal contraceptives. Two showed the study arms differed in body composition change by one year (Summary of findings 2), though they did not differ significantly for weight change. Within Dal'Ava 2012, participants using the LNG-IUC reportedly had a greater increase in fat mass (%) and a decrease in lean mass (%) compared with the non-hormonal IUC users. Similarly, in Napolitano 2015, the LNG-IUC group had a greater mean increase in fat mass (%) and a decrease in fat free mass (%) compared with the no-hormonal group.

One of two studies that examined P-O oral contraceptives showed an association between the OC and body composition change (Summary of findings 3). Napolitano 2015 compared use of an OC containing desogestrel 75 µg versus no hormonal contraceptive. The OC group showed the same pattern as the LNG-IUC group noted above, i.e. a greater mean increase in fat mass (%) and a decrease in fat free mass (%).

Overall completeness and applicability of evidence

Of the 22 included studies, 18 had data from a year or more of contraceptive use and 8 of those had data from two or more years. Weight gain (or the perception of weight gain) is frequently cited as a reason for discontinuing a contraceptive method. If a contraceptive method is associated with weight gain, a year is long enough to detect some change, though the amount may not be clinically significant. Of the eight studies with data from two or more years of contraceptive use, most showed the study groups did not differ significantly for weight gain, regardless of whether the comparison group used a progestin-only contraceptive or no hormonal method.

Within contraceptive method group, the studies varied in their comparison groups over time. Earlier DMPA studies generally compared DMPA with a hormonal contraceptive. Of the six studies added in this update, five compared DMPA with a non-hormonal IUC. For levonorgestrel-releasing intrauterine contraception, three of the four included studies were new. All four compared LNG-IUC with no hormonal method; in addition, two studied DMPA and one examined a P-O oral contraceptive. Overall, two studies of P-O pills met our inclusion criteria. We did not find any eligible studies of the progesterone-releasing vaginal ring. We do not have much evidence regarding weight change with currently marketed implants. Many studies of such implants did not meet our inclusion criteria, mainly due to the lack of comparative data on weight change. Two exceptions were a recent study of the etonogestrelreleasing implant and one that compared Norplant (six capsules) with a two-rod implant. Most of the weight change data for implants in this review came from studies of Norplant, which is no longer marketed.

Quality of the evidence

We assessed the quality of evidence as noted earlier (Assessment of risk of bias in included studies). Table 1 has a summary based on the factors used in our assessment. We considered the overall quality of evidence to be low, given that evidence from12 of the 22 studies was low. Three studies provided moderate quality evidence and seven had very low quality evidence. We downgraded the 17 NRS for lack of randomization and three RCTs for insufficient information on randomization and allocation concealment. Most studies had high loss to follow-up or were retrospective studies that may not have accounted for losses to follow-up or early discontinuation.

Potential biases in the review process

We selected studies that had data on mean change in weight or other body composition measure. Several excluded studies did not report the data we needed. For the initial review, many studies were older, which limited our ability to obtain additional information from the investigators.

Agreements and disagreements with other studies or reviews

As noted earlier, many concerns about weight gain with POC use are based on perceptions and discontinuation reasons rather than measures of actual weight change. We found limited evidence of significant change for POC users versus those who did not use hormonal contraceptives. Actual weight gain was less than 2 kg (4.4 lb) up to one year. Another review showed no clear evidence of weight gain with the use of combined hormonal contraceptives (Gallo 2014). People may gain weight over time regardless of contraceptive use.

Two studies compared perceived weight gain with actual weight gain among POC users. In a substudy of the CHOICE project, women who perceived weight gain of 5 lb or more had a mean weight change of 10 lb by 12 months, which is about 8 lb more than those who did not perceive a gain (Nault 2013). Risk of perceived gain was greater for the POC group versus the Cu IUC group. However, Vickery 2013, another substudy of CHOICE, did not show a difference in weight change by 12months between the POC groups and the Cu IUC group. A secondary analysis of data from an RCT compared users of a two-rod levonorgestrel implant versus women who did not yet receive the implant (Gallo 2016). Perceived weight gain was more common in the implant arm compared with the control group. Women with perceived weight gain in the implant group were more likely to have a gain of 2 kg by three months. The proportions of women who gained 2 kg did not differ significantly between the two groups nor did median weight gain.

Reviews have suggested that DMPA and weight gain may be a concern for women who are already obese, whether they are adolescents (Curtis 2009) or adults (Merki-Feld 2015). The review with three studies on adolescents considered the quality of evidence to be fair for two studies. They lacked numbers for those discontinued due to weight gain and did not have a non-hormonal comparison. The third study lost 37% of DMPA users by 18 months, thus producing high risk of bias. A retrospective study of adult women in this review showed that mean weight gain did not differ significantly between the DMPA and Cu IUC groups within the obese subgroup (Pantoja 2010). Within the normal and overweight subgroups though, weight gain was greater for the DMPA group. We included three studies of young women, but none with adolescents who were obese. One showed an increase in total body fat and a decrease in lean body mass for adolescents in the DMPA group compared with those in a non-hormonal group (Bonny 2009).

A review of adverse events examined whether early weight among DMPA users was associated with later weight gain (Steenland 2013). The researchers concluded weight change greater than 5% of baseline weight was associated with greater mean change in weight or BMI at follow-up. Two studies grouped DMPA users by early weight gain in secondary analysis from studies of bone mineral density. One study explored whether a 5% weight change in 6 months predicted weight change by 36 months (Le 2009). The original study lost 60% of DMPA users by 12 months and 76% by 36 months, leading to high risk of bias for the results. Predictors of early weight gain, among the 60% participating at six months, were a BMI < 30 and a reported increase in appetite. The second study was apparently based on a study that lost 24% by 12 months and 43% by 24 months; the report did not include the sample sizes at 6, 12, or 18 months (Bonny 2011). Of participants in the study at 12 or 18 months, baseline characteristics may have differed between those with 5% weight gain at six months and those with less weight gain.

AUTHORS' CONCLUSIONS

Implications for practice

We found limited evidence of weight gain when using progestin-only contraceptives. We identified some significant differences when a P-O method was compared with no hormonal contraceptive. This includes studies of a P-O oral contraceptive, DMPA, levonorgestrel IUC, and an older implant.

Overall, actual mean weight gain was low for 6 to 12 months, i.e. less than 2 kg (4.4 lb) in most studies. More weight gain was noted at two and three years, but the comparison groups did not differ much for weight change. People may gain weight over time regardless of contraceptive use. Appropriate and accurate counseling about typical weight gain may help reduce discontinuation of contraceptives due to perceptions of weight gain.

Implications for research

Five studies with moderate or low quality evidence showed a significant difference between study arms. They examined DMPA, LNG-IUC, and a P-O pill versus no hormonal method use. All included a group not using a hormonal method. The outcomes were change in

weight, percent body fat, or percent lean body mass. Three were prospective nonrandomized studies and two were retrospective chart reviews. The overall quality of evidence was low, largely due to the lack of randomization and high losses.

Weight change is rarely the focus of prospective contraceptive studies. Some of the newer prospective studies did focus on change in weight or body composition. Well-designed RCTs assessing weight change over time would better address this issue. Comparisons could be between a P-O method and a non-hormonal group. However, careful counseling and follow-up are needed to avoid the high losses to follow-up and discontinuation found in many contraceptive studies.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

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Internal sources

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No sources of support supplied

External sources

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APPENDICES

Appendix 1. Search 2016

MEDLINE via PubMed (1 January 2013 to 4 August 2016)

(contraceptive agents, female[Mesh] OR contraceptive devices, female[Mesh] OR contracept*) AND (progest* OR "progestin only" OR "progestin only" contracept* OR "progestin only pill" OR progestin* OR progesteron* OR progestational, hormones, synthetic OR progestogen* OR medroxyprogesterone OR DMPA OR (levonorgestrelreleasing AND intrauterine) OR "intrauterine device" OR "intrauterine contraception" OR IUD OR IUC OR (etonogestrel-releasing AND implant) OR (ETG AND implant) OR progesterone OR gestagen OR "progestogen only") AND (body mass index OR BMI OR weight) NOT (cancer[ti] OR polycystic [ti] OR exercise [ti] OR physical activity[ti] OR postmenopaus*[ti])

CENTRAL (Cochrane Central Register of Controlled Trials 2016, issue 1 (23 February 2016))

Abstract: weight OR body mass index OR BMI

AND Title, Abstract, Keywords: contraception OR contraceptive

NOT Record Title: premenstrual OR dysmenor* OR endometr* OR *androgen* OR HIV OR polycystic OR PCOS OR cancer OR exercise OR anorexi* OR bulimi*

NOT Record Title: postmenopausal OR post-menopausal OR hormone therapy OR male hormonal OR male contracept* OR testosterone

Publication Year from 2013 to 2015 in Trials

POPLINE (2013 to 2016 (23 February 2016))

Keyword: Body Weight

AND Keyword: Contraceptive Agents Progestin

Web of Science (24 March 2016)

TOPIC:(contracept*) AND TOPIC:(overweight OR obese OR obesity OR body mass index OR BMI)

NOT TITLE: (cancer OR polycystic OR PCOS OR diabetes OR exercise OR physical activity OR postmenopaus* OR hormone therapy OR replacement)

Refined by: RESEARCHAREAS: (OBSTETRICS GYNECOLOGY OR PHARMACOLOGY PHARMACY OR ENDOCRINOLOGY METABOLISM) AND DOCUMENT TYPES: (ARTICLE)

Timespan: 2013–2016.

Search language=Auto

LILACS via VHL Regional Portal (29 March 2016)

tw:((tw:(('contraceptive agents female")) OR (tw:(contracept*)))) AND (tw:(weight OR overweight OR obes* OR "body mass index"))) AND (instance:"regional") AND (db: ("LILACS")

AND limit:("female")

AND year cluster: ("2012" OR "2013" OR "2014" OR "2015"))

[Note: no listing for 2016]

ClinicalTrials.gov (7 March 2016)

Search terms: overweight OR obese OR obesity OR weight OR body mass index OR BMI

Condition: NOT (HIV OR polycystic OR PCOS OR cancer OR anorexia OR pulmonary OR metabolic OR amenorrhea)

Intervention: contraceptive OR contraception OR contraceptives

Gender: studies with female participants

First received: From 1 December 2012 to 16 October 2015

ICTRP (29 March 2016)

Condition: contraceptive OR contraception

AND Intervention: progestin OR progestin-only OR IUD OR implant OR medroxyprogesterone OR norethisterone

Recruitment status: All

Date of registration is between 1 December 2012 and 29 March 2016

Appendix 2. Previous searches

2013

MEDLINE via PubMed (1 January 2010 to 5 June 2013)

(contraceptive agents, female[Mesh] OR contraceptive devices, female[Mesh] OR contracept*) AND (progest* OR "progestin only" OR "progestin only" contracept* OR "progestin only pill" OR progestin* OR progesteron* OR progestational, hormones, synthetic OR progestogen* OR progesterone OR gestagen OR "progestogen only") AND (body weight changes OR weight gain OR weight loss OR body mass index OR BMI OR weight) NOT (cancer[ti] OR polycystic [ti] OR exercise [ti] OR physical activity[ti] OR postmenopaus*[ti])

limited to human, female

CENTRAL (2010 to 1 June 2013)

weight OR body mass index OR BMI in Abstract

AND contraception OR contraceptive in Title, Abstract or Keywords

NOT premenstrual OR dysmenor* OR endometr* OR *androgen* OR HIV OR polycystic OR PCOS OR cancer OR exercise OR anorexia OR bulimic in Record Title

NOT postmenopausal OR post-menopausal OR hormone therapy OR male hormonal OR male contracept* OR testosterone in Record Title

POPLINE (2010 to 26 December 2012)

Global: weight OR BMI OR body mass index

Keyword: contraceptive agents, progestin OR Low-Dose Progestins

Filter: research report

LILACS (2010 to 26 December 2012)

contraceptive agents or Agentes Anticonceptivos Femeninos or Anticoncepcionais Femininos or contraceptive devices, female or Dispositivos Anticonceptivos Femeninos or Dispositivos Anticoncepcionais Femininos or contraceptives or Anticonceptivos or Anticoncepcionais [Words]

AND weight or body weight or Peso Corporal or weight gain or Aumento de Peso or Ganho de Peso or weight reduction or weight loss or Pérdida de Peso or Perda de Peso or body weight changes or Cambios en el Peso Corporal or Alterações do Peso Corporal or body mass index or BMI [Words]

ClinicalTrials.gov (1 January 2010 to 26 December 2012)

Search terms: overweight OR obese OR obesity OR weight OR body mass index OR BMI

Condition: NOT (HIV OR polycystic OR PCOS OR cancer OR anorexia OR pulmonary OR metabolic OR amenorrhea)

Intervention: contraceptive OR contraception

Study type: interventional studies

Gender: studies with female participants

ICTRP (2010 to 26 December 2012)

- 1. Intervention: contraceptive OR contraception
- 2. Condition: contraceptive OR contraception

Intervention: progestin OR progestin-only OR IUD OR implant OR medroxyprogesterone OR norethisterone

2010

MEDLINE via PubMed (through 11 June 2010)

(contraceptive agents, female[Mesh] OR contraceptive devices, female[Mesh] OR contracept*) AND (progest* OR "progestin only" OR "progestin only" contracept* OR "progestin only pill" OR progestin* OR progesteron* OR progestational, hormones, synthetic OR progestogen* OR progesterone OR gestagen OR "progestogen only") AND

(body weight changes OR weight gain OR weight loss OR body mass index OR BMI OR weight) NOT (cancer[ti] OR polycystic [ti] OR exercise [ti] OR physical activity[ti] OR postmenopaus*[ti])

limited to human, female

POPLINE (through 19 May 2010)

(progestin only contracept*/ contraceptive agents, progestin/low-dose progestins) & (weight/ weight gain/weight loss/body weight/BMI/body mass index/ weight change)

CENTRAL (through 19 May 2010)

weight OR body mass index OR BMI in Abstract and contraception OR contraceptive in Title, Abstract or Keywords

NOT premenstrual OR dysmenor* OR endometr* OR *androgen* OR HIV OR polycystic OR PCOS OR cancer OR exercise OR anorexia OR bulimic in Record Title

NOT postmenopausal OR post-menopausal OR hormone therapy OR male hormonal OR male contracept* OR testosterone in Record Title

EMBASE (through 15 June 2010)

(contraceptive agent, progestin --side effects

or

((contraceptive device or contraceptives or contracept*) and (gestagen! or progest? or progestin? or progesterone? or progestational, hormones, synthetic or progestogen?or progestin()only or progestin()only()contracept or progestin()only()pill or progestogen()only)))

and

(body weight! or weight gain or weight reduction or weight()loss or body()mass()index)

not (cancer or polycystic or exercise or physical() activity or postmenopaus? or oral contraceptives, combined)

and body weight/de

limited to human

LILACS (through 1 July 2010)

contraceptive agents or Agentes Anticonceptivos Femeninos or Anticoncepcionais Femininos or contraceptive devices, female or Dispositivos Anticonceptivos Femeninos or Dispositivos Anticoncepcionais Femininos or contraceptives or Anticonceptivos or Anticoncepcionais [Words] and weight or body weight or Peso Corporal or weight gain or Aumento de Peso or Ganho de Peso or weight reduction or weight loss or Pérdida de Peso or Perda de Peso or body weight changes or Cambios en el Peso Corporal or Alterações do Peso Corporal or body mass index or BMI [Words]

ClinicalTrials.gov (through 27 April 2010)

Search terms: overweight OR obese OR obesity OR weight OR body mass index OR BMI

Condition: NOT (HIV OR polycystic OR PCOS OR cancer OR anorexia OR pulmonary OR metabolic OR amenorrhea)

Intervention: contraceptive OR contraception

Study type: interventional studies

Gender: studies with female participants

ICTRP (through 07 September 2010)

- **1.** Intervention: contraceptive OR contraception
- 2. Condition: contraceptive OR contraception

Intervention: progestin OR progestin-only OR IUD OR implant OR medroxyprogesterone OR norethisterone

Appendix 3. Newcastle-Ottawa Quality Assessment Scale for cohort studies

<u>Note:</u> A study can be awarded a maximum of one star (*) for each numbered item within the Selection and Outcome categories. A maximum of two stars can be given for Comparability.

Selection

1.	<u>Representativeness of the exposed cohort</u>		
	a.	truly representative of the average (describe) in the community *	
	b.	somewhat representative of the average \cdots in the community $\boldsymbol{*}$	
	с.	selected group of users eg nurses, volunteers	
	d.	no description of the derivation of the cohort	
2.	Selection of the non exposed cohort		
	a.	drawn from the same community as the exposed cohort $*$	
	b.	drawn from a different source	
	с.	no description of the derivation of the non exposed cohort	

3.	Ascertainment of exposure		
	a.	secure record (eg surgical records) *	
	b.	structured interview *	
	с.	written self report	
	d.	no description	
4.	Demonstration that outcome of interest was not present at start of study		
	a.	yes *	
	b.	no	

Comparability

1.

<u>Comparab</u>	ility of cohorts on the basis of the design or analysis
a.	study controls for \cdots (select the most important factor) *
b.	study controls for any additional factor* (This criteria could be modified to indicate specific control for a second important factor.)

Outcome

1.	Assessment of outcome		
	a.	independent blind assessment *	
	b.	record linkage *	
	c.	self report	
	d.	no description	
2.	Was follow-up long enough for outcomes to occur		
	a.	yes (select an adequate follow up period for outcome of interest) *	
	b.	no	
3.	Adequacy of follow up of cohorts		
	a.	complete follow up - all subjects accounted for $*$	
	b.	subjects lost to follow up unlikely to introduce bias - small number lost - > \cdots % (select an adequate %) follow up, or description provided of those lost) *	
	c.	follow up rate $< \cdots $ % (select an adequate %) and no description of those lost	
	d.	no statement	

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* Indicates the major publication for the study

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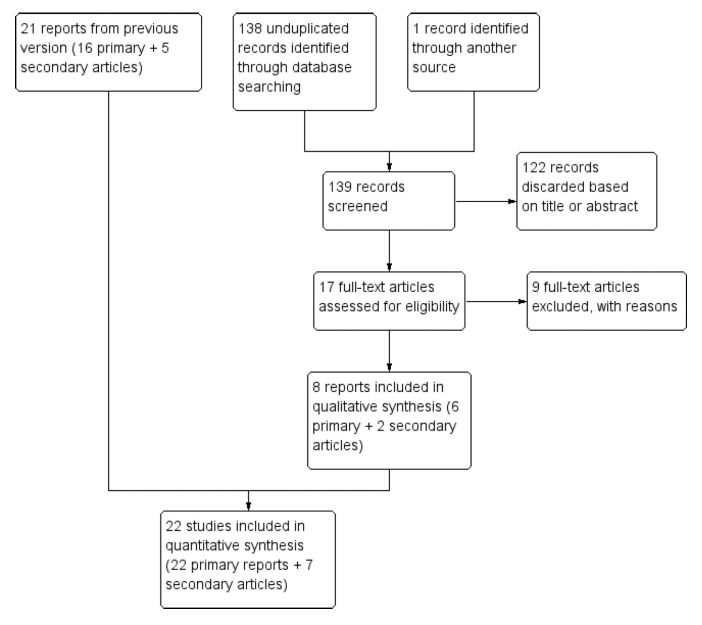
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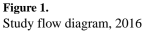
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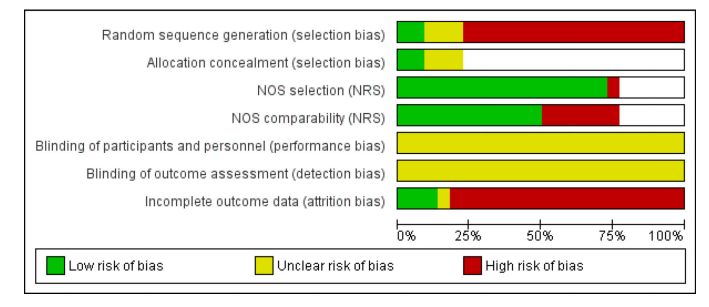


Figure 2.

Risk of bias graph: review authors' judgements about each risk of bias item presented as percentages across all included studies.



Figure 3.

Risk of bias summary: review authors' judgements about each risk of bias item for each included study.

Cochrane Database Syst Rev. Author manuscript; available in PMC 2017 August 28.

Review: Progestin-only contraceptives: effects on weight

Comparison: I Norethisterone 350 g OC versus levonorgestrel 30 g OC

Outcome: I Mean weight change (kg) at 6 months

Study or subgroup	Treatment		Control			Mean Difference		Weight	Mean Difference		
	N	Mean(SD)	Ν	Mean(SD)		IV,Fixed,95% CI					IV,Fixed,95% CI
Ball 1991	23	0 (1.7)	16	0.6 (1.9)			-			100.0 %	-0.60 [-1.76, 0.56]
Total (95% CI)	23		16							100.0 %	-0.60 [-1.76, 0.56]
Heterogeneity: not ap	plicable										
Test for overall effect: $Z = 1.01$ (P = 0.31)											
Test for subgroup diffe	erences: Not appl	icable									
					-4	-2	0	2	4		
				F	avors tr	eatment		Favors o	ontrol		

Analysis 1.1.

Comparison 1 Norethisterone 350 μ g OC versus levonorgestrel 30 μ g OC, Outcome 1 Mean weight change (kg) at 6 months.

Review: Progestin-only contraceptives: effects on weight

Comparison: 2 Desogestrel (DSG) OC versus control (no hormonal method)

Outcome: I Mean change in weight (kg) at I year

Study or subgroup	Treatment		Control			D	M oiffere	ean nce		Weight	Mean Difference
	Ν	Mean(SD)	Ν	Mean(SD)		IV,Fixed,95% CI				IV,Fixed,95% CI	
Napolitano 2015	42	0.3 (3.7)	26	-0.2 (3.6)			+	-		100.0 %	0.50 [-1.28, 2.28]
Total (95% CI)	42		26							100.0 %	0.50 [-1.28, 2.28]
Heterogeneity: not ap	plicable										
Test for overall effect:	Z = 0.55 (P = 0.	58)									
Test for subgroup diffe	erences: Not appl	icable									
					1						
					-2	-	0	1	2		
				Fa	ivors ti	reatment		Favors c	ontrol		

Analysis 2.1.

Comparison 2 Desogestrel (DSG) OC versus control (no hormonal method), Outcome 1 Mean change in weight (kg) at 1 year.

Review: Progestin-only contraceptives: effects on weight

Comparison: 2 Desogestrel (DSG) OC versus control (no hormonal method)

Outcome: 2 Mean change in BMI (kg/m²) at I year

Study or subgroup	Treatment		Control		Mean Difference	Weight	Mean Difference
	Ν	Mean(SD) N Me		Mean(SD)	IV,Fixed,95% CI		IV,Fixed,95% CI
Napolitano 2015	42	0.5 (1.6)	26	0.5 (1)		100.0 %	0.0 [-0.62, 0.62]
Total (95% CI)	42		26			100.0 %	0.0 [-0.62, 0.62]
Heterogeneity: not ap	plicable						
Test for overall effect:	Z = 0.0 (P = 1.0)						
Test for subgroup diffe	rences: Not appli	cable					
				-	-1 -0.5 0 0.5	I.	
				Favo	ors treatment Favors contr	ol	

Analysis 2.2.

Comparison 2 Desogestrel (DSG) OC versus control (no hormonal method), Outcome 2 Mean change in BMI (kg/m2) at 1 year.

Review: Progestin-only contraceptives: effects on weight

Comparison: 2 Desogestrel (DSG) OC versus control (no hormonal method)

Outcome: 3 Mean change in fat mass (%) at I year

Study or subgroup	Treatment	Control			Diff	Mean ference	Weight	Mean Difference	
	Ν	Mean(SD)	Ν	Mean(SD)	IV,Fixe	ed,95% Cl		IV,Fixed,95% CI	
Napolitano 2015	42	2.8 (3.5)	26	-0.5 (1.6)			100.0 %	3.30 [2.08, 4.52]	
Total (95% CI)	42		26			-	100.0 %	3.30 [2.08, 4.52]	
Heterogeneity: not ap	plicable								
Test for overall effect:	Z = 5.28 (P < 0.0)	(10000)							
Test for subgroup diffe	rences: Not appl	icable							
						<u> </u>			
				-	4 -2	0 2 4			
				Favo	ors treatment	Favors control			

Analysis 2.3.

Comparison 2 Desogestrel (DSG) OC versus control (no hormonal method), Outcome 3 Mean change in fat mass (%) at 1 year.

Review: Progestin-only contraceptives: effects on weight

Comparison: 2 Desogestrel (DSG) OC versus control (no hormonal method)

Outcome: 4 Mean change in fat free mass (%) at I year

Study or subgroup	Treatment N	Mean(SD)	Control N	Mean(SD)		Mean Difference IV,Fixed,95% CI		erence Weight		Weight	Mean Difference IV,Fixed,95% Cl
Napolitano 2015	42	-2.8 (3.5)	26	0.5 (1.6)	•	<u> </u>				100.0 %	-3.30 [-4.52, -2.08]
Total (95% CI)	42		26		-					100.0 %	-3.30 [-4.52, -2.08]
Heterogeneity: not applicable											
Test for overall effect: $Z = 5.28 (P < 0.00001)$											
Test for subgroup diffe	erences: Not app	licable									
					-4	-2	0	2	4		
					Favors	control		Favors t	treatmer	nt	

Analysis 2.4.

Comparison 2 Desogestrel (DSG) OC versus control (no hormonal method), Outcome 4 Mean change in fat free mass (%) at 1 year.

Review: Progestin-only contraceptives: effects on weight

Comparison: 3 DMPA 150 mg/mL versus COC

Outcome: I Weight gain \geq I kg at month 6

Study or subgroup	Treatment	Control	Odds Ratio	Weight	Odds Ratio	
	n/N	n/N	M-H,Fixed,95% CI		M-H,Fixed,95% CI	
Tankeyoon 1976	5/15	9/16		100.0 %	0.39 [0.09, 1.67]	
Total (95% CI)	15	16	-	100.0 %	0.39 [0.09, 1.67]	
Total events: 5 (Treatmen	t), 9 (Control)					
Heterogeneity: not applica	able					
Test for overall effect: Z =	= 1.27 (P = 0.20)					
Test for subgroup differen	ces: Not applicable					
600 x						
			0.01 0.1 1 10 100)		
		Fav	vors experimental Favors control			

Analysis 3.1. Comparison 3 DMPA 150 mg/mL versus COC, Outcome 1Weight gain 1 kg at month 6.

Review: Progestin-only contraceptives: effects on weight

Comparison: 3 DMPA 150 mg/mL versus COC

Outcome: 2 Weight gain \geq 1 kg at month 12

Study or subgroup	Treatment	Control	Odds Ratio	Weight	Odds Ratio
	n/N	n/N	M-H,Fixed,95% CI		M-H,Fixed,95% CI
Tankeyoon 1976	8/14	9/12		100.0 %	0.44 [0.08, 2.39]
Total (95% CI)	14	12		100.0 %	0.44 [0.08, 2.39]
Total events: 8 (Treatmen	t), 9 (Control)				
Heterogeneity: not applic	able				
Test for overall effect: Z =	= 0.95 (P = 0.34)				
Test for subgroup differen	ces: Not applicable				
			0.01 0.1 1 10 10	00	
		Fave	ors experimental Favors contro	ol	

Analysis 3.2.

Comparison 3 DMPA 150 mg/mL versus COC, Outcome 2 Weight gain 1 kg at month 12.

Review: Progestin-only contraceptives: effects on weight

Comparison: 3 DMPA 150 mg/mL versus COC

Outcome: 3 Weight loss \geq 1 kg at month 6

Study or subgroup	Treatment	Control	Odds Ratio	Weight	Odds Ratio
	n/N	n/N	M-H,Fixed,95% Cl		M-H,Fixed,95% Cl
Tankeyoon 1976	3/15	0/16		- 100.0 %	9.24 [0.44, 195.69]
Total (95% CI)	15	16		100.0 %	9.24 [0.44, 195.69]
Total events: 3 (Treatmen	it), 0 (Control)				
Heterogeneity: not applic	able				
Test for overall effect: Z =	= 1.43 (P = 0.15)				
Test for subgroup differer	nces: Not applicable				
				1	
			0.005 0.1 1 10 2	200	
			Favors experimental Favors contr	rol	

Analysis 3.3. Comparison 3 DMPA 150 mg/mL versus COC, Outcome 3 Weight loss 1 kg at month 6.

Review: Progestin-only contraceptives: effects on weight

Comparison: 3 DMPA 150 mg/mL versus COC

Outcome: 4 Weight loss \geq 1 kg at month 12

Study or subgroup	Treatment	Control	Odds Ratio	Weight	Odds Ratio
	n/N	n/N	M-H,Fixed,95% CI		M-H,Fixed,95% Cl
Tankeyoon 1976	2/14	1/12		100.0 %	1.83 [0.15, 23.15]
Total (95% CI)	14	12		100.0 %	1.83 [0.15, 23.15]
Total events: 2 (Treatmen	t), I (Control)				
Heterogeneity: not applic	able				
Test for overall effect: Z =	= 0.47 (P = 0.64)				
Test for subgroup differen	ices: Not applicable				
				1	
			0.01 0.1 1 10 10	00	
		Fa	vors experimental Favors contr	ol	

Analysis 3.4. Comparison 3 DMPA 150 mg/mL versus COC, Outcome 4 Weight loss 1 kg at month 12.

Review: Progestin-only contraceptives: effects on weight

Comparison: 3 DMPA 150 mg/mL versus COC

Outcome: 5 Mean weight change (kg) at 6 months

Study or subgroup	Treatment		Control			Mean Difference		Weight	Mean Difference		
	Ν	Mean(SD)	Ν	Mean(SD)		IV,Fi>	ked,9	5% CI			IV,Fixed,95% CI
Tuchman 2005	49	0.6 (5.8)	93	1.1 (3.4)				_		100.0 %	-0.50 [-2.26, 1.26]
Total (95% CI)	49		93					-		100.0 %	-0.50 [-2.26, 1.26]
Heterogeneity: not ap	plicable										
Test for overall effect:	Z = 0.56 (P = 0.	58)									
Test for subgroup diffe	rences: Not app	licable									
								- i			
					-4	-2	0	2	4		
				F	avors tr	eatment		Favors c	ontrol		

Analysis 3.5.

Comparison 3 DMPA 150 mg/mL versus COC, Outcome 5 Mean weight change (kg) at 6 months.

Review: Progestin-only contraceptives: effects on weight

Comparison: 3 DMPA 150 mg/mL versus COC

Outcome: 6 Mean weight change (kg) at 12 months

Study or subgroup	Treatment		Control			D	M oiffere	ean nce		Weight	Mean Difference	
	N	Mean(SD)	N	Mean(SD)		IV,F	ixed,9	5% CI			IV,Fixed,95% CI	
Tuchman 2005	29	1.7 (6.1)	52	I (5.I)					_	100.0 %	0.70 [-1.92, 3.32]	
Total (95% CI)	29		52						-	100.0 %	0.70 [-1.92, 3.32]	
Heterogeneity: not applicable												
Test for overall effect:	Test for overall effect: $Z = 0.52$ (P = 0.60)											
Test for subgroup diffe	erences: Not appl	icable										
									1			
					-4	-2	0	2	4			
				Fav	ors tr	reatment		Favors o	ontrol			

Analysis 3.6.

Comparison 3 DMPA 150 mg/mL versus COC, Outcome 6 Mean weight change (kg) at 12 months.

Review: Progestin-only contraceptives: effects on weight

Comparison: 3 DMPA 150 mg/mL versus COC

Outcome: 7 Mean percentage weight change at 6 months

Study or subgroup	Treatment		Control			Mean Difference		Weight	Mean Difference		
	Ν	Mean(SD)	Ν	Mean(SD)		IV,Fi>	ked,9	5% CI			IV,Fixed,95% CI
Tuchman 2005	49	1.2 (7.8)	93	1.9 (4.9)	,					100.0 %	-0.70 [-3.10, 1.70]
Total (95% CI)	49		93							100.0 %	-0.70 [-3.10, 1.70]
Heterogeneity: not applicable											
Test for overall effect: $Z = 0.57$ (P = 0.57)											
Test for subgroup diffe	rences: Not app	licable									
					ĩ			ĩ	1		
					-4	-2	0	2	4		
				F	avors t	reatment		Favors co	ontrol		

Analysis 3.7.

Comparison 3 DMPA 150 mg/mL versus COC, Outcome 7 Mean percentage weight change at 6 months.

Review: Progestin-only contraceptives: effects on weight

Comparison: 3 DMPA 150 mg/mL versus COC

Outcome: 8 Mean percentage weight change at 12 months

Study or subgroup	Treatment N	Mean(SD)	Control N	Mean(SD)		Mean ference ed,95% Cl	Weight	Mean Difference IV,Fixed,95% Cl
Tuchman 2005	29	2.9 (8.3)	52	2.9 (8.4)			100.0 %	0.0 [-3.79, 3.79]
Total (95% CI)	29		52				100.0 %	0.0 [-3.79, 3.79]
Heterogeneity: not ap	plicable							
Test for overall effect:	Z = 0.0 (P = 1.0)							
Test for subgroup diffe	rences: Not appli	cable						
					-4 -2	0 2 4		
				Fa	avors treatment	Favors control		

Analysis 3.8.

Comparison 3 DMPA 150 mg/mL versus COC, Outcome 8 Mean percentage weight change at 12 months.

Review: Progestin-only contraceptives: effects on weight

Comparison: 4 DMPA 150 mg/mL versus MPA + E₂C

Outcome: I Mean weight change (kg) at 6 months

Study or subgroup	Treatment N	Mean(SD)	Control N	Mean(SD)			fferer	ean nce 5% Cl		Weight	Mean Difference IV,Fixed,95% Cl
Tuchman 2005	49	0.6 (5.8)	21	1.2 (4.3)		-	-			100.0 %	-0.60 [-3.05, 1.85]
Total (95% CI)	49		21				-			100.0 %	-0.60 [-3.05, 1.85]
Heterogeneity: not ap	plicable										
Test for overall effect:	Z = 0.48 (P = 0.4)	.63)									
Test for subgroup diffe	erences: Not app	licable									
					r.			, i			
					-10	-5	0	5	10		
				Fa	avors tre	eatment		Favors	control		

Analysis 4.1.

Comparison 4 DMPA 150 mg/mL versus MPA + E2C, Outcome 1 Mean weight change (kg) at 6 months.

Review: Progestin-only contraceptives: effects on weight

Comparison: 4 DMPA 150 mg/mL versus MPA + E₂C

Outcome: 2 Mean weight change (kg) at 12 months

Study or subgroup	Treatment N	Mean(SD)	Control N	Mean(SD)			Me fferer ked,9!			Weight	Mean Difference IV,Fixed,95% Cl
Tuchman 2005	29	1.7 (6.1)	17	3 (9.6)				_		100.0 %	-1.30 [-6.37, 3.77]
Total (95% CI)	29		17					-		100.0 %	-1.30 [-6.37, 3.77]
Heterogeneity: not ap	plicable										
Test for overall effect:	Z = 0.50 (P = 0.)	62)									
Test for subgroup diffe	rences: Not app	licable									
					i.						
					-10	-5	0	5	10		
				Fav	vors tre	eatment		Favors	control		

Analysis 4.2.

Comparison 4 DMPA 150 mg/mL versus MPA + E2C, Outcome 2 Mean weight change (kg) at 12 months.

Review: Progestin-only contraceptives: effects on weight

Comparison: 4 DMPA 150 mg/mL versus MPA + E₂C

Outcome: 3 Mean percentage weight change at 6 months

Study or subgroup	Treatment		Control			D	Me iffere	ean nce		Weight	Mean Difference
	Ν	Mean(SD)	Ν	Mean(SD)		IV,Fi:	xed,9	5% CI			IV,Fixed,95% CI
Tuchman 2005	49	1.2 (7.8)	21	1.8 (6.2)				-		100.0 %	-0.60 [-4.04, 2.84]
Total (95% CI)	49		21				-	-		100.0 %	-0.60 [-4.04, 2.84]
Heterogeneity: not ap	plicable										
Test for overall effect:	Z = 0.34 (P = 0.34)	73)									
Test for subgroup diffe	rences: Not appl	licable									
					ī						
					-10	-5	0	5	10		
				Fa	vors tr	eatment		Favors o	ontrol		

Analysis 4.3.

Comparison 4 DMPA 150 mg/mL versus MPA + E2C, Outcome 3 Mean percentage weight change at 6 months.

Review: Progestin-only contraceptives: effects on weight

Comparison: 4 DMPA 150 mg/mL versus MPA + E₂C

Outcome: 4 Mean percentage weight change at 12 months

Study or subgroup	Treatment		Control			Di		lean ence		Weight	Mean Difference
	Ν	Mean(SD)	Ν	Mean(SD)		IV,Fi>	ked,	95% CI			IV,Fixed,95% CI
Tuchman 2005	29	2.9 (8.3)	17	3.6 (13)	-		•			100.0 %	-0.70 [-7.58, 6.18]
Total (95% CI)	29		17		-					100.0 %	-0.70 [-7.58, 6.18]
Heterogeneity: not ap	plicable										
Test for overall effect:	Z = 0.20 (P = 0.)	84)									
Test for subgroup diffe	erences: Not app	licable									
								ī			
					-10	-5	0	5	10		
				Fa	avors tre	eatment		Favors o	ontrol		

Analysis 4.4.

Comparison 4 DMPA 150 mg/mL versus MPA + E2C, Outcome 4 Mean percentage weight change at 12 months.

Review: Progestin-only contraceptives: effects on weight

Comparison: 5 DMPA 150 mg/mL + placebo versus DMPA 150 mg/mL + E₂C

Outcome: I Mean change in total body fat (%) at 6 months

Study or subgroup	Treatment		Control			D	M iffere	lean ence		Weight	Mean Difference
	Ν	Mean(SD)	Ν	Mean(SD)		IV,Fi	xed,9	95% CI			IV,Fixed,95% CI
Bonny 2009	8	10.3 (10.49)	7	2.8 (4.4)				+	-	100.0 %	7.50 [-0.47, 15.47]
Total (95% CI)	8		7				-		-	100.0 %	7.50 [-0.47, 15.47]
Heterogeneity: not ap	plicable										
Test for overall effect:	Z = 1.85 (P = 0	0.065)									
Test for subgroup diffe	erences: Not app	olicable									
					1	- T			- T		
					-20	-10	0	10	20		
				Fa	avors tr	reatment		Favors o	ontrol		

Analysis 5.1.

Comparison 5 DMPA 150 mg/mL + placebo versus DMPA 150 mg/mL + E2C, Outcome 1 Mean change in total body fat (%) at 6 months.

Review: Progestin-only contraceptives: effects on weight

Comparison: 5 DMPA 150 mg/mL + placebo versus DMPA 150 mg/mL + E₂C

Outcome: 2 Mean change in lean body mass (%) at 6 months

Study or subgroup	Treatment		Control		D	Mean ifference	Weight	Mean Difference
	Ν	Mean(SD)	Ν	Mean(SD)	IV,Fi	xed,95% Cl		IV,Fixed,95% CI
Bonny 2009	8	-3.4 (3.57)	7	-1.2 (1.77)	←	-	100.0 %	-2.20 [-5.00, 0.60]
Total (95% CI)	8		7			-	100.0 %	-2.20 [-5.00, 0.60]
Heterogeneity: not ap	plicable							
Test for overall effect:	Z = 1.54 (P = 0.)	12)						
Test for subgroup diffe	erences: Not app	licable						
					-4 -2	0 2	4	
					Favors control	Favors tre	atment	

Analysis 5.2.

Comparison 5 DMPA 150 mg/mL + placebo versus DMPA 150 mg/mL + E2C, Outcome 2 Mean change in lean body mass (%) at 6 months.

Review: Progestin-only contraceptives: effects on weight

Comparison: 6 DMPA 150 mg/mL versus DMPA 450 mg/mL

Outcome: I Mean weight change (kg) at 6 months

Study or subgroup	Treatment		Control			D	M Differe	ean nce		Weight	Mean Difference
	Ν	Mean(SD)	Ν	Mean(SD)		IV,F	ixed,9	95% CI			IV,Fixed,95% CI
Castle 1978	285	0.33 (3.38)	366	0.32 (2.87)						100.0 %	0.01 [-0.48, 0.50]
Total (95% CI)	285		366				-			100.0 %	0.01 [-0.48, 0.50]
Heterogeneity: not ap	plicable										
Test for overall effect:	Z = 0.04 (P = 0.04)	.97)									
Test for subgroup diffe	erences: Not app	licable									
					ï						
					-1	-0.5	0	0.5	T		
				F	avors t	reatment		Favors co	ontrol		

Analysis 6.1.

Comparison 6 DMPA 150 mg/mL versus DMPA 450 mg/mL, Outcome 1 Mean weight change (kg) at 6 months.

Review: Progestin-only contraceptives: effects on weight

Comparison: 7 DMPA 150 mg/mL initiation after pregnancy: interval (≥ 20 weeks) versus postpartum (5 to 8 weeks)

Outcome: I Mean weight gain (lb) at I year

Study or subgroup	Treatment		Control		D	Mean lifference		Weight	Mean Difference
	Ν	Mean(SD)	Ν	Mean(SD)	IV,Fi	xed,95% Cl			IV,Fixed,95% CI
Espey 2000	115	9.3 (11)	57	7 (9.8)				100.0 %	2.30 [-0.94, 5.54]
Total (95% CI)	115		57			-		100.0 %	2.30 [-0.94, 5.54]
Heterogeneity: not ap	plicable								
Test for overall effect:	Z = 1.39 (P = 0.	16)							
Test for subgroup diffe	erences: Not appl	icable							
							1		
					-10 -5	0 5	10		
				Fav	vors treatment	Favors of	ontrol		

Analysis 7.1.

Comparison 7 DMPA 150 mg/mL initiation after pregnancy: interval (20 weeks) versus postpartum (5 to 8 weeks), Outcome 1 Mean weight gain (lb) at 1 year.

Review: Progestin-only contraceptives: effects on weight

Comparison: 7 DMPA 150 mg/mL initiation after pregnancy: interval (≥ 20 weeks) versus postpartum (5 to 8 weeks)

Outcome: 2 Mean weight gain (lb) at 2 years

Study or subgroup	Treatment N	Mean(SD)	Control N	Mean(SD)			ffere	ean nce 5% Cl		Weight	Mean Difference IV,Fixed,95% Cl
Espey 2000	35	15.8 (13.2)	29	14.2 (12.8)			-		-	100.0 %	1.60 [-4.79, 7.99]
Total (95% CI)	35		29				-		-	100.0 %	1.60 [-4.79, 7.99]
Heterogeneity: not ap	plicable										
Test for overall effect:	Z = 0.49 (P = 0.4)	62)									
Test for subgroup diffe	erences: Not appl	icable									
									- 1		
					-10	-5	0	5	10		
				Fa	avors treat	tment		Favors co	ontrol		

Analysis 7.2.

Comparison 7 DMPA 150 mg/mL initiation after pregnancy: interval (20 weeks) versus postpartum (5 to 8 weeks), Outcome 2 Mean weight gain (lb) at 2 years.

Review: Progestin-only contraceptives: effects on weight

Comparison: 8 DMPA-IM 150 mg versus DMPA-SC 104 mg

Outcome: I Mean weight change (kg) at 36 months

Study or subgroup	Treatment		Control		Diffe	Mean erence		Weight	Mean Difference
	Ν	Mean(SD)	Ν	Mean(SD)	IV,Fixe	ed,95% Cl			IV,Fixed,95% CI
Westhoff 2007	56	5.8 (8.7)	65	4.5 (8.5)	_			100.0 %	1.30 [-1.78, 4.38]
Total (95% CI)	56		65		-	-		100.0 %	1.30 [-1.78, 4.38]
Heterogeneity: not ap	plicable								
Test for overall effect:	Z = 0.83 (P = 0.4)	41)							
Test for subgroup diffe	erences: Not appl	icable							
					-10 -5	0 5	10		
				Fav	vors treatment	Favors c	ontrol		

Analysis 8.1.

Comparison 8 DMPA-IM 150 mg versus DMPA-SC 104 mg, Outcome 1 Mean weight change (kg) at 36 months.

Review: Progestin-only contraceptives: effects on weight

Comparison: 9 DMPA 150 mg/mL versus NET-EN 200 mg (60-day intervals)

Outcome: I Mean weight change (kg) at I2 months

Study or subgroup	Treatment		Control			D	Me Differer			Weight	Mean Difference
	Ν	Mean(SD)	Ν	Mean(SD)		IV,F	ixed,9.	5% CI			IV,Fixed,95% CI
WHO 1983	764	1.9 (8.29)	398	1.7 (5.98)		-	-	_		100.0 %	0.20 [-0.63, 1.03]
Total (95% CI)	764		398			_	-			100.0 %	0.20 [-0.63, 1.03]
Heterogeneity: not ap	plicable										
Test for overall effect:	Z = 0.47 (P = 0.	64)									
Test for subgroup diffe	rences: Not appl	licable									
					i	. T			Ţ.		
					-2	- [0	I	2		
				Fa	avors tr	eatment		Favors c	ontrol		

Analysis 9.1.

Comparison 9 DMPA 150 mg/mL versus NET-EN 200 mg (60-day intervals), Outcome 1 Mean weight change (kg) at 12 months.

Review: Progestin-only contraceptives: effects on weight

Comparison: 9 DMPA 150 mg/mL versus NET-EN 200 mg (60-day intervals)

Outcome: 2 Mean weight change (kg) at 24 months

Study or subgroup	Treatment		Control			D	Me ifferer	ean nce		Weight	Mean Difference
	Ν	Mean(SD)	Ν	Mean(SD)		IV,Fi	xed,9	5% CI			IV,Fixed,95% CI
WHO 1983	390	3.3 (9.87)	214	3.3 (7.31)						100.0 %	0.0 [-1.39, 1.39]
Total (95% CI)	390		214							100.0 %	0.0 [-1.39, 1.39]
Heterogeneity: not ap	plicable										
Test for overall effect:	Z = 0.0 (P = 1.0)										
Test for subgroup diffe	erences: Not appli	cable									
					ī						
					-2	-1	0	1	2		
				Fav	vors tre	eatment		Favors c	ontrol		

Analysis 9.2.

Comparison 9 DMPA 150 mg/mL versus NET-EN 200 mg (60-day intervals), Outcome 2 Mean weight change (kg) at 24 months.

Review: Progestin-only contraceptives: effects on weight

Comparison: 9 DMPA 150 mg/mL versus NET-EN 200 mg (60-day intervals)

Outcome: 3 Mean weight change at I year

Study or subgroup	Treatment		Control			Mean Difference		Weight	Mean Difference
	Ν	Mean(SD)	Ν	Mean(SD)	IV,	Fixed,95% C			IV,Fixed,95% CI
Salem 1988	107	3.5 (3.4)	94	2.7 (3.1)				100.0 %	0.80 [-0.10, 1.70]
Total (95% CI)	107		94					100.0 %	0.80 [-0.10, 1.70]
Heterogeneity: not ap	plicable								
Test for overall effect:	Z = 1.74 (P = 0.)	081)							
Test for subgroup diffe	rences: Not app	licable							
					<u>і </u>				
					-2 -1	0 I	2		
				Fav	vors treatmen	t Favor	s control		

Analysis 9.3.

Comparison 9 DMPA 150 mg/mL versus NET-EN 200 mg (60-day intervals), Outcome 3 Mean weight change at 1 year.

Review: Progestin-only contraceptives: effects on weight

Comparison: 10 NET-EN 200 mg: 60-day intervals versus 3 intervals of 60 days then 84-day intervals

Outcome: I Mean weight change (kg) at I2 months

Study or subgroup	Treatment		Control			l Differ	Mean rence		Weight	Mean Difference
	Ν	Mean(SD)	Ν	Mean(SD)		IV,Fixed	I,95% CI			IV,Fixed,95% CI
WHO 1983	398	1.7 (5.98)	424	1.7 (4.12)		-	-		100.0 %	0.0 [-0.71, 0.71]
Total (95% CI)	398		424						100.0 %	0.0 [-0.71, 0.71]
Heterogeneity: not ap	plicable									
Test for overall effect:	Z = 0.0 (P = 1.0)									
Test for subgroup diffe	rences: Not appl	cable								
					-2 -	0	I	2		
				Fav	ors treatn	nent	Favors of	ontrol		

Analysis 10.1.

Comparison 10 NET-EN 200 mg: 60-day intervals versus 3 intervals of 60 days then 84-day intervals, Outcome 1 Mean weight change (kg) at 12 months.

Review: Progestin-only contraceptives: effects on weight

Comparison: 10 NET-EN 200 mg: 60-day intervals versus 3 intervals of 60 days then 84-day intervals

Outcome: 2 Mean weight change (kg) at 24 months

Study or subgroup	Treatment		Control			D	Miffere	ean hce		Weight	Mean Difference
	Ν	Mean(SD)	Ν	Mean(SD)		IV,F	ixed,9	5% CI			IV,Fixed,95% CI
WHO 1983	214	3.3 (7.31)	239	3.4 (6.18)			-			100.0 %	-0.10 [-1.35, 1.15]
Total (95% CI)	214		239				-			100.0 %	-0.10 [-1.35, 1.15]
Heterogeneity: not ap	plicable										
Test for overall effect:	Z = 0.16 (P = 0.16)	88)									
Test for subgroup diffe	erences: Not app	licable									
					,	- 1		ï	- 1		
					-2	- [0	T	2		
				F	avors ti	reatment		Favors o	ontrol		

Analysis 10.2.

Comparison 10 NET-EN 200 mg: 60-day intervals versus 3 intervals of 60 days then 84-day intervals, Outcome 2 Mean weight change (kg) at 24 months.

Review: Progestin-only contraceptives: effects on weight

Comparison: 11 DMPA 150 mg/mL versus Cu IUC

Outcome: I Mean weight change (kg) at 10 years

Study or subgroup	Treatment N	Mean(SD)	Control N	Mean(SD)	Mean Difference IV.Fixed.95%	CI	Weight	Mean Difference IV.Fixed,95% Cl
	1.8	r leari(SD)		r lean(SD)	14,11XC0,7570			11,11,11,11,11,11,11,11
Taneepanichskul 1998	50	10.9 (1.2)	50	11.2 (1.5)			100.0 %	-0.30 [-0.83, 0.23]
Total (95% CI)	50		50		-		100.0 %	-0.30 [-0.83, 0.23]
Heterogeneity: not applical	ble							
Test for overall effect: $Z =$	1.10 (P = 0.27)							
Test for subgroup difference	es: Not applicat	ble						
				-2	-1 0	1 2		
				Favor	s treatment Fav	ors control		

Analysis 11.1.

Comparison 11 DMPA 150 mg/mL versus Cu IUC, Outcome 1 Mean weight change (kg) at 10 years.

Review: Progestin-only contraceptives: effects on weight

Comparison: 11 DMPA 150 mg/mL versus Cu IUC

Outcome: 2 Mean weight change (kg) at I year by baseline BMI

Study or subgroup	Treatment		Control		Mean Difference	Weight	Mean Difference
	Ν	Mean(SD)	Ν	Mean(SD)	IV,Fixed,95% CI		IV,Fixed,95% Cl
BMI < 25 kg/m ²							
Pantoja 2010	226	2.2 (3.4)	226	-0.3 (3.1)		66.7 %	2.50 [1.90, 3.10]
Subtotal (95% CI)	226		226		-	66. 7 %	2.50 [1.90, 3.10]
Heterogeneity: not applical	ole						
Test for overall effect: Z =	8.17 (P < 0.000	01)					
2 BMI 25 to 29.9 kg/m 2							
Pantoja 2010	109	1.4 (3.6)	109	-0.7 (3.5)		27.0 %	2.10 [1.16, 3.04]
Subtotal (95% CI)	109		109			27.0 %	2.10 [1.16, 3.04]
Heterogeneity: not applical	ole						
Test for overall effect: Z =	4.37 (P = 0.000	013)					
3 BMI \geq 30 kg/m 2							
Pantoja 2010	44	0.4 (4.8)	44	-0.3 (4.6)		6.2 %	0.70 [-1.26, 2.66]
Subtotal (95% CI)	44		44			6.2 %	0.70 [-1.26, 2.66]
Heterogeneity: not applical	ole						
Test for overall effect: Z =	0.70 (P = 0.48)						
Total (95% CI)	379		379		-	100.0 %	2.28 [1.79, 2.77]
Heterogeneity: Chi ² = 3.14	4, df = 2 (P = 0.2)	21); I ² =36%					
Test for overall effect: $Z =$	9.12 (P < 0.000	OI)					
Test for subgroup differenc	es: Chi ² = 3.14,	df = 2 (P = 0.2)), I ² =36%				

Favors treatment Favors control

Analysis 11.2.

Comparison 11 DMPA 150 mg/mL versus Cu IUC, Outcome 2 Mean weight change (kg) at 1 year by baseline BMI.

Review: Progestin-only contraceptives: effects on weight

Comparison: 11 DMPA 150 mg/mL versus Cu IUC

Outcome: 3 Mean weight change (kg) at 2 years by baseline BMI

Mea Differenc	Weight	Mean Difference		Control		Treatment	Study or subgroup
IV,Fixed,95% C		IV,Fixed,95% CI	Mean(SD)	Ν	Mean(SD)	Ν	
							I BMI < 25 kg/m ²
2.70 [2.02, 3.38	74.4 %		0.8 (3.7)	226	3.5 (3.7)	226	Pantoja 2010
2.70 [2.02, 3.38	74.4 %	•		226		226	Subtotal (95% CI)
						ble	Heterogeneity: not applicat
)))	7.76 (P < 0.0000	Test for overall effect: Z = 1 2 BMI 25 to 29.9 kg/m ²
3.00 [1.73, 4.27	21.3 %		-0.2 (4.9)	109	2.8 (4.7)	109	Pantoja 2010
3.00 [1.73, 4.27	21.3 %	•		109		109	Subtotal (95% CI)
						ble	Heterogeneity: not applicat
))	4.61 (P < 0.0000	Test for overall effect: Z = $\frac{1}{2}$ 3 BMI \geq 30 kg/m ²
1.50 [-1.33, 4.33	4.3 %		-0.1 (7.7)	44	1.4 (5.7)	44	Pantoja 2010
1.50 [-1.33, 4.33	4.3 %	-		44		44	Subtotal (95% CI)
						ble	Heterogeneity: not applicat
						I.04 (P = 0.30)	Test for overall effect: $Z =$
2.71 [2.12, 3.30	100.0 %	•		379		379	Total (95% CI)
					54); I ² =0.0%), df = 2 (P = 0.6	Heterogeneity: $Chi^2 = 0.90$
)))	9.03 (P < 0.0000	Test for overall effect: $Z = 2$
				, I ² =0.0%	df = 2 (P = 0.64)	es: Chi ² = 0.90,	Test for subgroup difference

-10 -5 0 5 10 Favors treatment Favors control

Analysis 11.3.

Comparison 11 DMPA 150 mg/mL versus Cu IUC, Outcome 3 Mean weight change (kg) at 2 years by baseline BMI.

Review: Progestin-only contraceptives: effects on weight

Comparison: 11 DMPA 150 mg/mL versus Cu IUC

Outcome: 4 Mean weight change (kg) at 3 years by baseline BMI

Study or subgroup Tr	reatment N	Mean(SD)	Control N	Mean(SD)	Mean Difference IV.Fixed,95% CI	Weight	Mean Difference IV,Fixed,95% Cl
	IN	rieari(5D)	IN	Triedin(SD)	IV,FIXEd,75% CI		IV, FIXED, 7576 CI
MI < 25 kg/m ²							
Pantoja 2010	226	4.5 (4.5)	226	1.2 (4)		71.6 %	3.30 [2.52, 4.08]
btotal (95% CI)	226		226		•	71.6 %	3.30 [2.52, 4.08]
terogeneity: not applicable							
t for overall effect: Z = 8.24 Ml 25 to 29.9 kg/m ²	(P < 0.0000)))					
Pantoja 2010	109	3.4 (5.5)	109	0.2 (4.9)		23.1 %	3.20 [1.82, 4.58]
btotal (95% CI)	109		109		•	23.1 %	3.20 [1.82, 4.58]
terogeneity: not applicable							
t for overall effect: Z = 4.54 MI \geq 30 kg/m ²	(P < 0.0000)))					
Pantoja 2010	44	1.9 (6.7)	44	0.6 (7)		5.4 %	1.30 [-1.56, 4.16]
btotal (95% CI)	44		44		-	5.4 %	1.30 [-1.56, 4.16]
terogeneity: not applicable							
t for overall effect: $Z = 0.89$	(P = 0.37)						
tal (95% CI)	379		379		•	100.0 %	3.17 [2.51, 3.83]
terogeneity: Chi ² = 1.75, df	= 2 (P = 0.4	42); I ² =0.0%					
t for overall effect: Z = 9.35	(P < 0.0000	01)					
t for subgroup differences: ($Chi^2 = 1.75,$	df = 2 (P = 0.42)), l ² =0.0%				

Favors treatment Favors control

Analysis 11.4.

Comparison 11 DMPA 150 mg/mL versus Cu IUC, Outcome 4 Mean weight change (kg) at 3 years by baseline BMI.

Mean weight change (kg) by year

Study	Year	Mean change ± SD DMPA	Mean change ± SD IUC
Pantoja 2010	1	1.76 ± 3.6	-0.42 ± 3.4
Pantoja 2010	2	3.1 ± 4.3	0.4 ± 4.7
Pantoja 2010	3	3.9 ± 5.1	0.8 ± 4.7

Analysis 11.5.

Comparison 11 DMPA 150 mg/mL versus Cu IUC, Outcome 5 Mean weight change (kg) by year.

Mean changes in body composition by 12 months

Study	Body composi- tion assessment	Reported mean ± SD DMPA	N	Reported mean ± SD Cu T380A IUC	N	Reported P
Dal'Ava 2014	Total body mass (kg)	1.9 ± 3.5	26	1.1 ± 3.2	26	0.38
Dal'Ava 2014	Total fat mass (kg)	1.6 ±3.4	26	-0.9 ± 7.2	26	0.14
Dal'Ava 2014	Total lean mass (kg)	0.3 ± 1.8	26	1.2 ± 2.3	26	0.11
Dos Santos 2014	Total body mass (kg)	1.4 ± 3.13	20	0.3 ± 2.24	20	0.183
Dos Santos 2014	Total body fat (kg)	1.57 ± 3.29	20	0.52 ± 2.5	20	0.256
Dos Santos 2014	Total lean mass (kg)	-0.31 ± 1.7	20	-0.26 ± 0.94	20	0.909

Analysis 11.6.

Comparison 11 DMPA 150 mg/mL versus Cu IUC, Outcome 6 Mean changes in body composition by 12 months.

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Mean	weight change	(kg) a	at 1	year
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Study	Method	Reported adjusted mean ± SE	N	Reported P
Modesto 2015	DMPA	1.3 ± 0.15	675	< 0.0001
Modesto 2015	LNG-IUS	0.7 ± 0.18	602	0.1719
Modesto 2015	Cu IUC	0.2 ± 0.17	602	Referent

Analysis 12.1.

Comparison 12 DMPA, ENG implant, or LNG-IUC versus Cu IUC, Outcome 1 Mean weight change (kg) at 1 year.

Mean weight change (kg) at 4 years

Study	Method	Reported adjusted mean ± SE	N	Reported P
Modesto 2015	DMPA	3.5 ± 0.23	540	< 0.0001
Modesto 2015	LNG-IUS	2.7 ± 0.27	563	0.2310
Modesto 2015	Cu IUC	1.9 ± 0.23	625	Referent

Analysis 12.2.

Comparison 12 DMPA, ENG implant, or LNG-IUC versus Cu IUC, Outcome 2 Mean weight change (kg) at 4 years.

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Mean	weight	change	(kg)	at 10 years
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Study	Method	Reported adjusted mean ± SE	N	Reported P
Modesto 2015	DMPA	6.6 ± 0.61	125	0.0350
Modesto 2015	LNG-IUS	4.0 ± 0.97	68	0.3475
Modesto 2015	Cu-IUD	4.9 ± 0.60	154	Referent

Analysis 12.3.

Comparison 12 DMPA, ENG implant, or LNG-IUC versus Cu IUC, Outcome 3 Mean weight change (kg) at 10 years.

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Weight change	(kg)	at	12	months	
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Study	Method	Reported adjusted beta (95% CI)	N	Reported P
Vickery 2013	DMPA	1.37 (-0.44 to 3.18)	67	0.14
Vickery 2013	ENG implant	1.37 (-0.16 to 2.91)	130	0.08
Vickery 2013	LNG-IUC	0.46 (-1.04 to 1.95)	130	0.55
Vickery 2013	Cu IUC	Referent	100	Referent

Analysis 12.4.

Comparison 12 DMPA, ENG implant, or LNG-IUC versus Cu IUC, Outcome 4 Weight change (kg) at 12 months.

Review: Progestin-only contraceptives: effects on weight Comparison: 13 DMPA 150 mg/mL versus no hormonal method Outcome: I Mean change in total body fat (%) at 6 months

Study or subgroup	DMPA N	Mean(SD)	No hormonal N	Mean(SD)		Mean ference ed,95% Cl	Weight	Mean Difference IV,Fixed,95% Cl
Bonny 2009	8	10.3 (10.49)	18	-0.7 (8.94)			100.0 %	.00 [2.64, 9.36]
Total (95% CI)	8		18				- 100.0 %	11.00 [2.64, 19.36]
Heterogeneity: not ap	plicable							
Test for overall effect:	Z = 2.58 (P	= 0.0099)						
Test for subgroup diffe	erences: No	t applicable						
					1 1			
					-20 -10	0 10	20	
					Favors DMPA	Favors no	hormonal	

Analysis 13.1.

Comparison 13 DMPA 150 mg/mL versus no hormonal method, Outcome 1 Mean change in total body fat (%) at 6 months.

Review: Progestin-only contraceptives: effects on weight Comparison: 13 DMPA 150 mg/mL versus no hormonal method Outcome: 2 Mean change in lean body mass (%) at 6 months

Study or subgroup	DMPA N	Mean(SD)	No hormonal N	Mean(SD)			Me fferen ked,95			Weight	Mean Difference IV,Fixed,95% Cl
Bonny 2009	8	-3.4 (3.57)	18	0.6 (3.39)	•					100.0 %	-4.00 [-6.93, -1.07]
Total (95% CI)	8		18							100.0 %	-4.00 [-6.93, -1.07]
Heterogeneity: not ap	plicable										
Test for overall effect:	Z = 2.68 (P	= 0.0074)									
Test for subgroup diffe	erences: Not	applicable									
					1						
					-4	-2	0	2	4		
				Favo	rs no ha	ormonal		Favors I	OMPA		

Analysis 13.2.

Comparison 13 DMPA 150 mg/mL versus no hormonal method, Outcome 2 Mean change in lean body mass (%) at 6 months.

Review: Progestin-only contraceptives: effects on weight Comparison: 13 DMPA 150 mg/mL versus no hormonal method Outcome: 3 Mean change in weight (lb) at 12 months postpartum

Study or subgroup	DMPA N	Mean(SD)	Sterilization N	Mean(SD)	M Differe IV,Fixed,9		Weight	Mean Difference IV,Fixed,95% Cl
Nyirati 2013	35	2.14 (13.39)	12	0.6 (15.7)	-	-	100.0 %	1.54 [-8.39, 11.47]
Total (95% CI)	35		12		+	-	100.0 %	1.54 [-8.39, 11.47]
Heterogeneity: not ap	plicable							
Test for overall effect:	Z = 0.30 (P	= 0.76)						
Test for subgroup diffe	rences: Not	applicable						
							1	
					-50 -25 0	25	50	
					Favors DMPA	Favors s	terilization	

Analysis 13.3.

Comparison 13 DMPA 150 mg/mL versus no hormonal method, Outcome 3 Mean change in weight (lb) at 12 months postpartum.

Review: Progestin-only contraceptives: effects on weight Comparison: 13 DMPA 150 mg/mL versus no hormonal method Outcome: 4 Mean change in BMI (kg/m²) at 12 months postpartum

Study or subgroup	DMPA N	Mean(SD)	Sterilization N	Mean(SD)			iffere	lean ence 95% Cl		Weight	Mean Difference IV,Fixed,95% Cl
Nyirati 2013	35	0.36 (2.33)	12	0.04 (2.71)		_	+			100.0 %	0.32 [-1.40, 2.04]
Total (95% CI)	35		12				-			100.0 %	0.32 [-1.40, 2.04]
Heterogeneity: not app	olicable										
Test for overall effect:	Z = 0.37 (P	= 0.71)									
Test for subgroup diffe	rences: Not	applicable									
					ï			i.	T		
					-4	-2	0	2	4		
					Favor	s DMPA		Favors st	erilizatio	on	

Analysis 13.4.

Comparison 13 DMPA 150 mg/mL versus no hormonal method, Outcome 4 Mean change in BMI (kg/m2) at 12 months postpartum.

Review: Progestin-only contraceptives: effects on weight Comparison: 13 DMPA 150 mg/mL versus no hormonal method Outcome: 5 Mean change in body fat (%) at 12 months postpartum

Study or subgroup	DMPA N	Mean(SD)	Sterilization N	Mean(SD)	Mean Difference IV,Fixed,95% Cl	Weight	Mean Difference IV,Fixed,95% Cl
Nyirati 2013	35	-0.58 (3.88)	12	-1.56 (5.51)	-	100.0 %	0.98 [-2.39, 4.35]
Total (95% CI)	35		12		•	100.0 %	0.98 [-2.39, 4.35]
Heterogeneity: not app	olicable						
Test for overall effect:	Z = 0.57 (P	= 0.57)					
Test for subgroup diffe	rences: Not	applicable					
					-20 -10 0 10 20		
					Favors DMPA Favours steriliz	zation	

Analysis 13.5.

Comparison 13 DMPA 150 mg/mL versus no hormonal method, Outcome 5 Mean change in body fat (%) at 12 months postpartum.

Review: Progestin-only contraceptives: effects on weight Comparison: 14 Norplant versus non-hormonal IUC Outcome: I Mean weight change (kg) at 6 months

Study or subgroup	Treatment		Control		-	Mean ference	Weight	Mean Difference
	N	Mean(SD)	N	Mean(SD)	IV,Fixe	ed,95% Cl		IV,Fixed,95% CI
Salem 1984	49	1.39 (0.53)	50	0.92 (0.34)			100.0 %	0.47 [0.29, 0.65]
Total (95% CI)	49		50				100.0 %	0.47 [0.29, 0.65]
Heterogeneity: not app	plicable							
Test for overall effect:	Z = 5.24 (P < 0.0	(1000						
Test for subgroup diffe	rences: Not appl	icable						
					-0.5 -0.25	0 0.25 0.5		
				I	Favors treatment	Favors control		

Analysis 14.1.

Comparison 14 Norplant versus non-hormonal IUC, Outcome 1 Mean weight change (kg) at 6 months.

Review: Progestin-only contraceptives: effects on weight Comparison: 14 Norplant versus non-hormonal IUC Outcome: 2 Mean weight change (kg) at 1 year

Study or subgroup	Treatment N	Mean(SD)	Control N	Mean(SD)		Mean erence ed,95% Cl	Weight	Mean Difference IV,Fixed,95% Cl
Sule 2005	188	2.5 (3.6)	136	1.4 (3.2)			100.0 %	1.10 [0.36, 1.84]
Total (95% CI)	188		136			-	100.0 %	1.10 [0.36, 1.84]
Heterogeneity: not app	olicable							
Test for overall effect: 2	Z = 2.90 (P = 0.0)	0038)						
Test for subgroup diffe	rences: Not appl	icable						
					-2 -1	0 1 2		
				E	avors treatment	Favors control		

Analysis 14.2.

Comparison 14 Norplant versus non-hormonal IUC, Outcome 2 Mean weight change (kg) at 1 year.

Review: Progestin-only contraceptives: effects on weight Comparison: 14 Norplant versus non-hormonal IUC Outcome: 3 Mean weight change (kg) at 3 years

Study or subgroup	Treatment N	Mean(SD)	Control N	Mean(SD)			ffere	ean nce 5% Cl		Weight	Mean Difference IV,Fixed,95% Cl
Sule 2005	130	4.8 (4.3)	60	3.9 (4.2)		-			→	100.0 %	0.90 [-0.39, 2.19]
Total (95% CI)	130		60							100.0 %	0.90 [-0.39, 2.19]
Heterogeneity: not app	olicable										
Test for overall effect:	Z = 1.36 (P = 0.)	17)									
Test for subgroup diffe	rences: Not appl	icable									
					-2	-1	0	1	2		
				F	avors trea	atment		Favors co	ontrol		

Analysis 14.3.

Comparison 14 Norplant versus non-hormonal IUC, Outcome 3 Mean weight change (kg) at 3 years.

Review: Progestin-only contraceptives: effects on weight

Comparison: 15 Norplant versus barrier, 'local', or no contraceptive method

Outcome: I Mean weight change (kg) at 6 months

Study or subgroup	Treatment		Control		-	Mean erence	Weight	Mean Difference
	N	Mean(SD)	Ν	Mean(SD)	IV,Fixe	ed,95% Cl		IV,Fixed,95% CI
Salem 1984	49	1.39 (0.53)	48	0.65 (0.56)			100.0 %	0.74 [0.52, 0.96]
Total (95% CI)	49		48			-	100.0 %	0.74 [0.52, 0.96]
Heterogeneity: not ap	plicable							
Test for overall effect:	Z = 6.68 (P < 0.01)	(1000						
Test for subgroup diffe	rences: Not appl	icable						
					-1 -0.5	0 0.5 I		
				F	avors treatment	Favors control		

Analysis 15.1.

Comparison 15 Norplant versus barrier, 'local', or no contraceptive method, Outcome 1 Mean weight change (kg) at 6 months.

Review: Progestin-only contraceptives: effects on weight Comparison: 16 Norplant versus DMPA 150 mg/mL Outcome: 1 Mean weight change (kg) at 1 year

Study or subgroup	Treatment		Control				iffere	lean ence		Weight	Mean Difference
	Ν	Mean(SD)	N	Mean(SD)		IV,Fi	xed,	95% CI			IV,Fixed,95% CI
Moore 1995	50	-0.81 (3.1)	50	0.06 (1.8)	-	-	+			100.0 %	-0.87 [-1.86, 0.12]
Total (95% CI)	50		50				-			100.0 %	-0.87 [-1.86, 0.12]
Heterogeneity: not app	plicable										
Test for overall effect: 2	Z = 1.72 (P = 0.1)	086)									
Test for subgroup diffe	rences: Not app	licable									
					ī			ï			
					-2	- [0	I.	2		
				F	avors tr	eatment		Favors c	ontrol		

Analysis 16.1.

Comparison 16 Norplant versus DMPA 150 mg/mL, Outcome 1 Mean weight change (kg) at 1 year.

Review: Progestin-only contraceptives: effects on weight Comparison: 17 Norplant versus 2-rod LNG Outcome: I Mean weight change (kg) at I year

Study or subgroup	Treatment	Marri (CD)	Control	Marra (CD)		Mean Difference	Weight	Mean Difference
	Ν	Mean(SD)	Ν	Mean(SD)	IV,F	ixed,95% Cl		IV,Fixed,95% CI
Sivin 1998	596	0.99 (3.66)	600	0.9 (3.67)		-	100.0 %	0.09 [-0.33, 0.51]
Total (95% CI)	596		600			•	100.0 %	0.09 [-0.33, 0.51]
Heterogeneity: not app	olicable							
Test for overall effect: 2	Z = 0.42 (P = 0.	67)						
Test for subgroup diffe	rences: Not app	licable						
					-2 -1	0 1	2	
				Fa	vors treatment	Favors con	trol	

Analysis 17.1.

Comparison 17 Norplant versus 2-rod LNG, Outcome 1 Mean weight change (kg) at 1 year.

Review: Progestin-only contraceptives: effects on weight

Comparison: 17 Norplant versus 2-rod LNG Outcome: 2 Mean weight change (kg) at 3 years

Study or subgroup	Treatment		Control			Me Differer			Weight	Mean Difference
	Ν	Mean(SD)	Ν	Mean(SD)	IV	V,Fixed,9	5% CI			IV,Fixed,95% CI
Sivin 1998	462	3.12 (4.51)	460	3.12 (4.72)		-	-		100.0 %	0.0 [-0.60, 0.60]
Total (95% CI)	462		460			-	-		100.0 %	0.0 [-0.60, 0.60]
Heterogeneity: not app	licable									
Test for overall effect: Z	C = 0.0 (P = 1.0))								
Test for subgroup differ	ences: Not appl	icable								
					-2 -1	0	1	2		
				Fav	ors treatme	nt	Favors co	ontrol		



Comparison 17 Norplant versus 2-rod LNG, Outcome 2 Mean weight change (kg) at 3 years.

Review: Progestin-only contraceptives: effects on weight Comparison: 17 Norplant versus 2-rod LNG Outcome: 3 Mean weight change (kg) at 5 years

Study or subgroup	Treatment		Control			_	iffen	1ean ence		Weight	Mean Difference
	Ν	Mean(SD)	N	Mean(SD)		IV,Fi	xed,	95% CI			IV,Fixed,95% CI
Sivin 1998	302	4.14 (4.69)	312	3.54 (4.59)			+	-		100.0 %	0.60 [-0.13, 1.33]
Total (95% CI)	302		312							100.0 %	0.60 [-0.13, 1.33]
Heterogeneity: not app	olicable										
Test for overall effect: 2	Z = 1.60 (P = 0.	11)									
Test for subgroup diffe	rences: Not app	licable									
					ī.			ī	ī		
					-2	-1	0	1	2		
				E	avors tr	reatment		Favors co	ntrol		

Analysis 17.3.

Comparison 17 Norplant versus 2-rod LNG, Outcome 3 Mean weight change (kg) at 5 years.

Review: Progestin-only contraceptives: effects on weight Comparison: 18 Norplant versus COC

Outcome: I Mean weight change (kg) at I year

Study or subgroup	Treatment		Control			Mean ference	Weight	Mean Difference
	N	Mean(SD)	Ν	Mean(SD)	IV,Fix	ed,95% Cl		IV,Fixed,95% CI
Sule 2005	188	2.5 (3.6)	38	1.4 (3.5)			100.0 %	1.10 [-0.13, 2.33]
Total (95% CI)	188		38				100.0 %	1.10 [-0.13, 2.33]
Heterogeneity: not app	olicable							
Test for overall effect:	Z = 1.76 (P = 0.1)	079)						
Test for subgroup diffe	rences: Not appl	icable						
					-2 -1	0 1 2		
				Fa	vors treatment	Favors control		

Analysis 18.1.

Comparison 18 Norplant versus COC, Outcome 1 Mean weight change (kg) at 1 year.

Study	Body composi- tion assessment	Reported mean ± SD LNG-IUC	N	Reported mean ± SD Cu T380A IUC	N	Reported P
Dal'Ava 2012	Change in weight (kg)	2.9 + 5.7	38	1.4 + 4.4	38	0.068
Dal'Ava 2012	Percent change in fat mass	2.5 + 8.0	38	-1.3 + 6.9	38	0.029
Dal'Ava 2012	Percent change in lean mass	-1.4 + 4.7	38	1.0 + 3.8	38	0.027

Mean changes in body composition by 12 months

Analysis 19.1.

Comparison 19 LNG-IUC versus no hormonal contraceptive, Outcome 1 Mean changes in body composition by 12 months.

Review: Progestin-only contraceptives: effects on weight Comparison: 19 LNG-IUC versus no hormonal contraceptive Outcome: 2 Mean change in weight (kg) at 1 year

Study or subgroup	Treatment N	Mean(SD)	Control N	Mean(SD)			fere	lean ence 95% Cl		Weight	Mean Difference IV,Fixed,95% Cl
Napolitano 2015	34	0.6 (2.1)	26	-0.2 (3.6)					→	100.0 %	0.80 [-0.75, 2.35]
Total (95% CI)	34		26			-			-	100.0 %	0.80 [-0.75, 2.35]
Heterogeneity: not ap	plicable										
Test for overall effect:	Z = 1.01 (P = 0.)	31)									
Test for subgroup diffe	rences: Not appl	icable									
					-2	-1	0	1	2		
				F	avors trea	tment		Favors co	ntrol		

Analysis 19.2.

Comparison 19 LNG-IUC versus no hormonal contraceptive, Outcome 2 Mean change in weight (kg) at 1 year.

Review: Progestin-only contraceptives: effects on weight Comparison: 19 LNG-IUC versus no hormonal contraceptive Outcome: 3 Mean change in BMI (kg/m²) at 1 year

Study or subgroup	Treatment N	Mean(SD)	Control N	Mean(SD)	I	N Differe V,Fixed,			Weight	Mean Difference IV,Fixed,95% Cl
Napolitano 2015	34	0.2 (0.7)	26	0.5 (1)					100.0 %	-0.30 [-0.75, 0.15]
Total (95% CI)	34		26			-			100.0 %	-0.30 [-0.75, 0.15]
Heterogeneity: not app	plicable									
Test for overall effect:	Z = 1.30 (P = 0.	19)								
Test for subgroup diffe	erences: Not app	licable								
					-2 -1	0	1	2		
				Fa	avors treatme	ent	Favors o	ontrol		

Analysis 19.3.

Comparison 19 LNG-IUC versus no hormonal contraceptive, Outcome 3 Mean change in BMI (kg/m2) at 1 year.

Review: Progestin-only contraceptives: effects on weight Comparison: 19 LNG-IUC versus no hormonal contraceptive Outcome: 4 Mean change in fat mass (%) at 1 year

Study or subgroup	Treatment N	Mean(SD)	Control N	Mean(SD)		-	Mean ifference xed,95% Cl		Weight	Mean Difference IV,Fixed,95% Cl
Napolitano 2015	34	1.1 (2.9)	26	-0.5 (1.6)			-		100.0 %	1.60 [0.45, 2.75]
Total (95% CI)	34		26				-		100.0 %	1.60 [0.45, 2.75]
Heterogeneity: not app	olicable									
Test for overall effect: 2	Z = 2.72 (P = 0.0)	0065)								
Test for subgroup diffe	rences: Not appl	icable								
						T				
					-4	-2	0 2	4		
				F	avors tr	reatment	Favors o	ontrol		

Analysis 19.4.

Comparison 19 LNG-IUC versus no hormonal contraceptive, Outcome 4 Mean change in fat mass (%) at 1 year.

Review: Progestin-only contraceptives: effects on weight Comparison: 19 LNG-IUC versus no hormonal contraceptive Outcome: 5 Mean change in fat free mass (%) at 1 year

Study or subgroup	Treatment N	Mean(SD)	Control N	Mean(SD)		Mean erence ed,95% Cl	Weight	Mean Difference IV,Fixed,95% Cl
Napolitano 2015	34	-1.1 (2.9)	26	0.5 (1.6)			100.0 %	-1.60 [-2.75, -0.45]
Total (95% CI)	34		26		-		100.0 %	-1.60 [-2.75, -0.45]
Heterogeneity: not ap	plicable							
Test for overall effect:	Z = 2.72 (P = 0)	.0065)						
Test for subgroup diffe	erences: Not app	licable						
					-4 -2 (0 2	4	
					Favors control	Favors treatr	ment	

Analysis 19.5.

Comparison 19 LNG-IUC versus no hormonal contraceptive, Outcome 5 Mean change in fat free mass (%) at 1 year.

Table 1

Evidence quality

Study	Randomiza- tion methods or NRS	NRS: NOS se- lection criterion	NRS: NOS comparability	Follow-up period	Loss or chart re- view	Evidence qual- ity ^a
Ball 1991	-1	NA	NA	-1	-1	Very low
Bonny 2009	-1	-	-	-1	-	Low
Castle 1978	-1	-	-1	-1	-1	Very low (poor)
Dal'Ava 2012	-1	-	-	-	-	Moderate
Dal'Ava 2014	-1	-	-	-	-1	Low
Dos Santos 2014	-1	-	-	-	-1	Low
Espey 2000	-1	-	-	-	-1	Low
Modesto 2015	-1	-	-	-	-1	Low
Moore 1995	-1	-	-	-	-1	Low
Napolitano 2015	-1	-	-1	-	-	Low
Nyirati 2013	-1	-	-1	-	-1	Very low
Pantoja 2010	-1	-	-	-	-1	Low
Salem 1984	-1	-	-1	-1	-	Very low
Salem 1988	-	NA	NA	-	-1	Moderate
Sivin 1998	-	NA	NA	-	-1	Moderate
Sule 2005	-1	-	-1	-	-1	Very low
Taneepanichskul 1998	-1	-1	-	-	-1	Very low
Tankeyoon 1976	-1	-	-	-	-1	Low
Tuchman 2005	-1	-	-1	-	-1	Very low
Vickery 2013	-1	-	-	-	-1	Low
Westhoff 2007	-1	NA	NA	-	-1	Low
WHO 1983	-1	NA	NA	-	-1	Low

^aDowngraded for the following: (1) Risk of bias high for randomization sequence generation or allocation concealment, or no information provided on either, or study not randomized (NRS); (2) NRS: high risk of bias in selection; (3) NRS: no stars for comparability, i.e. not controlling for relevant confounding; (4) follow-up < 12 months for change in weight or BMI; (6) loss to follow-up > 20%, combined loss to follow-up and discontinuation > 50%, major differential losses between groups, or retrospective chart review of selected cases

NA = not applicable

NOS = Newcastle-Ottawa Quality Assessessment Scale

Table 2

Overview of interventions, outcome measures, mean changes

Study	N	Intervention groups	Time frame	Outcome measures	Mean change
Progestin-only oral contraceptive	ntracepti	ive			
Ball 1991	51	NET 350 µg vs lev- onorgestrel 30 µg	6 months	Weight (kg)	0 vs 0.6
Napolitano 2015	68	Desogestrel 75 µg vs no hormonal method	12 months	Weight (kg); BMI (kg/m ²); fat mass (%); fat-free mass (%)	0.3 vs -0.2 ; 0.5 vs 0.5 ; 2.8 vs -0.5^{a} ; -2.8 vs 0.5^{a}
Depot medroxyprogest	terone ac	Depot medroxyprogesterone acetate (DMPA) 150 mg/mL			
Tuchman 2005	222	DMPA vs MPA + E_2C	6 months; 12 months	Weight (kg)	0.6 vs 1.2; 1.7 vs 3.0
		DMPA vs COC	-		0.6 vs 1.1; 1.7 vs 1.0
Tankeyoon 1976	32	DMPA vs COC	12 months	Weight (kg)	1.8 vs 3.1 (estimated)
Bonny 2009	15	DMPA + placebo vs $DMPA + E_2C$	6 months	Total body fat (%); lean body mass (%)	10.3 vs 2.8; -3.4 vs -1.2
Castle 1978	1000	DMPA 150 vs DMPA 450	6 months	Weight (kg)	0.33 vs 0.32
Espey 2000	172	DMPA: interval vs postpartum	1 year; 2 years	Weight (kg)	4.2 vs 3.2; 7.2 vs 6.5
Westhoff 2007	534	DMPA-intramuscular 150 vs DMPA-subcuta- neous 104	3 years	Weight (kg)	5.8 vs 4.5
WHO 1983	3172	DMPA vs NET-EN (60 days)	1 year; 2 years	Weight (kg)	1.9 vs 1.7; 3.3 vs 3.3
		NET-EN: 60 days vs 84 days	-		1.7 vs 1.7; 3.3 vs 3.4
Salem 1988	400	DMPA vs NET-EN	1 year	Weight; no unit (kg or Ib)	3.5 vs 2.7
Taneepanichskul 1998	100	DMPA vs Cu IUC	10 years	Weight (kg)	10.9 vs 11.2
Pantoja 2010	758	DMPA vs Cu IUC	1 year; 2 years; 3 years	Weight (kg)	1.76 vs -0.42; ^a 3.1 vs 0.4; ^a 3.9 vs 0.8 ^a
Modesto 2015	1277	DMPA 150 vs Cu IUC	1 year; 4 years;	Weight (kg)	1.3 vs 0.2; ^{<i>a</i>}

Study	z	Intervention groups	Time frame	Outcome measures	Mean change
			10 years		3.5 vs 1.9; ^{<i>a</i>} 6.6 vs 4.9 ^{<i>a</i>}
Vickery 2013	167	DMPA 150 vs Cu IUC	12 months	Weight (kg)	2.20 vs 0.16
Dal' Ava 2014	110	DMPA 150 vs Cu IUC	12 months	Weight (kg); fat mass (kg); lean mass (kg)	1.9 vs 1.1; 1.6 vs -0.9; 0.3 vs 1.2
Dos Santos 2014	71	DMPA 150 vs Cu IUC	12 months	Weight (kg); body fat (kg); lean mass (kg)	1.4 vs 0.3; 1.57 vs 0.52; -0.31 vs -0.26
Bonny 2009	26	DMPA + placebo vs no hormonal method	6 months	Total body fat (%); lean body mass (%)	10.3 vs $-0.1; a$ -3.4 vs 0.6^{a}
Nyirati 2013	78	DMPA 150 vs surgical sterilization	12 months	Weight (lb); BMI (kg/m ²); body fat (%)	2.14 vs 0.6; 0.36 vs 0.04; -0.58 vs -1.56
Implants					
Salem 1984	150	Norplant vs Cu IUC	6 months	Weight (kg)	1.39 vs 0.92 ^{<i>a</i>}
		Norplant vs non-hor- monal method	6 months		1.39 vs 0.65 ^{<i>a</i>}
Moore 1995	100	Norplant vs DMPA	12 months	Weight (kg)	-0.81 vs 0.06
Sivin 1998	1200	Norplant vs 2-rod lev- onorgestrel	1 year; 3 years; 5 years	Weight (kg)	0.99 vs 0.90; 3.12 vs 3.12; 4.14 vs 3.54
Sule 2005	516	Norplant vs non-hor- monal IUC	1 year; 3 years	Weight (kg)	2.5 vs 1.4; ^{<i>a</i>} 4.8 vs 3.9
		Norplant vs COC			2.5 vs 1.4; 4.8 vs 0.0
Vickery 2013	230	Etonogestrel implant vs Cu IUC	12 months	Weight (kg)	2.12 vs 0.16
Levonorgestrel (LNG)	intraute	Levonorgestrel (LNG) intrauterine contraception (IUC)			
Dal'Ava 2012	76	LNG-IUC vs non-hor-	12 months	Weight (kg)	2.9 vs 1.4
				Total body fat; lean body mass	2.5% vs -1.3%; ^{<i>a</i>} -1.4% vs 1.0% ^{<i>a</i>}
Vickery 2013	230	LNG-IUC vs Cu IUC	12 months	Weight (kg)	1.03 vs 0.16
Modesto 2015	1204	LNG-IUC vs Cu IUC	1 year; 4 years; 10 years	Weight (kg)	0.7 vs 0.2; 2.7 vs 1.9; 4.0 vs 4.9

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- 1					
	Z	Intervention groups	Time frame	Time frame Outcome measures	Mean change
	60	LNG-IUC vs no	12 months	Weight (kg);	0.6 vs -0.2;
		method		BMI (kg/m ²);	0.2 vs 0.5;
				fat mass (%);	$1.1 \text{ vs} - 0.5;^{a}$
				fat-free mass (%)	$-1.1 \text{ vs } 0.5^{a}$
1					

Napolitano 2015

Study

 a Significant difference between comparison groups

Cu IUC: copper intrauterine contraception

COC: combination oral contraception

DMPA: depot medroxyprogesterone acetate

E2C: estradiol cypionate

LNG-IUC: levonorgestrel-releasing intrauterine contraception

MPA: medroxyprogesterone acetate NET: norethisterone NET-EN: norethisterone enanthate

Table 3

Results by contraceptive method

Study	Comparison groups	Mean difference (95% CI)	Quality of evidence
Progestin-only oral co	ntraceptive		•
Ball 1991	NET 350 µg vs levonorgestrel 30 µg	NS	Very low
Napolitano 2015	Desogestrel 75 µg vs no hormonal contraceptive	Fat mass (%) 3.30 (2.08 to 4.52); fat-free mass (%) -3.30 (-4.52 to - 2.08)	Low
Depot medroxyproges	terone acetate (DMPA) 150 mg/mL		•
Comparison: combinati	on contraceptive		
Tankeyoon 1976	DMPA vs COC	NS	Low
Tuchman 2005	DMPA vs COC	NS	Very low
	DMPA vs MPA + E ₂ C	NS	
Bonny 2009	$DMPA + placebo vs DMPA + E_2C$	NS	Low
Comparison: another pr	ogestin-only injectable formulation or	regimen	•
Castle 1978	DMPA 150 vs DMPA 450	NS	Very low (poor)
Espey 2000	DMPA: interval (20 weeks) vs postpartum (5 to 8 weeks)	NS	Low
Westhoff 2007	DMPA-intramuscular 150 vs DMPA-subcutaneous 104	NS	Low
WHO 1983	DMPA vs NET-EN (60 days)	NS	Low
	NET-EN: 60 days vs 84 days	NS	
Salem 1988	DMPA vs NET-EN	NS	Moderate
Comparison: no hormor	nal contraceptive (retrospective studies)	
Taneepanichskul 1998	DMPA vs Cu IUC	NS	Very low
Pantoja 2010	DMPA vs Cu IUC	Weight (kg): 2.28 (1.79 to 2.77); 2.71 (2.12 to 3.30); 3.17 (2.51 to 3.83)	Low
Modesto 2015	DMPA vs Cu IUC	Reported mean weights (kg): 1.3 vs 0.2 (P < 0.0001); 3.5 vs 1.9 (P < 0.0001); 6.6 vs 4.9 (P < 0.0350)	Low
Comparison: no hormon	nal contraceptive (prospective studies)	-	
Vickery 2013	DMPA vs Cu IUC	NS	Low
Dal'Ava 2014	DMPA vs Cu IUC	NS	Low
Dos Santos 2014	DMPA vs Cu IUC	NS	Low
Bonny 2009	DMPA + placebo vs no hormonal	Total body fat (%) 11.00 (2.64 to 19.36); lean body mass (%) -4.00 (-6.93 to -1.07)	Low
Nyirati 2013	DMPA: 6 weeks postpartum vs sterilization	NS	Very low
Implants			
Vickery 2013	ENG implant vs Cu IUC	NS	Low

Study	Comparison groups	Mean difference (95% CI)	Quality of evidence ^a
Salem 1984	Norplant vs other non-hormonal	Weight (kg) 0.74 (0.52 to 0.96)	Very Low
	Norplant vs Cu IUC	Weight (kg) 0.47 (0.29 to 0.65)	
Sule 2005	Norplant vs non-hormonal IUC	Weight (kg) 1.10 (0.36 to 1.84)	Very low
	Norplant vs COC	NS	
Moore 1995	Norplant vs DMPA	NS	Low
Sivin 1998	Norplant vs 2-rod LNG	NS	Moderate
Levonorgestrel-rele	asing IUC vs no hormonal method		•
Dal'Ava 2012	LNG-IUC vs non-hormonal IUC	Reported mean weights (kg): Total body fat (%) 2.5 vs -1.3 (P = 0.029); lean body mass (%) -1.4 vs 1.0 (P = 0.027)	Moderate
Vickery 2013	LNG-IUC vs Cu IUC	NS	Low
Modesto 2015	LNG-IUC vs Cu IUC	NS	Low
Napolitano 2015	LNG-IUC vs no contraceptive	Fat mass (%) 1.60 (0.45 to 2.75); fat-free mass (%) –1.60 (–2.75 to – 0.45)	Low

^a From Table 1

Cu IUC: copper intrauterine contraception

COC: combination oral contraception

DMPA: depot medroxyprogesterone acetate

LNG-IUC: levonorgestrel-releasing intrauterine contraception

MPA: medroxyprogesterone acetate

NET: norethisterone

NET-EN: norethisterone enanthate

 $\mathbf{NS} = \mathbf{no}$ significant difference between groups

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SUMMARY OF FINDINGS FOR THE MAIN COMPARISON [Explanation]

I	DMPA c	ompared	with no	hormonal	contraceptiv	e for	effect on	weight
I	Dunne	umparcu	with no	normonai	contractput	C 101	uncer on	weight

Change in body fat (%) MD 11.00 (2.64 to 19. 26 by 6 months; 36); (Bonny 2009) change in lean body MD -4.00 (-6.93 to -1. (Bonny 2009) mass (%) by 6 months 07 758	Low	DMPA + placebo vs no hormonal:
Change in weight (l_{12}) , MD 2 28 (1 70 to 2 77), 758		adolescents 15 to 18 years old
MD 2.71 (2.12 to 3.30); (Pantoja 2010) 2 years; MD 3.17 (2.51 to 3.83); 3 years	Low	DMPA vs Cu IUC; women of child-bearing age
$\begin{array}{llllllllllllllllllllllllllllllllllll$	Low	DMPA vs Cu IUC; women 18 to 40 years old

may change the estimate.

Low quality: Further research is very likely to have an important impact on our confidence in the estimate of effect and is likely to change the estimate. Very low quality: We are very uncertain about the estimate.

DMPA: depot medroxyprogesterone acetate

Cu IUC: copper intrauterine contraception

ADDITIONAL SUMMARY OF FINDINGS [Explanation]

Outcomes by 1 year	Relative effect (95% CI)	Participants (study)	Quality of the evidence (GRADE)	Comments
Change in fat mass (%) ; change in lean mass (%)	Reported mean \pm SD (reported P): 2.5 \pm 8.0 vs -1.3 \pm 6.9 (P = 0.029); -1.4 \pm 4.7 vs 1.0 \pm 3.8 (P = 0.027)	76 (Dal'Ava 2012)	Moderate	LNG-IUC vs non-hor- monal IUC; women 18 to 45 years old
Change in fat mass (%) ; change in fat free mass (%)	MD 1.60 (0.45 to 2.75); MD -1.60 (-2.75 to -0. 45)	60 (Napolitano 2015)	Low	LNG-IUC vs no contra ceptive; perimenopausal wome
CI: Confidence interval;	MD = mean difference; SD	= standard deviation		

IUC: intrauterine contraception

LNG-IUC: levonorgestrel-releasing intrauterine contraception

Patient or population: perim Settings: clinic Intervention: OC containing Comparison: no hormonal c		• contraception		
Outcomes by 1 year	Relative effect (95% CI)	Participants (study)	Quality of the evidence (GRADE)	
Change in fat mass (%); change in fat free mass (%)	MD 3.30 (2.08 to 4.52); MD -3.30 (-4.52 to -2.08)	68 (Napolitano 2015)	Low	
CI: Confidence interval; MD = mean difference				
Moderate quality: Further reamay change the estimate.	h is very unlikely to change our co search is likely to have an importa is very likely to have an importan	int impact on our confide	nce in the estimate of effect and	

OC: oral contraceptive

CHARACTERISTICS OF STUDIES

Characteristics of included studies [ordered by study ID]

Ball 1991		
Methods	Design: randomized controlled trial Location: likely in Oxford, England Time frame: no information Sample size estimation and outcome of focu	us: no information
Participants	51 women, 17 to 41 years old, requesting or Inclusion criteria: new OC users had not us switchers were changing from low-dose con Exclusion criteria: hypertension (diastolic b BP > 140 mm Hg); smoking > 20 cigarettes	ed OC or hormone therapy for 3 months; mbined OC blood pressure (BP) > 100 mm Hg, systolic
Interventions	Progestin-only pills	
	1 Norethisterone (NE	T) 350 μg (N = 23)
	2 Levonorgestrel (LN	IG) $30 \ \mu g \ (N = 23)$
	6 treatment cycles	
Outcomes	Primary: mean change in weight; lipoprotei tors, and blood pressure Time frame: 6 months	n levels, glucose tolerance, coagulation fac-
Notes		
Risk of bias		
Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	No information other than stratified ac- cording to prior OC use
Allocation concealment (selection bias)	Unclear risk	No information
Blinding of participants and personnel (performance bias) All outcomes	Unclear risk	Single-blind (unspecified)
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	No information; objective outcome mea- sure
Incomplete outcome data (attrition bias) All outcomes	High risk	Loss to follow-up: 24% (12/51); analysis for weight included 39 women at 6 months (23 NET; 16 LNG) 5 did not return for follow-up and were ex- cluded (groups not specified); 9 withdrew after 3 months (1 NET; 8 LNG)
Bonny 2009		•
Methods	Design: prospective study; part of larger 2-y ception and bone mineral density Location: 4 urban adolescent health clinics OH (USA) Time frame: enrollment 2002 to 2003 Sample size estimation and outcome of focu	in large metropolitan area, likely Cleveland
Participants	alcohol or drug dependence; medical condit (e.g. corticosteroids) associated with outcor	ible for control group in past 6 months; OC use in past 3 months; tion (e.g. renal disease) or medication use

Interventions	Choice of study group		
	1 Depot medroxypro	gesterone acetate (DMPA) (N = 15): randomized to additional of placebo (N = 8) or estradiol cypionate 5 mg (E_2C) (N = 7)	
	2 Control (no hormo	nal contraception) (N = 18)	
		OCs (N = 18); not included here. Type of OC not specified; may estin-only OCs and combination OCs	
Outcomes	Percent change in total body fat and in lear Follow-up: 6 months	body mass	
Notes			
Risk of bias			
Bias	Authors' judgement	Support for judgement	
Random sequence generation (selection bias)	High risk	NRS: participants chose contraceptive method	
NOS selection (NRS)	Low risk	Exposed: clinic population; volunteers for study Non-exposed: same population as exposed but chose different contraceptive Exposure: adherence to DMPA appoint- ments	
NOS comparability (NRS)	Low risk	Analysis: multivariate models for change in total body fat and change in lean body mass; adjusted for potential confounders, e.g. age, race or ethnicity, caloric intake	
Blinding of participants and personnel (performance bias) All outcomes	Unclear risk	Not feasible due to women having chosen contraceptive method	
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	No information; objective outcome mea- sure	
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	Loss to follow-up: unknown; analysis table does not specify N To remain in study, participants had to ad- here to DMPA by appointment. No infor- mation on controls	
Castle 1978		•	
Methods	Design: prospective study Location: family planning center in Salisbu Time frame: enrolment June to December Sample size estimation and outcome of foc	1976	
Participants	Sample size estimation and outcome of focus: no information 1000 women; age range: "under 20" to 40 years or older Inclusion criteria: Black women seeking contraception at specific clinic Exclusion criteria: no mention		
Interventions	Ũ	ry 3 months (N = 500) ry 6 months (N = 500)	
Outcomes	Mean increase in weight Timeframe: 6 months Weight measured at each visit while partic	pant wore only a gown	
Notes			
Risk of bias			
Bias	Authors' judgement	Support for judgement	

Random sequence generation (selection bias)	High risk	NRS; allocation by participant request
NOS selection (NRS)	Low risk	Exposed: clinic population; volunteers for study Non-exposed: same population as exposed (different DMPA) Exposure: injections given during clinic ap- pointments
NOS comparability (NRS)	High risk	Analysis: no adjustment
Blinding of participants and personnel (performance bias) All outcomes	Unclear risk	Not feasible due to women having chosen contraceptive method
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	No information; objective outcome mea- sure
Incomplete outcome data (attrition bias) All outcomes	High risk	Loss to follow-up: 39% DMPA 150 mg and 23% DMPA 450 mg; differential losses between groups Withdrawals from study: 21 for DMPA 150 mg and 17 for DMPA 450 mg
Dal'Ava 2012		
Methods	Design: prospective study Location: single site in Campinas, Brazil Time frame: enrolled October 2009 to May 2010 Sample size estimation and outcome of focus: based on weight gain in LNG-IUC users from previous study; 37 needed for each group	
Participants	76 women Inclusion criteria: 18 to 45 years of age; initiated contraceptive use (either LNG-IUC or Cu T380A IUC) Exclusion criteria: currently breastfeeding or breastfeeding during 6 months before en- rollment; used corticosteroids, thiazide diuretics, or drugs for treatment of thyroid dis- ease; eating disorder; chronic disease	
Interventions	1 LNG-IUC (N = 38) 2 Cu T380A IUC (N = 38) 2	
Outcomes	Change in weight (kg); percent change in total fat mass, total lean mass, central-peripheral fat ratio Follow-up: 12 months	
Notes	Limitations: physical activity and daily caloric intake were not monitored during study	
Risk of bias	•	
Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	High risk	Women apparently chose contraceptive method
NOS selection (NRS)	Low risk	Exposed: study volunteers; recruitment methods not specified unclear if they were clinic attendees Non-exposed: same population as exposed but chose non-hormonal IUC Exposure: clinic inserted IUC
NOS comparability (NRS)	Low risk	Design: paired by age $(\pm 2 \text{ years})$ and BMI $(\pm 2 \text{ kg/m}^2)$ for intervention groups Analysis: no mention of using control vari- ables in analysis; listed sociodemographic variables; obstetric and gynaecologic his- tory; physical activity; consumption of cof- fee, alcohol, milk; smoking; family history

	1	of osteoporosis
Blinding of participants and personnel (performance bias) All outcomes	Unclear risk	Not feasible due to women having chosen contraceptive method
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	No information; objective outcome mea- sure
Incomplete outcome data (attrition bias) All outcomes	Low risk	Loss to follow-up: no discontinuation at 12 months
Dal'Ava 2014		
Methods	Design: prospective matched Location: single site in Campinas, Brazil Time frame: enrolled October 2009 to June 2011 Sample size estimation and outcome of focus: reported change in baseline percentage of body fat 12 months after initiation of contraception (primary outcome), 1.75% increase in DMPA users vs 0.31% reduction in Cu IUC users; required 20 complete cases in each group, using repeated measures ANOVA with 5% significance level and 80% power	
Participants	97 women Inclusion criteria: 18 to 50 years of age; new DMPA or Cu IUC users Exclusion criteria: breastfeeding; use of DMPA in prior 6 months; history of diabetes, pituitary disorder, liver or kidney disease, cancer; use of corticosteroids, diuretics, hor- mone therapy; eating disorder	
Interventions	1 DMPA 150 mg (N = 55)	
	2 Cu T380A IUC (N =	42)
Outcomes	Change in weight (kg), fat mass (kg), and lean mass (kg) Follow-up: 12 months	
Notes	Women using Cu IUC may overlap with those in Dal'Ava 2012; study periods overlap and inclusion criteria are similar. Unable to obtain further information from investigator	
Risk of bias		
Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	High risk	Participants chose method
NOS selection (NRS)	Low risk	Exposed: volunteered for study; unclear if they were clinic attendees Non-exposed: same population as exposed but chose different contraceptive method Exposure: clinic records (inserted IUC and administered DMPA)
NOS comparability (NRS)	Low risk	Design: paired by age (± 2 years) and weight (± 2 kg) for intervention groups Analysis: control variables of physical ac- tivity, consumption of coffee and alcohol, smoking in regression; reportedly not sig- nificant
Blinding of participants and personnel (performance bias) All outcomes	Unclear risk	Not feasible due to women having chosen contraceptive method

No mention; objective outcome measure

Loss to follow-up and discontinuation: DMPA 53% (29/55); IUC 24% (10/42) From IUC group, only 26 of 32 matched

Unclear risk

High risk

Blinding of outcome assessment (detection

Incomplete outcome data

(attrition bias) All outcomes

bias) All outcomes

D 0 . 2014		Major differential losses between groups	
Dos Santos 2014			
Methods	Design: prospective matched Location: Campinas, Brazil Time frame: enrolled February 2011 to February 2013 Sample size estimation and outcome of focus: none; exploratory study		
Participants	71 women Inclusion criteria: age 18 to 40 years; BMI < 30; never used DMPA; fasting glucose < 100 mg/dL and glucose level < 140 mg/dL at 120 minutes after 75 g oral glucose load Exclusion criteria: breastfeeding; family history of first-degree relative with diabetes mel- litus; type 1 or 2 diabetes, metabolic syndrome, hypertension, hyper- or hypothyroidism, chronic renal failure, hirsutismor hyperandrogenism, polycystic ovary syndrome or acan- thosis nigricans; transplant recipient; had undergone bariatric surgery		
Interventions	1 DMPA 150 mg intramuscular (IM) (N = 44)		
	2 Cu T380A IUC (N	Cu T380A IUC (N = 27)	
Outcomes	Change in body weight, total fat mass, percent body fat, and total lean mass; change in bone mineral density Follow-up: 12 months		
Notes			
Risk of bias			
Bias	Authors' judgement	Support for judgement	
Random sequence generation (selection bias)	High risk	Women chose method	
NOS selection (NRS)	Low risk	Exposed: clinic population, volunteered for study Non-exposed: same population as exposed but chose different contraceptive Exposure: clinic records; clinic inserted IUC and administered DMPA	
NOS comparability (NRS)	Low risk	Design: paired for age (± 1 year) and BMI (± 1 kg/m ²) Analysis: multiple linear regression in- cluded age, weight, BMI, schooling, skin color, social class, pregnancies, deliveries, smoking, alcohol, coffee (secondary report Modesto 2014)	
Blinding of participants and personnel (performance bias) All outcomes	Unclear risk	Not feasible due to women having chosen contraceptive method	
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	No mention; objective outcome measures	
Incomplete outcome data (attrition bias) All outcomes	High risk	Loss to follow-up and discontinuation: overall 24% (17/71); DMPA 34% (15/44) and CU IUC 7% (2/27) Major differential losses across groups	
Espey 2000	•	-	
Methods	Design: retrospective chart review Location: 3 Indian Health Service facilities in southwestern USA Time frame: first injection of DMPA from December 1992 to June 1995 Sample size estimation and outcome of focus: no information		
Participants	Female members of Navajo tribe Inclusion criteria: 18 to 40 years old; completed at least 5 consecutive injections of DMPA at 10- to 14-week intervals; had weights recorded at 1- or 2-year intervals Exclusion criteria: history of diabetes or thyroid disease; women in postpartum group		

	who had pre-eclampsia or multiple gestations within index pregnancy		
Interventions	DMPA initiation		
	1 Interval (N = 115): first injection 20 weeks past pregnancy of 20 weeks gestation		
	2 Postpartum (N = 57): first injection at 5 to 8 weeks after delivery of singleton pregnancy of 20 weeks gestation		
Outcomes	Mean weight gain (lb) for DMPA by initiation group Time frame: 1 and 2 years		
Notes	For another group, not included in this review, investigators reportedly extracted method of contraception from charts but did not provide specifics. Discussion noted that group "more frequently used" IUC or tubal ligation and included COC users		

Risk of bias	-		
Bias	Authors' judgement Support for judgement		
Random sequence generation (selection bias)	High risk	DMPA users by choice	
NOS selection (NRS)	Low risk	Exposed: clinic population of Navajo women Non-exposed: same population as exposed Exposure: clinic records of DMPA injec- tions	
NOS comparability (NRS)	Low risk	Analysis: adjusted for age, parity, and initial weight	
Blinding of participants and personnel (performance bias) All outcomes	Unclear risk	Not feasible due to differences in insertion times	
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	No information; objective outcome mea- sure	
Incomplete outcome data (attrition bias) All outcomes	High risk	Loss to follow-up: for 1 year, only com- plete records included in retrospective re- view; does not account for discontinuation and loss; at 2 years, no weight data for 70% of interval group and 49% of postpartum group	
Modesto 2015		·	
Methods		of focus: based on difference in weight between 5% and beta error 20% using repeated measures	
Participants	2138 women Inclusion criteria: 18 to 40 years of age; started using DMPA, Cu IUC or LNG-IUC in 1990 and used uninterruptedly for 10 years; Cu IUC and LNG-IUC users monitored annually at clinic and DMPA users attended clinic every 90 days for injection Exclusion criteria: chronic medical disease including dyslipidemia, diabetes, thyroid dis- ease, and renal failure; history of bariatric surgery or organ transplant; LNG-IUC use for other than contraception		
Interventions	1 DMPA 150 mg IM (N = 714)		
	2 Cu IUC (N = 723)		
	3 LNG-IUC (N	= 701)	
Outcomes	Mean weight gain (kg) Time frame: 1 year and 10 years		
Notes			

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	High risk	Women chose method; charts reviewed un- til 700 met eligibility criteria for each group
NOS selection (NRS)	Low risk	Exposed: clinic population, using contra- ceptive method for 10 years, beginning in 1990 Non-exposed: same population as exposed but chose different contraceptive Exposure: clinic records (clinic inserted IUC and administered DMPA)
NOS comparability (NRS)	Low risk	Analysis: generalized linear mixed model adjusted for years of school and number of children
Blinding of participants and personnel (performance bias) All outcomes	Unclear risk	Not feasible due to women having chosen contraceptive method
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	No mention; objective outcome measure
Incomplete outcome data (attrition bias) All outcomes	High risk	Loss to follow-up: retrospective review of charts with relevant data; does not account for discontinuation and loss Overall loss: at 1 year 12% (DMPA 5%, LNG-IUC 14%, Cu IUC 17%); at 4 years 19% (DMPA 24%, LNG-IUC 20%, Cu IUC 14%); at 10 years 84% (DMPA 82%, LNG-IUC 90%, Cu IUC 79%)
Moore 1995	-	•
Methods	Design: retrospective chart review Location: rural obstetrics and gynecology clinic in Arizona (USA) Time frame: no information Sample size estimation and outcome of focus: none; selected 50 women in each group who met inclusion criteria	
Participants	150 women, 15 to 30 years old Inclusion criteria: users of OCs, Norplant, or DMPA Exclusion criteria: prior hormonal contraceptive therapy; height < 62 inches (152.4 cm) or > 70 inches (177.8 cm); weight < 100 lb (45.5 kg) or > 180 lb (81.8 kg); presence of diabetes; history of thyroid disease; < 12 months postpartum	
Interventions	1 Norplant (N = 50)	
	2 DMPA 150 mg (N =	50)
	Third group of OC users excluded from this review; type of OC not specifie may have included progestin-only OCs and combination OCs	
Outcomes	Weight gain (kg) Timeframe: 1 year	
Notes		
Risk of bias		
Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	High risk	Women presumably chose contraceptive method; charts reviewed until 50 met eli- gibility criteria for each group
NOS selection (NRS)	Low risk	Exposed: rural clinic population Non-exposed: same population as exposed but used different contraceptive Exposure: clinic records

NOS comparability (NRS)	Low risk	Analysis: model adjusted for age, height, weight, and parity at beginning of study period	
Blinding of participants and personnel (performance bias) All outcomes	Unclear risk	Not feasible because women presumably chose contraceptive method	
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	No information; objective outcome mea- sure	
Incomplete outcome data (attrition bias) All outcomes	High risk	Loss to follow-up: retrospective review of charts with relevant data; does not account for discontinuation and loss	
Napolitano 2015			
Methods	Design: prospective study Location: menopause clinic in Italy Time frame: enrolled January 2011 to Janua Sample size estimation and outcome of focu needed per group to document between-grou 0.6; no power analysis reported	is: based on prior experience, 11 subjects	
Participants	110 women Inclusion criteria: perimenopausal based on serum FSH > 15 IU/ml and irregular men- strual cycles or amenorrhea < 3 months (aged 45 to 55 years) Exclusion criteria: current hormone use; sterilization; BMI < 18 or > 30; vascular disease or coagulation disorder; hypersensitivity to study drug ingredient; thyroid dysfunction; fasting glucose > 110 mg/dl; breast or gynecologic disease		
Interventions	1 Desogestrel (DSG) 75 μg OC (N = 44)		
	2 LNG-IUC (N = 35)		
	3 Control, no contrace	eption (N = 31)	
Outcomes	Change in weight (kg), BMI, fat mass (%), fat free mass (%), waist (cm), waist to hip ratio, resting metabolic rate (kJ/24 h) Follow-up: 12 months		
Notes			
Risk of bias			
Bias	Authors' judgement	Support for judgement	
Random sequence generation (selection bias)	High risk	Women chose method	
NOS selection (NRS)	Low risk	Exposed: menopause clinic population, volunteered for study Non-exposed: same population as exposed Exposure: clinic records (clinic inserted IUC)	
NOS comparability (NRS)	High risk	Analysis: unadjusted	
Blinding of participants and personnel (performance bias) All outcomes	Unclear risk	Not feasible due to women having chosen contraceptive method	
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	No mention; objective outcome measure	
Incomplete outcome data (attrition bias) All outcomes	Low risk	Loss to follow-up or discontinuation: 7% (8/110); DSG OC 4.5% (2/44), LNG- IUC 3% (1/35), and control 16% (5/31); control had 2 lost to follow-up (6%)	

Nyirati 2013			
Methods	Design: prospective study using convenience sample Location: large Midwestern city, possibly Columbus OH (USA) Time frame: 18-month period, specific dates not given Sample size estimation and outcome of focus: not reported; body composition changes at 1 year postpartum		
Participants	78 postpartum women Inclusion criteria: 18 years of age; elected to use DMPA or surgical sterilization Exclusion criteria: hormone replacement therapy, e.g. OC, thyroid hormone, steroid therapy; significant prenatal and postpartum medical illness including gestational or type II diabetes; BMI > 35		
Interventions	1DMPA 150 mg every 12 weeks, beginning at 6 weeks postpartum (N = 61)2Surgical sterilization and no other contraceptive (N = 17)		
Outcomes	Change in weight (lb), BMI, fat%, body dimensions (fat folds and circumferences) from 6 weeks to 1 year postpartum Follow-up: every 3 months for 1 year		
Notes			

Support for judgement

Risk of bias		
Bias	Authors' judgement	
Random sequence generation (selection bias)	High risk	
NOS selection (NRS)	Low risk	
NOS comparability (NRS)	High risk	
Blinding of participants and	Unclear risk	

Random sequence generation (selection bias)	High risk	Women chose method
NOS selection (NRS)	Low risk	Exposed: clinic population; postpartum women Non-exposed: same population as exposed but chose different contraceptive Exposure: clinic records (clinic adminis- tered DMPA)
NOS comparability (NRS)	High risk	Analysis: no adjustment for potential con- founding of outcome measures
Blinding of participants and personnel (performance bias) All outcomes	Unclear risk	Not feasible due to women having chosen contraceptive method
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	No mention; objective outcome measure
Incomplete outcome data (attrition bias) All outcomes	High risk	Loss to follow-up or discontinuation: DMPA 43%(26/61); sterilization 29%(5/ 17)
Pantoja 2010		
Methods	Design: retrospective chart review Location: university department of obstetrics and gynecology in Campina, Brazil	

5		
Methods	Design: retrospective chart review Location: university department of obstetrics and gynecology in Campina, Brazil Time frame: chart data from January 1991 to December 2000 Sample size estimation and outcome of focus: difference in weight between DMPA users and nonusers using analysis of variance for repeat measures; based on mean increase in fat mass in users, 150 DMPA users estimated; significance 5% and power 80%	
Participants	Women who accepted contraceptive method Inclusion criteria: chose DMPA and used continuously 3 years or who used Cu T380A for similar time period (mean age 29 years) Exclusion criteria: diabetes mellitus; hyperthyroidism or hypothyroidism; chronic renal failure; rheumatic diseases requiring chronic use of corticoids; organ transplant	
Interventions	1 DMPA (N = 379)	
	2 Cu T380A IUC (N = 379)	
	After pairing for age and baseline BMI, 379 for e	ach contraceptive group

Outcomes	Change in weight (kg) by contraceptive group and by baseline BMI (kg/m ²) (< 25; 25 to 29.9; 30) Time frame: 1, 2, and 3 years Weight and height measured at baseline (method initiation) and annually		
Notes			
Risk of bias			
Bias	Authors' judgement	Support for judgement	
Random sequence generation (selection bias)	High risk	Women chose contraceptive method	
NOS selection (NRS)	Low risk	Exposed: clinic attendees Non-exposed: same population as controls but chose different contraceptive Exposure: from clinic records	
NOS comparability (NRS)	Low risk	Design: paired for age and baseline BMI	
Blinding of participants and personnel (performance bias) All outcomes	Unclear risk	Not feasible due to women having chosen contraceptive method	
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	No information; objective outcome mea- sure	
Incomplete outcome data (attrition bias) All outcomes	High risk Loss to follow-up: chart review of those with 3 years continuous use; does not account for discontinuation and loss		
Salem 1984		•	
Methods	Design: prospective study; focused on effect of Norplant use on lactating women and on lactation performance and infant growth Location: postpartum clinic of university hospital in Assiut, Egypt Time frame: no information Sample size estimation and outcome of focus: no information; recruited 50 women for each group		
Participants	150 lactating women; mean age 29 years Inclusion criteria: normal delivery of normal living baby and exclusively breastfeeding; 1 month after delivery; infant weight 3500 gm Exclusion criteria: no mention		
Interventions	Acceptors (50 in each group)		
	1 Norplant	1 Norplant	
	2 Barrier, 'local' or no contraceptive method		
	3 Cu T380A IUC		
Outcomes	Weight gain by study group Follow-up: 6 months		
Notes			
Risk of bias	1		
Bias	Authors' judgement Support for judgement		
Random sequence generation (selection bias)	High risk	Women apparently chose contraceptive method	
NOS selection (NRS)	Low risk	Exposed: attendees at postpartum clinic; volunteered to participate Non-exposed: same population and time- frame as controls but chose different con- traceptive Exposure: obtained during monthly fol-	

	1	low-up visits	
NOS comparability (NRS)	High risk	Analysis: no adjustment	
Blinding of participants and personnel (performance bias) All outcomes	Unclear risk	Not feasible due to women having chosen contraceptive method	
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	No information; objective outcome mea- sure	
Incomplete outcome data (attrition bias) All outcomes	Low risk	Loss to follow-up: only those excluded (1 Norplant group, lost baby and wanted to get pregnant; 2 pregnancies in group with barrier, 'local', or no contraceptive)	
Salem 1988			
Methods	Design: randomized controlled trial; examine side effects, continuation, and termination Location: family planning center in Assiut, Eg Time frame: no information Sample size estimation and outcome of focus	gypt	
Participants	400 women attending family planning clinic Inclusion criteria: 18 to 40 years old; proven fertility and frequent risk of pregnancy; regular menstrual cycles; willing to rely on one method Exclusion criteria: breast-feeding; cardiovascular disease; liver disease; known or sus- pected breast malignancy, genital malignancy, uterine fibroids; undiagnosed vaginal bleeding; suspected pregnancy		
Interventions	200 in each group		
	1 DMPA 150 mg every 3 months		
	2 Norethisterone enanthate (NET-EN) 200 mg every 2 months		
Outcomes	Mean change in weight by contraceptive group; units not specified (kg or lb) Follow-up: 1 year Report had mean change for those with increase, decrease, or no change in weight We calculated combined weight change means and standard deviations		
Notes			
Risk of bias			
Bias	Authors' judgement	Support for judgement	
Random sequence generation (selection bias)	Low risk	Random numbers table prepared by WHO	
Allocation concealment (selection bias)	Low risk Sealed envelopes contained assignments		
Blinding of participants and personnel (performance bias) All outcomes	Unclear risk Not feasible due to different injection schedules		
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk No information; objective outcome mea- sure		
Incomplete outcome data (attrition bias) All outcomes	High risk	Loss to follow-up: reportedly 19% for DMPA and 13.3% for NET-EN One-year method continuation rates: 68. 8% DMPA and 57.1% NET-EN Finished study: 54% DMPA and 47% NET-EN	

Sivin 1998 Methods

Design: randomized controlled trial; focused on effectiveness of reformulated 2-rod LNG

	implant versus 6-rod implant Location: 7 centers including USA and Finland Time frame: enrollment 1990 to 1994 Sample size estimation and outcome of focus: assumed 50/100 acceptors would continue; pregnancy rate 2.0/100 at 3 years with SE 0.66/100; sample size could distinguish difference in pregnancy of 2/100 between 2 implant types		
Participants	1200 healthy women, 18 to 40 years old, who sought implant contraception Inclusion criteria: no contraindication to implant use; willing to undergo study proce- dures Exclusion criteria: cancer; severe cardiovascular problem; hyperlipidemia; diabetes melli- tus; mental illness; epilepsy; severe or frequent headaches; undiagnosed genital bleeding; hyperprolactinemia or bloody breast discharge; pelvic inflammatory disease since last pregnancy or ectopic pregnancy		
Interventions	Levonorgestrel implants		
	1 Norplant: 6 capsules	containing levonorgestrel 216 mg (total)	
	2 LNG rod (Jadelle): 2 elastomer in core that	rods containing levonorgestrel 150 mg (total); different in earlier implant	
	Follow-up: 1, 3, 6 m	onths; then semi-annually to 5 years	
Outcomes	Mean weight change by implant group Weight change for 10th, 50th, and 90th percentiles of body weight (at admission) Follow-up: 5 years Weighing method not specified		
Notes			
Risk of bias			
Bias	Authors' judgement	Support for judgement	
Random sequence generation (selection bias)	Low risk	Randomization by "linear congruential method"; blocks of 50 per clinic	
Allocation concealment (selection bias)	Low risk	Implants in sealed opaque envelopes num- bered sequentially according to randomiza- tion lists	
Blinding of participants and personnel (performance bias) All outcomes	Unclear risk	Not feasible due to apparent differences in interventions	
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	No information; objective outcome mea- sures	
Incomplete outcome data (attrition bias) All outcomes	High risk	Loss to follow-up: year 3, 2.7% each group; year 5, 7.2% LNG rod and 10.2% Nor- plant Discontinuation: end of year 3, LNG rod 31.6% (190/600) and Norplant 31.2% (187/598); end of year 5, LNG rod 54.7% (328/600) and Norplant 60% (359/598) 2 sets of Norplant contaminated and not used (1198/2000 analyzed)	
Sule 2005			
Methods	Design: retrospective chart review Study examined hormonal contraceptives and weight changes. Location: family planning clinic of university hospital in Zaria, Nigeria Time frame: registered from 01 January 1993 to 31 December 1995 Sample size estimation and outcome of focus: no information		
Participants	516 new clients Inclusion criteria: used hormonal contraceptive (COC, injectable (DMPA or NET-EN), Norplant); non-hormonal IUC users as controls; followed for 1 year (mean age hormonal users 30.5 years and non-hormonal IUD 29.1 years) Exclusion criteria: used barrier methods; had bilateral tubal ligation; chose no contra- ceptive method		

Interventions	Method users		
	1	Norplant (N = 188)	
	2	non-hormonal IUC (N	= 136)
Outcomes	Mean weight gain or loss by contraceptive group Time frame: 1 and 3 years Report had mean change for those with increase, decrease, or no change in weight. We calculated combined weight change means and standard deviations		
Notes	Injectable users not u analysis	sed in this review; DMPA	A and NET-EN had been grouped for
Risk of bias	·		
Bias	Authors' judgement	t	Support for judgement
Random sequence generation (selection bias)	High risk		Users of contraceptive method; women presumably chose method
NOS selection (NRS)	Low risk		Exposed: clinic attendees Non-exposed: same population as exposed but chose different contraceptive Exposure: clinic records
NOS comparability (NRS)	High risk		Analysis: no adjustment
Blinding of participants and personnel (performance bias) All outcomes	Unclear risk		Not feasible due to women presumably having chosen method
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk		No information; objective outcome mea- sure
Incomplete outcome data (attrition bias) All outcomes	High risk		Loss to follow-up: investigators selected charts with 1 year of data, so no loss by 1 year; by 3 years, overall loss by 3 years of 54% (Norplant 31%; COC 95%; IUC 56%)
Taneepanichskul 1998			
Methods	IUC Location: family plan Time frame: no infor	ning clinic at a hospital i	-
Participants	100 women, age 37 to 50 years, attending family planning clinic Inclusion criteria: used DMPA or IUC for 120 months (10 years); followed "regularly"; no history of smoking or alcohol intake IUC users had not used any hormonal contraceptive Exclusion criteria: developed chronic disease or metabolic disorder during DMPA or IUC use		
Interventions	Method chosen		
	1	DMPA (N = 50)	
	2	Cu T380A IUC (N = 5	50)
Outcomes	Mean change in body weight Time frame: 120 months Weight measured in standard manner at 120 months; prior method not specified		
Notes			
Risk of bias			
Bias	Authors' judgement	t	Support for judgement

Random sequence generation (selection bias)	High risk	Users of DMPA "recruited randomly"; users of Cu T380A selected as controls	
NOS selection (NRS)	High risk	Exposed: clinic population, older contra- ceptive users; used method for 10 years and regularly attended clinic Non-exposed: same as exposed group but chose different contraceptive Exposure: presumably from clinic records	
NOS comparability (NRS)	Low risk	Design: matched for age, parity, income, weight	
Blinding of participants and personnel (performance bias) All outcomes	Unclear risk	Not feasible due to women presumably having chosen method	
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	No information; objective outcome mea- sure	
Incomplete outcome data (attrition bias) All outcomes		Loss to follow-up: retrospective study of women who used method for 10 years and regularly attended clinic; may have re- cruited women with relevant data in charts Exclusions: developed chronic disease or disorder during method use; may have bi- ased results because weight gain is asso- ciated with development of some diseases and disorders	
Tankeyoon 1976		-	
Methods	Design: prospective metabolic study; focused on metabolic effects of contraceptive meth- ods Location: Bangkok, Thailand Time frame: no information Sample size estimation and outcome of focus: no sample size calculation; focused on metabolic effects of contraceptive methods		
Participants	32 healthy women attending the family planning clinic; age 18 to 38 years Inclusion criteria: > 6 weeks postpartum and no other steroid use for past 3 months Exclusion criteria: no information		
Interventions	1 DMPA 150 mg (3-m	onth intervals) (N = 16)	
	2 COC: d-norgestrel 5	$0 \ \mu g + EE \ 50 \ \mu g \ (N = 16)$	
Outcomes	Percent of cases with 1 kg increase or decr Follow-up: 1, 2, 3, 6, 9, 12 months	ease in body weight by contraceptive method	
Notes			
Risk of bias			
Bias	Authors' judgement	Support for judgement	
Random sequence generation (selection bias)	High risk	Women receiving contraceptive method; presumably chose method	
NOS selection (NRS)	Low risk	Exposed: clinic population, volunteered for study Non-exposed: same as exposed group but chose different contraceptive Exposure: presumably from clinic records	
NOS comparability (NRS)	Low risk	Analysis: adjusted for pretreatment value	
Blinding of participants and personnel (performance bias) All outcomes	Unclear risk	Not feasible due to women presumably having chosen method	

Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	No information; objective outcome mea- sure	
Incomplete outcome data (attrition bias) All outcomes	High risk	Loss to follow-up: 19% by 12 months; DMPA 2/16 (13%) and COC 4/16 (25%) Differential losses between groups though sample sizes are small Reasons for missing data not specified	
Tuchman 2005	•	·	
Methods	Design: retrospective chart review Location: urban, hospital-based, teen health center (USA) Time frame: enrollment 01 January 2001 to 31 December 2001 Sample size estimation and outcome of focus: no sample size calculation; emphasis on weight change		
Participants	222 females, aged 12 to 21 years, attending health center for contraception Inclusion criteria: first-time use of oral or injectable contraceptive 'New start' defined as no OC in past 3 months or DMPA in past 6 months prior to new method initiation Exclusion criteria: no other information		
Interventions	Choice of method		
	1 DMPA every 3 months		
	2 Medroxyprogesteror	he acetate + estradiol cypionate 5 mg (MPA + E_2C) monthly	
	3 COC		
Outcomes	Mean weight change (kg) and mean percent Time frame: 3, 6, 9, 12 months Standardized weight and height measures de		
Notes			
Risk of bias			
Bias	Authors' judgement	Support for judgement	
Random sequence generation (selection bias)	High risk	Young women presumably chose contra- ceptive method	
NOS selection (NRS)	Low risk	Exposed: clinic population of new users Non-exposed: same as exposed group but chose different contraceptive Exposure: clinic records	
NOS comparability (NRS)	High risk	Analysis: no adjustment for weight out- come	
Blinding of participants and personnel (performance bias) All outcomes	Unclear risk	Not feasible due to differences in inter- ventions and women presumably chose method	
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	No information; objective outcome mea- sure	
Incomplete outcome data (attrition bias) All outcomes	High risk	Loss to follow-up or discontinuation: may have selected charts with relevant data for retrospective review; at 12 months, discon- tinuation 54% to 58%	

Vickery 2013

Methods

Design: substudy of CHOICE (see Notes below)

Location: St. Louis, MO (USA)

Time frame: enrolled between June 2009 and May 2011 Sample size estimation and outcome of focus: for weight change (kg) at 12 months, assumed mean weight gain 0.6 kg over 12 months in Cu IUC users and 2.0 kg in

	progestin-only users; assumed alpha 0.05, power 80%, and SD 3.0 kg in all groups; needed 73 women in each arm; for SD of 5.0 kg for Cu IUC and 6.0 kg for progestin- only users, increased sample size to 100 in Cu IUC group and 130 in progestin-only groups		
Participants	427 women enrolled in CHOICE Inclusion criteria: 18 to 45 years old; continuous user for 11 months of LNG-IUC, Cu IUC, implant, or DMPA; enrolled at university research site; had height and weight measured at enrollment visit Exclusion criteria: did not speak English; < 18 years old; metabolic disorder known to affect body weight, e.g. hypothyroidism or diabetes		
Interventions	1	DMPA (N = 67)	
	2	LNG-IUC (N = 130)	
	3	Cu IUC (N = 100)	
	4	ENG implant (N = 130)	
Outcomes	Weight change Follow-up: 12		
Notes		CHOICE, prospective cohort study of 9256 women: promote the use of LARC, remove financial barriers by providing contraceptives at no cost, and evaluate method continu-	

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	High risk	Women selected method of contraception
NOS selection (NRS)	Low risk	Exposed: clinic population; volunteered to participate in substudy Non-exposed: same population as exposed but chose different contraceptive Exposure: clinic records (inserted IUC or ENG implant and administered DMPA)
NOS comparability (NRS)	Low risk	Analysis: stratified by race (associated with weight change); final adjusted linear regres- sion model included age (LNG-IUC and Cu-IUC users were slightly older) as well as covariates associated with outcome and ex- posure and those that altered effect 10%
Blinding of participants and personnel (performance bias) All outcomes	Unclear risk	Not feasible due to women having chosen contraceptive method
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	No mention; objective outcome measure
Incomplete outcome data (attrition bias) All outcomes		Loss to follow-up: selected women who completed 11 months of use; 35% of women screened met eligibility criteria (749/2145), of which 57% enrolled (427/ 749)

Westhoff 2007	
Methods	Design: analysis of data from 3 RCTs Location: sites in North and South America Time frame: no information Sample size estimation and outcome of focus: no information
Participants	534 women, 18 to 49 years old, sexually active and wanting long-term contraception Inclusion criteria: no OC use for past 2 months; regular menstruation in past 3 months; willing to rely on DMPA for year Exclusion criteria: used OCs, implants, or hormonal IUC in past 2 months or DMPA-IM in past 10 months; pregnant or infertile; abnormal Pap; undiagnosed genital bleeding;

Interventions	1	DMPA-SC 104 mg (1	N = 266)
	2	DMPA-IM 150 mg (1	
	_	Injections every 3 mo	
Outcomes	Weight change (as safety endpoint) Follow-up: 36 months		
Notes	Investigators also analyzed weight change by BMI group: 25; 25 to 30 ; > 30 kg/m ² . Report notes no consistent differences by BMI groups		
Risk of bias			
Bias	Authors' judgen	nent	Support for judgement
Random sequence generation (selection bias)	Unclear risk		No information
Allocation concealment (selection bias)	Unclear risk		No information
Blinding of participants and personnel (performance bias) All outcomes	Unclear risk		Not feasible due to differences in interven- tions (intramuscular vs subcutaneous)
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk		No information for 2 trials; evaluator blinded in 1 trial, but unclear if relevant to weight outcome
Incomplete outcome data (attrition bias) All outcomes	High risk Loss to follow-up or discontinuation: DMPA-SC 24% (201/266); DMPA-IM 79% (212/268); reasons for discontinua- tion unclear		DMPA-SC 24% (201/266); DMPA-IM 79% (212/268); reasons for discontinua-
WHO 1983			
Methods	Location: 13 cent Time frame: recru	randomized controlled trial ters in Africa, Asia, Central uitment began 1977; final fo nation and outcome of focus	and South America, and Europe llow-up March 1982 : no information
Participants	3172 women; mean age 27.4 years Inclusion criteria: non-breastfeeding women who chose injectable contraception Exclusion criteria: contraindication for long-acting contraceptive methods		
Interventions	1 DMPA 150 mg at 90-day intervals (N = 1587)		
	2	Norethisterone enant	nate (NET-EN) 200 mg at 60-day intervals (N = 789)
	3	NET-EN 200 mg at 6	0-day intervals for 6 months then 84-day intervals ($N = 796$)
Outcomes	Mean weight change by contraceptive group Follow-up: 12 and 24 months		
Notes	Method for measure	uring weight not mentioned	
Risk of bias			
Bias	Authors' judgement Support for judgement		Support for judgement
Random sequence generation (selection bias)	Unclear risk		"Randomly allocated"
Allocation concealment (selection bias)	Unclear risk		No information
Blinding of participants and personnel (performance bias) All outcomes	Unclear risk Not feasible due to differences in injection schedules for interventions		

Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	No information
Incomplete outcome data (attrition bias) All outcomes	High risk	Loss to follow-up: reportedly 10.7% DMPA; 8.9% NET-EN 60 days; 9.8% NET-EN 84 days Life-table rates for total discontinuation: 71% to 74%

BMI: body mass index

DMPA: depot medroxyprogesterone acetate

FSH: follicle stimulating hormone

LNG-IUC: levonorgestrel-releasing intrauterine contraception

MPA: medroxyprogesterone acetate

NET-EN: norethisterone enanthate

NRS: non-randomized study

OC: oral contraceptive

SD: standard deviation

Characteristics of excluded studies [ordered by study ID]

Study	Reason for exclusion		
Agoestina 1978	Insufficient weight change data: presented in figure without any specific numbers, other than mean gain for DMPA group in text		
Bahamondes 2010	Mean weight change not reported. BMI included as control variable for examining bone mineral density Study examined weight change among participants who had been using the method (LNG-IUC) for at least 7 years before the study		
Barsivala 1974	Insufficient data: study duration not reported (our criteria was 3 months); also, investigators did not specify whether the variance reported is standard deviation or standard error		
Beksinska 2010	Analysis combined users of DMPA, NET-EN, or both. Investigators noted the subgroups were too small to analyze separately and that differences in weight gain were not significant		
Berenson 1997	Insufficient weight change data: means reported without any variance measure		
Berenson 2009	Insufficient weight change data: mean change reported without any variance. Of 240 who chose DMPA, 60% discontinued the method by 12 months and 76% discontinued by 36 months. DMPA users with > 5% weight increase at 6-month visit were more likely lost to follow-up by next visit than those who had not gained such weight Secondary report (Rahman 2012) provided weight gain (by kg categories) over any 6-month period; categories overlapped		
Bonny 2006	Analysis combined groups that received DMPA + placebo or DMPA + estradiol supplement; reportedly DMPA groups did not differ in weight gain DMPA discontinuation 37% at 18 months; data from 2000 to 2003		
Bonny 2011	Not comparative		
Bonny 2015	Not comparative		
Casey 2013	Insufficient data: no weight change reported		
Chen 2011	Abstract notes that weight gain was reported as side effect. Wrote to investigator regarding whether weigh change was measured. Unable to obtain further information		
Clark 2005	Insufficient weight change data: means (not mean change) presented in a figure. Text mentions mean change for DMPA at 30 months (no variance measure) and that the control group was basically unchanged Due to discontinuations of DMPA and initiation of hormonal contraception among controls, the samples sizes were 17% (DMPA) and 19% (controls) of baseline by the last visit		
Costa 2012	Mean change in weight was not reported. Participants were 1.5 months postpartum		
Dahlberg 1982	Insufficient weight change data: means reported without any variance measure		
El Mahgoub 1980	Insufficient weight change data: mean change reported without any variance measure. Also, percent that lost or gained weight was reported, but no specific amount of weight was provided		
Gallo 2016	Two arms: immediate and delayed insertion of Sino-Implant II. Secondary analysis of data from RCT tha examined condom use. Delayed group was offered OC prescription; no information on how many used it		
Hall 1980	Insufficient weight change data: mean change in 'ideal body weight' shown in figure, except for mean change for progestin-only group reported in text		
Havranek 1972	Insufficient weight data: mean change for one group reported without any variance measure		
Hernandez-Juarez 2014	Insufficient weight data; weight change not reported		
Kaunitz 2009	Insufficient weight data. Mean change was reported within adverse event section without any variance measure. Data from 2001 to 2004		
Mangan 2002	Comparison groups were DMPA users and OC users. Types of OC were not specified and might have included progestin-only as well as combination OC		
Modesto 2014b	Insufficient data: counseling only; no weight change measured		
Nault 2013	Insufficient data: weight gain not broken down by contraceptive type		
Olsson 1988	Insufficient data for analysis: no N per group for analysis. First-year continuation rate was 59% for Norpla and 77% for Norplant-2; methods suggest these were life-table rates		
Ortayli 2001	Insufficient data: report does not provide sample sizes used for analysis. Outcome data are from a pilot conducted in 1995 and the main study conducted from 1996 to 1998		

Study	Reason for exclusion	
Risser 1999	Comparison groups were DMPA users and OC users. Types of OC were not specified and might have included progestin-only as well as combination OC	
Segall-Gutierrez 2012	Single-arm study comparing normal weight and obese women. The one intervention was subcutaneous DMPA	
Veisi 2013	Insufficient weight data: self-reported weight only	
WHO 1978	Insufficient weight data: mean gains reported without any variance measure	
Yela 2006	Insufficient weight change data: tables show weight and BMI means (not change) by year; text mentions mean change per group over 5 years without any variance measure. Study began in 1998	
Zheng 1999	Insufficient weight change data: means reported without any variance measure	

BMI: body mass index

DMPA: depot medroxyprogesterone acetate

LNG-IUC: levonorgestrel-releasing intrauterine contraception

NET-EN: norethisterone enanthate

OC: oral contraceptive

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Characteristics of studies awaiting assessment [ordered by study ID]

Madden 2014

Madden 2014	_		
Methods	Purpose: learn in Location: St Lor Time frame: Ap	tional prospective cohort; non-probability sample f women gain weight using progestin-only methods of contraception and if so, how much uis, MO (USA) vril 2010 to August 2014 imation and outcome of focus: no information	
Participants	345 women Inclusion criteria: aged 18 to 45 years; starting copper IUC or implant through provider; first study visit must occur within 14 days of method insertion Exclusion criteria: DMPA in past 16 weeks; POPs, LNG-IUC, or implant in past 4 weeks; thyroid disease, autoim- mune disease, diabetes (excluding gestational); history of eating disorder; currently taking antidepressants for < 6 months, antipsychotics; oral glucocorticoids (steroids, i.e. prednisone) for > 6 months; currently breastfeeding or < 6 months postpartum		
Interventions	1	Levonorgestrel-containing intrauterine contraceptive (LNG-IUC)	
	2	Etonogestrel (ENG) subdermal implant	
	3	Copper IUC	
Outcomes	Primary: weight change; BMI change Secondary: body composition, including fat mass and percentage and central-to-peripheral fat ratio Body composition with dual-energy x-ray absorptiometry (DEXA); diet and activity via validated questionnaires Follow-up: 12 months		
Notes	Will consider for inclusion when full report is available		
Schreiber 2014			
Methods	Location: Philac Time frame: Ap	ized; open label; mixed methods delphia PA (USA) rril 2011 to October 2012 imation and outcome of focus: no information	
Participants	100 women Inclusion criteria: English-speaking; age 18 to 45 years; immediately postpartum of live singleton infant, 37 weeks gestation; desire to delay another pregnancy for next 6 months; willing and able to follow protocol Exclusion criteria: breastfeeding; plans to relocate outside of Philadelphia area in next 6 months; plans for use of weight loss medication or diet pills in next 6 months; wish to use Implanon or DMPA prior to discharge but does not want to be randomized		
Interventions	1	DepoProvera (DMPA) 150 mg	
	2	Implanon; 68 mg etonogestrel	
	3	Control: select own method of contraception or no contraception	
Outcomes	Primary: weight change by 6 months postpartum Secondary: pregnancy; contraception satisfaction Follow-up: 3, 6, and 12 months postpartum		
Notes	XX711 1 1 C	or inclusion when full report is available	

BMI: body mass index

DMPA: depot medroxyprogesterone acetate

LNG-IUC: levonorgestrel-releasing intrauterine contraception

POP: progestin-only oral pills

Characteristics of ongoing studies [ordered by study ID]

Bonny 2016

Trial name or title	Drug Exposure and Depot Medroxyprogesterone Acetate (DMPA) in Adolescent Subjects		
Methods	Design: randomized, open-label; pharmacokinetic study Purpose: learn whether DMPA affects weight gain and bone mineral density in teens Location: Columbus, OH (USA) Time frame: September 2011 to April 2015 Sample size estimation and outcome of focus: no information		
Participants	45 healthy young women Inclusion criteria: healthy, postmenarchal females; age 12 to 21 years; self-selected to initiate DMPA; willing to use barrier method of contraception in addition to DMPA Exclusion criteria: chronic disease known to affect weight or bone mineral density (BMD) (e.g. diabetes, kidney); medication known to affect weight or BMD (e.g. corticosteroids); DMPA use in past 12 months; pregnancy in past 6 months; etonogestrel implant, LNG-IUC or combined contraceptive in past 3 months (OC, transdermal patch, vaginal ring); weight > 450 lb; need for confidential contraceptive care for individuals < 18 years of age		
Interventions	DMPA, intramuscular injection		
	• 150 mg; approved for use (US Food and Drug Administration (FDA)) when given into muscle		
	• 104 mg; FDA-approved for use only when given under the skin; considered experimental		
	• 75 mg; considered experimental; not FDA-approved regardless of how given		
Outcomes	Primary: > 5% weight gain at 24 weeks Secondary: > 10% weight gain at 48 weeks Follow-up: 24 and 48 weeks		
Starting date	September 2011; planned completion April 2016		
Contact information	Andrea Bonny, MD; Nationwide Children's Hospital; Columbus, Ohio (USA)		
Notes			
Halpern 2017			
Trial name or title	Pharmacodynamics and Pharmacokinetics Study of Existing DMPA Contraceptive Methods		
Methods	Design: randomized, multicenter, open label Location: Portland OR (USA); Santo Domingo, Dominican Republic Time frame: September 2015 to August 2017 Sample size estimation and outcome of focus: no information		
Participants	48 women Inclusion criteria: in good general health; aged 18 to 40 years; willing to provide consent and follow study requirements; negative urine pregnancy test on day of injection; does not become pregnant in next 24 months; regular menstrual cycle (27 to 35 days); confirmed ovulation in 2 consecutive samples during pre-treatment phase; low risk of pregnancy; BMI 18 to 35; hemoglobin 10.5 g/L Exclusion criteria: medical contraindication to DMPA use; use of any investigational drug, prohibited drugs, OCs, LNG-IUC or implant within 1 month prior to enrollment; use of DMPA in past 12 months; use of combined injectable contraceptive in past 6 months; recent pregnancy (within 3 months); current lactation; ongoing or anticipated use of prohibited drugs; known sensitivity to MPA; plan to move to another location in next 18 months		
Interventions	1 1 subcutaneous injection of 150 mg/mL DMPA in abdomen		
	2 1 subcutaneous injection of 300 mg/mL DMPA in abdomen		
	3 2 injections of 104 mg/0.65 mL DMPA in abdomen, given at 3-month intervals		
Outcomes	Primary: time to ovulation Secondary: weight at follow-up; Cmax; aggregate of individual Cmax measurements and parameters; adverse events Follow-up: 18 months		
Starting date	September 2015; planned completion August 2017		
Contact information	Vera Halpern, MD: FHI 360; vhalpernfhi360.org; 919-544-7040 x11390		
Notes			

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DMPA: depot medroxyprogesterone acetate LNG-IUC: levonorgestrel-releasing intrauterine contraception MPA: medroxyprogesterone acetate

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