BMJ Open Systematic review of copper intrauterine contraception continuation in young nulliparous women based on intrauterine device type

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ABSTRACT

Objectives No copper intrauterine device (IUD) type is known to better suit young nulliparous women who tend to experience higher rates of IUD discontinuation compared with their older parous counterparts. A systematic review to determine which IUDs have higher continuation rates in young nulliparous women was undertaken.

Design Systematic review and meta-analyses of available evidence based on IUD type.

Data sources AMED, BNI, CINAHL, DARE, EMBASE, EMCARE, HMIC, MEDLINE, PsycINFO, PubMed, TRIP, and the Cochrane Library electronic databases were searched from inception to 11 May 2022; as well as the Bandolier, Medicines and Healthcare products Regulatory Agency, Faculty of Sexual and Reproductive Healthcare, Royal College of Obstetricians and Gynaecologists, Department of Health, National Institute for Health and Care Excellence, Scottish Intercollegiate Guidelines, WHO and Google Scholar websites.

Eligibility criteria All studies on IUDs currently available in the UK or comparable (same design and size) to those available in the UK, involving nulliparous women of any age including those aged under 30.

Data extraction and synthesis Independently extracted data were assessed as low risk of bias using the Mixed Methods Appraisal Tool. Random effects meta-analyses of proportions were performed where data, including subgroups, were amenable to quantitative synthesis. Heterogeneity was reported using tau² and I² statistics, and sensitivity analyses were also performed.

Results Nineteen studies involving 13045 nulliparous women were included but the heterogeneity of participant ages, parity and IUD types made quantitative synthesis of outcome data in totality inappropriate. The highest continuation rate obtained was 91.02% (95% CI 88.01% to 93.64%) for the smaller TCu 380A at 12 months post insertion.

Conclusions Evidence for IUD use in young nulliparous women based on IUD type remains limited. Smaller sized IUD types appear better suited to this group of IUD users, however, more research is needed.

PROSPERO registration number CRD42019120969.

INTRODUCTION

The highest rates of unintended pregnancy terminations of pregnancy, which

STRENGTHS AND LIMITATIONS OF THE STUDY

- ⇒ The first reported systematic review exploring intrauterine device (IUD) types in young nulliparous women.
- ⇒ A wide range of data sources, unrestricted to randomised controlled trials, was reviewed —an approach more representative of the real world.
- ⇒ Articles for inclusion were limited to publications in the English language.
- ⇒ Some data were obtained by calculation and measurements of graphs or figures where these data were not numerically specified in reports.
- ⇒ Most studies did not differentiate between nulligravid and nulliparous participants.

contribute to poor sexual health, are in women aged 20-24 followed by those aged 25–29. Increasing uptake of long-acting reversible contraceptives (LARCs), such as copper intrauterine contraception, in these women is yet to yield a proportional reduction in pregnancy terminations. This is attributable to their higher LARC discontinuation

Copper intrauterine contraception is the LARC with the greatest number of brands, with 21 copper intrauterine devices (IUDs) available in the UK.3 IUDs are of various shapes, sizes, total copper surface area and copper distribution on the IUD frame. They have changed little over the last 40 years. No IUD type has been shown to be associated with better outcomes regarding unwanted effects that lead to early IUD discontinuation. This early IUD discontinuation excludes discontinuation due to IUD user choice alone or the wish to conceive. IUD continuation rates tend to be surrogate for IUD satisfaction and/or acceptability. Studies have shown IUD discontinuation rates to be higher in adolescents and women in their 20s compared with their



older counterparts, as well as in nulliparous compared with parous women. $^{4-8}$

Previous systematic reviews and guidance suggest that IUD size and shape may be a factor in discontinuation, and have recommended future research investigate which IUD types are associated with less pain, bleeding and discontinuation.^{7 9-11} The identification and use of IUDs with higher continuation rates and fewer unwanted effects could improve outcomes including IUD satisfaction for young nulliparous women. A systematic review and meta-analysis were therefore undertaken to investigate continuation rates and reasons for discontinuation of IUDs, currently available, or comparable to those currently in use in the UK, based on IUD type involving women aged under 30.

OBJECTIVES

This study aimed to determine which currently available IUDs have higher continuation rates, in nulliparous women aged under 30, by systematically reviewing published studies. Discontinuation rates and reasons for discontinuation were secondary outcomes.

METHODS

An appraisal of previous systematic reviews, including publications by the Cochrane Collaboration Fertility Regulation Group, Faculty of Sexual and Reproductive Healthcare (FSRH) and National Institute for Health and Care Excellence (NICE), was performed. A search strategy was developed in conjunction with an Electronic Services Librarian. These informed the design of this systematic review and its protocol.

This study is reported as per the Preferred Reporting Items for Systematic Reviews and Meta-Analyses (PRISMA) guideline (see online supplemental material 1). Its protocol was registered on the International Prospective Register of Systematic Reviews database (see online supplemental material 2). The protocol included other studies besides randomised controlled trials (RCTs) reporting on IUD continuation, in case the RCTs determined eligible for inclusion in the systematic review were too few to address the review question.

Selection criteria

Inclusion criteria

Inclusion criteria are as follows: articles published in English, on studies in women who are nulliparous and aged under 30, that involved IUDs available or of the same design and size, to those available in the UK.

Exclusion criteria

Exclusion criteria are as follows: articles not published in English, studies solely in parous women aged 30 or over 30, that involved IUDs not available, or not of the same design and size to those available in the UK.

Where studies on IUDs currently available in the UK were lacking, studies with IUDs comparable in shape, size, total copper surface area or distribution on the IUD frame to those currently available in the UK were included. Where studies involving only nulliparous women aged under 30 were lacking, studies with nulliparous women of all ages (incorporating those aged under 30) were also included in the review.

Search strategy

Nine electronic databases—the Allied and Complementary Medicine (AMED), British Nursing Index (BNI), Cumulative Index to Nursing and Allied Health Literature (CINAHL), Excerpta Medica Database (EMBASE), Nursing and Allied Health Professionals Database (EMCARE), Health Management Information Consortium (HMIC), General Medical Database (MEDLINE), Psychology and Allied Fields (PsycINFO) and PubMed were searched. The search terms were (copper intrauterine).ti,ab OR (copper intrauterine device).ti,ab OR (copper coil).ti,ab OR (copper IUD).ti,ab OR (copper T).ti,ab from database inception to 7 February 2021 (updated to 11 May 2022). The following additional sources were searched using the term 'copper intrauterine': the Cochrane Library, Database of Abstracts and Reviews of Effects (DARE), Turning Research into Practice (TRIP) database, National Electronic Library of Health (merged with MEDLINE), Bandolier, Medicines and Healthcare products Regulatory Agency, FSRH, Royal College of Obstetricians and Gynaecologists, Department of Health, NICE, Scottish Intercollegiate Guidelines and WHO websites. A Google Scholar search was also undertaken using the term 'copper intrauterine device young nulliparous'. The full search strategy is provided as a supplementary file (online supplemental material 3).

Relevant articles published in English were identified by two authors and these were exported into an Endnote library on completion of all the searches. Following deduplication, the relevant articles obtained from the searches were exported to Rayyan, a web app for systematic reviews (rayyan.ai). In Rayyan, further deduplication yielded unique entries of which abstracts, and then full texts, were screened independently by two authors to assess eligibility for inclusion in the systematic review based on the inclusion/exclusion criteria. Additional citation screening of reference lists of both included and excluded studies was performed. Screening was initially done in batches of 20, then later increased to 50. Agreements were obtained between the first two authors and did not require a third review. Selected articles were RCTs and observational studies published in English, involving IUDs available or comparable to those in the UK, and involving nulliparous women aged under 30.

Quality assessment and data summary

All articles selected for inclusion in the systematic review underwent a quality assessment using the Mixed Methods Appraisal Tool (MMAT), v.2018. The MMAT risk of bias

tool was chosen because it was applicable to all the study types selected for inclusion. The highest total MMAT score conforming with best quality was seven, while the lowest possible score equating with poorest quality was zero. Included articles were initially quality assessed by the two authors separately and then agreement was reached.

Data extracted from articles included IUD type, study location(s) and year of publication, age of women, gravidity/parity of women, IUD continuation and discontinuation rates and reasons for IUD discontinuation. Where a rate was not specified but could be reliably calculated, this was done to one decimal place. If a continuation rate was not specified, this was obtained by subtracting the discontinuation rate from 100, or adding all stated rates for reasons for discontinuation (where these were mutually exclusive) and subtracting from 100, if the report suggested such a calculation to be valid. If a discontinuation rate was not specified, this was obtained by subtracting a stated continuation rate from 100, or by adding all stated rates for reasons for discontinuation (where these were mutually exclusive), if the report suggested such a calculation was valid. Gross rates (obtained after excluding participants lost to follow-up or removals to conceive) were used, except where only net cumulative rates were reported. Measurements were performed to obtain data from published graphs or figures where rates had been reported in this format but not numerically specified.

An Excel data collection form was developed, piloted with three articles selected for inclusion by one author, then revised and amended by the second author before proceeding to data extraction. Data from the 19 selected articles included in the review were extracted by one author into the Excel spreadsheet and checked by the second author.

Data analysis

Where available, data were amenable to quantitative synthesis, random effects meta-analyses of proportions were performed using the metaprop suite of

commands on STATA 16. Variances were stabilised using the Freeman-Tukey double arcsine transformation. This approach provides better approximation and leads to results between 0% and 100% when synthesising proportions from small samples and multiple studies in meta-analyses. 14 Where possible, subgroup analysis was performed to examine differences between nulliparous women aged ≤30 years and nulliparous women of any age. Statistical heterogeneity was reported using I² and tau² statistics, since random effects meta-analyses were being performed. The I² value describes the percentage of the variability in effect estimates that is due to statistical heterogeneity (reflecting methodological diversity among the included studies) as opposed to chance. Conventionally, while an I² value <40% may not be significant, a value >50% may represent substantial heterogeneity and a value >75% may indicate considerable heterogeneity. 15 The tau² statistic measure of 'between-study variance', unlike the I² statistic, is not affected by size of included studies in a meta-analysis and hence may be considered more appropriate for estimating heterogeneity. 16 The effect of removing individual studies on the overall effect size (ES) was explored in sensitivity analyses (online supplemental material 4). Publication bias was examined by producing Doi plots and generating LFK index values, being considered a more appropriate measure of publication bias than funnel plots/Egger's test when performing meta-analyses of proportions.

Patient and public involvement

The FSRH is the UK organisation committed to meeting the highest SRH standards, ensuring improvements in population SRH and supporting SRH professionals. The FSRH's Contraceptive Priority Setting Partnership in liaison with the James Lind Alliance vielded over 700 responses from patients, practitioners and the public that identified: 'Which interventions increase uptake and continuation of effective contraception including longacting methods...?' as the top SRH research priority. 18 This influenced the research aims. IUD users attending a

| IUD brand/name | Copper (mm²) | Shape/design | Width (mm) | Arms' flexibility |
|---|--------------|---------------------|------------|-------------------|
| Currently available in the UK | | | | |
| Cu T380A/TCu 380 A/TT380 Slimline | 380 | T with arm bands | >30 | No |
| TCu 380A Nul/Mini TT380 slimline | 380 | T with arm bands | 23.2 | No |
| Multiload Cu 375 | 375 | Ω | 16–20.5 | Yes, flex down |
| Nova T380 | 380 | T without arm bands | >30 | Yes, flex up |
| Comparable to those available in the UK | | | | |
| Nova T200 | 200 | T without arm bands | ≥30 | Yes, flex up |
| TCu 300 | 300 | T without arm bands | >30 | No |
| Cu T200/TCu 200 | 200 | T without arm bands | >30 | No |
| TCu 220C | 220 | T without arm bands | >30 | No |

| Table 2 Characte | eristics of | f the included s | tudies | | | |
|---------------------------------------|-------------|---|---------------------------------|--|--|----------------------------|
| Study/authors | Year | Country | Study design | Study objectives | IUDs in study | Quality (MMAT score) |
| Abraham <i>et al</i> ¹⁹ | 2015 | USA | Prospective cohort | Relationship among young age, nulliparity and continuation of long-acting reversible contraceptives | Copper T380A | Good (7) |
| Akintomide <i>et al³⁰</i> | 2019 | UK | Retrospective records review | Discontinuation rates and reasons for discontinuation at 1 year of the small-sized Mini TT380 Slimline IUD compared with the standard-sized TT380 Slimline | Mini TT380 slimline TT380 slimline | Good (6) |
| Allonen <i>et al</i> ³¹ | 1980 | Denmark, Finland Sweden | RCT—double blind | Continuation rates and reasons for discontinuation at 2 years of the Nova T200 and Copper T200 | Nova T200 Copper T200 | Good (6) |
| Elkhateeb <i>et al</i> ³² | 2020 | Egypt | Prospective cohort | Acceptability of IUD use in nulliparous women by both women and healthcare providers | Copper T380A | Good (7) |
| Fugere ³³ | 1990 | Canada | Prospective cohort | Clinical performance of the Nova T200 IUD over 5 years | Nova T200 | Good (7) |
| Hall and Kutler ³⁴ | 2016 | USA | Prospective cohort | Experience and satisfaction of nulliparous intrauterine contraception users at 1, 6, 12 and 18 months | Copper T380A | Good (7) |
| Kaislasuo <i>et al³⁵</i> | 2015 | Finland | Prospective cohort | Menstrual characteristics and ultrasonographic uterine cavity measurements predict bleeding and pain in nulligravid women using intrauterine contraception | Nova T380 | Good (7) |
| Larsen <i>et al</i> ³⁶ | 1981 | Denmark | RCT—patient blind | Comparison of clinical performances of Progestasert and Copper T200 at 12 months | Copper T200 | Good (5) |
| Lewit ³⁷ | 1973 | USA | Prospective cohort | Two years' experience of the Copper T200 | Copper T200 | Good (7) |
| Liedholm and Sjöberg ³⁸ | 1974 | Sweden | Prospective cohort | Two years' experience with the Copper T200 and comparison between nulliparous and parous women | Copper T200 | Good (7) |
| Luukkainen <i>et al</i> ³⁹ | 1979 | Denmark, Finland Sweden | RCT—double blind | Experience and clinical performance of the Nova T200 and Copper T200 at 12 months | Nova T200 Copper T200 | Good (6) |
| Luukkainen <i>et al⁴⁰</i> | 1987 | Denmark, Finland, Hungary, Norway, Sweden | RCT—no blinding | Use-effectiveness and clinical performance of levonorgestrel-releasing and copper-releasing intrauterine devices at 12 months | Nova T200 | Good (6) |
| Mishell et al ⁴¹ | 1973 | USA | Prospective cohort | Continuation and clinical performance of TCu 200 in nulliparous women | Copper T200 | Good (7) |
| Nygren <i>et al</i> ⁴² | 1981 | Denmark, Finland Sweden | RCT—double blind | Continuation rates and reasons for discontinuation at 3 years of the Nova T200 and Copper T200 | Nova T200 Copper T200 | Good (7) |

Continued



Table 2 Continued

| Study/authors | Year | Country | Study design | Study objectives | IUDs in study | Quality (MMAT score) |
|---|------|---------|---|---|--|----------------------------|
| Ostergard and Gunning ⁴³ | 1979 | USA | RCT—blinding not stated | Continuation and clinical performances of Copper T200 and Dalkon Shield in nulligravid women at 12 months | Copper T200 | Good (5) |
| Otero-Flores <i>et</i> <i>al</i> ⁴⁴ | 2003 | Mexico | RCT—single (patient) blind | Comparison of clinical performance of three different IUDs in nulliparous women | Copper T380A Copper T380A Nul Multiload 375 sl | Good (6) |
| Roy et al ⁴⁵ | 1974 | USA | Prospective cohort | Experience with three different IUD models in nulliparous women at 1 year | Copper T380A Copper T300 Copper T200 | Good (7) |
| Sivin and Stern ⁴⁶ | 1979 | USA | RCT—double blind | Experience of three different IUDs in nulliparous and parous women | Copper T380A Copper T220C Copper T200 | Good (5) |
| Timonen et al ⁴⁷ | 1974 | Finland | Prospective, single (patient) blind | Use-effectiveness of Copper T300 at 1 year | Copper T300 | Good (7) |

sexual health clinic over a 4-week period were consulted about improving access to and use of intrauterine contraception. Their suggestions, which included studying women's experiences with IUDs, were used in developing the research question, aim and study design. The Consumer Panel of the North East Research Design Service was also consulted and the proposed research presented to them. The research plan was modified in line with their feedback.

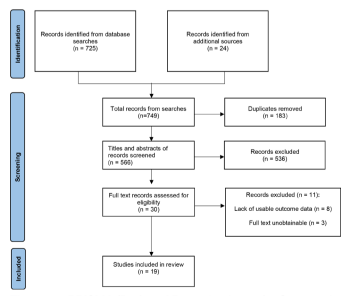


Figure 1 PRISMA (Preferred Reporting Items for Systematic Reviews and Meta-Analyses) flow diagram.

RESULTS

Only one study, a prospective (non-RCT) cohort study, provided information on an IUD available in the UK, solely involving nulliparous users aged under 30.19 This was inadequate to address the review question. As per the systematic review protocol, other studies on IUDs currently available in the UK or IUDs comparable to those available in the UK (table 1) involving nulliparous women of all ages (so not limited to those aged under 30) were also screened. An IUD was considered comparable if at least two out of its four characteristics (copper surface area, shape/design, width and arms flexibility) equated with IUDs currently used in the UK. So, for example, the Nova T200 was comparable because it has the same shape/design as a Nova T380, the same width as a Nova T380/Cu T380A/TCu 380A and TT380 slimline, and the same flexible arms as a Nova T380 (table 1).

Thirty records were obtained and their full texts assessed where possible. Eleven records were excluded, either for lack of usable outcome data (n=8⁵ ²⁰⁻²⁶) or because their full texts were unobtainable (n=3²⁷⁻²⁹) (see online supplemental material 5). A total of 19 studies on IUDs available or comparable to those available in the UK, involving 13 045 nulliparous women, were included in the systematic review (table 2). ^{19 30-47} Figure 1 depicts a PRISMA flow diagram detailing the search and selection process. ⁴⁸

All included studies were generally of good quality (mean 6.42 [5-7]; see online supplemental material 6 for quality and risk of bias assessments). The lowest MMAT score of five obtained was awarded to three RCTs

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| Table 3 Summary of findings | findings | | | | | | | |
|--|--|----------------------------------|-------------------------|--|-------------------------------------|--------------------------------------|-----------------------------------|----------------------------------|
| Study | IUD types (N*) | Age at insertion (years) | Study period | Continuation rates % (n) | Discontinuation rates % (n) | Removal for bleeding/pain % (n) | Expulsion % (n) | Pregnancy % (n) |
| Studies of IUD types currently available in the UK only involving nulliparous wo | ly available in the UK only in | volving nulliparous w | vomen aged ≤30 | | | | | |
| RCT | | | | | | | | |
| Otero-Flores et al ⁴⁴ *† | TCu 380A (375) TCu 380A Nul (367) ML Cu 375 sl (374) | 23.2±6.8 22.4±6.6 22.6±6.4 | 12 months | 30.7 (115) 91.3 (335) 89.0 (333) | 69.3 (260) 8.7 (32) 11.0 (41) | 61.6 (231) 3.81 (14) 6.68 (25) | 3.47 (13) 1.91 (7) 1.87 (7) | 1.07 (4) 0.54 (2) 0.00 (0) |
| Non-RCT | | | | | | | | |
| Abraham et al¹9 | Cu T380A (201) Cu T380A (44) Cu T380A (201) Cu T380A (44) | 20–25 <20 20–25 <20 | 12 months 24 months | 82 [95% CI 76-87] 79 [95% CI 64-89] 73 [95% CI 66-79] 64 [95% CI 48-77] | SU SU | su su | su su | su su |
| Hall and Kutler ³⁴ | Cu T 380A (21) | 18–30 | 12 months | 73.7 (14) | 26.3 (5) | 10.5 (2) | 10.5 (2) | 5.26 (1) |
| Studies of IUD types currently available in the UK involving nulliparous women | ly available in the UK involvir | ng nulliparous wome | n of all ages | | | | | |
| RCTs | | | | | | | | |
| Sivin and Stern ⁴⁶ ‡§ | TCu 380A (2254) TCu 220C (1301) TCu 200 (4215) | <20-35+ <20-35+ <20-35+ | 2 years | 55.7 57.8 54.2 | 44.3 42.2 45.8 | 21.9 19.5 16.8 | 7.8 9.8 9.8 | 0.8 1.6 5.1 |
| Non-RCTs | | | | | | | | |
| Akintomide et al³0 | TT380 Slimline (27) Mini TT380 Slimline (53) | 15–37 16–37 | 1 year | 66.7 (18) 86.8 (46) | 33.3 (9) 13.2 (7) | ns ns | 3.7 (1) 3.77 (2) | (0) 0 |
| Elkhateeb <i>et al</i> ³² | TCu 380A (90) | 16->30 | 6 months | 94.4 (85) | 5.6 (5) | ns | 0 (0) | ns |
| Kaislasuo <i>et al</i> ³⁵ † | Nova T380 (42) | 18–43 | 1 year | 83.3 (35) | 16.7 (7) | ns | 4.76 (2) | ns |
| Roy et al ⁴⁵ | TCu 380A (785) TCu 300 (347) TCu 200 (472) | <14->33 15->33 <14->33 | | 81.9 80.7 74.2 | 18.1 19.3 25.8 | 9.1 9.2 10.7 | 3.8 6.1 5.4 | 0.2 0.6 1.7 |
| Studies of IUD types comparable to those available in the UK involving nulliparous women of RCTs | rable to those available in the | e UK involving nullip | arous women of all ages | Ø | | | | |
| Luukkainen <i>et al</i> ³9§¶ | Nova T200 (ns) Cu T200 (ns) | <19->35 | 12 months | SU SU | ns ns | 15.3 23.4 | 6 10.8 | 0.53 |
| Allonen <i>et al</i> ³¹§¶ | Nova T200 (ns) Cu T200 (ns) | <19 ->35<19 ->35 | 24 months | ns ns | ns ns | 23.5 24 | 6.5 14 | 1.14 5.28 |
| Nygren <i>et al</i> ⁴² § | Nova T200 (ns) Cu T200 (ns) | <20 ->35 | 36 months | 36.9 31.0 | ns ns | 28.3 (74) 28.2 (68) | 10.3 (27) 10.7 (26) | 1.5 (4) 6.5 (15) |
| Larsen <i>et al</i> ³6§ | Cu T200 (99) | 15-44 | 12 months | 73 | 27** | 16 | 5 | - |
| Luukkainen <i>et al</i> ⁴⁰ | Nova T200 (77) | 17–40 | 12 months | 73.1 | 26.9** | 10.4 | 9.2 | 0 |
| Ostergard and Gunning ⁴³ | TCu 200 (117) TCu 200 (115) | 18–34 | 6 months 12 months | 88.9 (104) 73.0 (84) | 11.1 (13) 27.0 (31) | 6.0 (7) 12.2 (14) | 3.41 (4) 6.09 (7) | (0) 0 (0) |
| Non-RCTs | | | | | | | | |
| Fugere ³³ | Nova T200 (54) | 17–42 | 24 months | ns | ns | 17.2 | 1.9 | 0 |
| | | | | | | | | |

| 0 | |
|---|--|

| Study IUD types (N*) Age at insertion (years) Study period Lewit ³⁷ TCu-200 (2099) 15-49 1 year Nulligravid subgroups: TCu-200 (1585)† 15-49 1 year TCu-200 (130) 15-19 1 year TCu-200 (130) 15-19 1 year TCu-200 (1513) 20-24 1 year TCu-200 (1513) 25-29 1 year TCu-200 (153) 30-34 1 year TCu-200 (499) 35-49 1 year TCu-200 (208) 14-40 12 months Liedholm and Sjöberg ³⁸ T-Cu 200 (208) 14-40 12 months | | hates Discontinuation rates Removal for % (n) bleeding/pai 26.7 9.4 24.1 9.6 | Removal for bleeding/pain % (n) Expulsion % (n) | | |
|--|----------------|--|---|-------------------|--------------------|
| TCu-200 (2099) 15–49 Nulligravid subgroup: 15–49 TCu-200 (1585)† Age subgroups: 15–19 TCu-200 (1130) 20–24 TCu-200 (1513) 25–29 TCu-200 (1543) 30–34 TCu-200 (449) 35–49 | | 26.7 24.1 | | Expulsion % (n) | Fregnancy % (n) |
| Nulligravid subgroup: 15–49 TCu-200 (1585)† Age subgroups: 15–19 TCu-200 (2468) 20–24 TCu-200 (1513) 25–29 TCu-200 (449) 35–49 T-Cu 200 (208) 14–40 | | 24.1 | 9.4 | 10.7 | 1.3 |
| Age subgroups: 15–19 TCu–200 (1130) 20–24 TCu–200 (1513) 25–29 TCu–200 (683) 30–34 TCu–200 (449) 35–49 T-Cu 200 (208) 14–40 | | | 9.6 | 8.7 | 8.0 |
| TCu-200 (2468) 20-24 TCu-200 (1513) 25-29 TCu-200 (683) 30-34 TCu-200 (449) 35-49 T-Cu 200 (208) 14-40 | | 32.7 | 7 | ر ت | 83 |
| TCu-200 (1513) 25–29 TCu-200 (683) 30–34 TCu-200 (449) 35–49 T-Cu 200 (208) 14–40 | | 26.2 | 8.3 | 8.5 | 2.8 |
| TCu-200 (683) 30–34 TCu-200 (449) 35–49 T-Cu 200 (208) 14–40 | | 22.4 | 5.8 | 8.7 | 1.5 |
| TCu-200 (449) 35–49 T-Cu 200 (208) 14–40 | | 18.3 | 7.9 | 9 | 0.4 |
| T-Cu 200 (208) 14–40 | | 14.8 | 8.9 | 3.1 | 0.3 |
| | | 29.8 | 18.1 | 0.5 | 2.9 (6) |
| | 24 months 60.3 | 39.7 | 28 | 0.5 | 2.9 (6) |
| Mishell et al ⁴¹ § TCu 200 (471) 14–33 3 months | | 7.4 | 2.8 | 2.6 | 0.2 |
| | 6 months 84.5 | 15.5 | 5.8 | 4.7 | 0.4 |
| 12 months | | 25.8 | 10.7 | 5.4 | 1.7 |
| Timonen et al ⁴⁷ T Cu-300 (138) <25-40+ 12 months | 12 months 84.7 | 15.3 | 7.2 | 1.6 | 1.6 |

"Sample size or participants excluding those lost to follow-up or removals.

†Nulligravid women only.

‡A combination of double blind studies.

§Net cumulative rate and such some such as the summary of the such as the such

Estimated continuation rates at 12 months of IUD types from the included studies Table 4

| | Continuation rates with numbers of patier | Continuation rates with numbers of patients (n) and statistical heterogeneity (tau² and l²) values of studies included in subgroup | s of studies included in subgroup |
|----------------------|--|--|---|
| IUD type | Nulliparous women aged <30 | Nulliparous women of any age | Overall effect size (all studies) |
| TCu 380A* | TCu 380A* 81.60% (95% CI 76.52% to 86.21%)† (n=264; tau^2 =0.0; t^2 =0.0%, p =0.69) ^{19.34} | 80.97% (95% CI 76.04% to 85.48%) (n=971; tau^2 =0.005; t^2 =27.6%, p=0.25) ^{19 30 45} | 81.93% (95% CI 79.66% to 84.09%) (n=1235; tau ² =0.0; l ² =0.0%, p=0.62) ^{19 30 34 45} |
| Smaller TCu 380A‡ | Not applicable—only one study group | 91.02% (95% CI 88.01% to 93.64%) (n=420; tau ² =0.0; I^2 =0.0%, p=0.51) ^{30.44} | 91.02% (95% CI 88.01% to 93.64%) (n=420; tau ² =0.0; I ² =0.0%, p=0.51) ^{30.44} |
| TCu 300 | Not applicable—no study | 81.92% (95% CI 78.35% to 85.24%) (n=485; tau²=0.0; l²=17.3%, p=0.27) ^{45,47} | 81.92% (95% CI 78.35% to 85.24%) (n=485; tau²=0.0; l²=17.3%, p=0.27) ^{45,47} |
| TCu 200 | 73.03% (95% CI 67.63% to 78.10%) (n=5111; \tan^2 =0.010; l^2 =94.2%, p =<0.01) ³⁷ | 76.51% (95% CI 72.67% to 80.14%) (n=3277; \tan^2 =0.012; l^2 =84.0%, p=<0.01) ^{37-39 41 43 45} | 75.44% (95% CI 72.32% to 78.43%) (n=8388; tau^2 =0.012; l^2 =89.9%, p =<0.01) ^{37-3941 43 45} |
| Nova T200 | Not applicable—no study | 73.21% (95% CI 70.10% to 76.22%) (n=818; tau²=0.0; l²=0.0%, p=0.94) ^{39.40} | 73.21% (95% CI 70.10% to 76.22%) (n=818; tau^2 =0.0; I^2 =0.0%, p =0.94) ^{39.40} |
| | | | |

†Includes women aged 30 from Hall and Kutler's study data. ‡TCu 380A Nul/Mini TT380 Slimline IUDs. *Excludes Otero-Flores et al's study data. IUD, intrauterine device. BMJ Open: first published as 10.1136/bmjopen-2021-060606 on 3 October 2022. Downloaded from http://bmjopen.bmj.com/ on August 27, 2023 by guest. Protected by copyright.

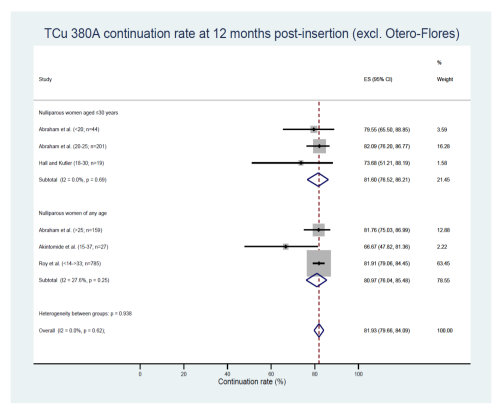


Figure 2 TCu 380A continuation rates (excluding Otero-Flores). ES, effect size.

published in 1979 and 1981 and may relate to inade-quate reporting. $^{36\ 43\ 46}$ Their reports did not confirm that randomisation had been appropriately performed, $^{36\ 46}$ randomised groups were comparable at baseline, $^{43\ 46}$ nor that outcome assessors were blinded to the intervention provided. $^{36\ 43}$

Although the outcome data obtained were considered homogeneous, studies' designs, participant ages and parity, and IUD types were not; making a quantitative synthesis of the outcome data in totality inappropriate. Results were therefore grouped into three to include studies involving: (1) IUD types currently available in the

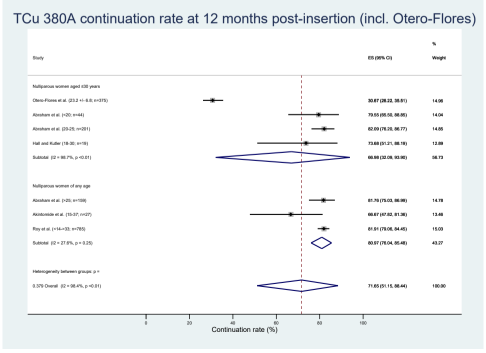


Figure 3 TCu 380A continuation rates (including Otero-Flores). ES, effect size.

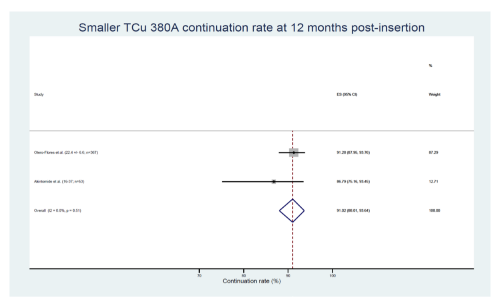


Figure 4 Smaller TCu 380A continuation rates. ES, effect size.

UK and only nulliparous women aged ≤30; (2) IUD types currently available in the UK and nulliparous women of all ages; (3) IUD types comparable to those available in the UK and nulliparous women of all ages (table 3). The estimated continuation rates at 12 months by IUD type, obtained from the included studies with data amenable to synthesis, is reported in table 4. Tau² values for heterogeneity of the included studies are provided separately (see online supplemental material 7).

Studies of IUD types currently available in the UK only involving nulliparous women aged ≤30

Three studies—Abraham et al¹⁹, Hall and Kutler³⁴ and Otero-Flores et al⁴⁴—reported on IUDs in women aged ≤30 involving the Copper T380A IUD (TCu 380A or Cu T380A). 19 34 44 The TCu 380A data obtained from Otero-Flores et al⁴⁴ was an outlier, with 30.7% reported as the

continuation rate at 12 months. 44 This was much lower than for the other two studies with a pooled estimate of 81.60% (95% CI 76.52% to 86.21%) ^{19 34} (figure 2). When the Otero-Flores et al data were included in this TCu 380A meta-analysis, nulliparous women ≤30 years of age at 12 months had a continuation rate of 66.98% (95% CI 32.09% to 93.90%) (figure 3).

Continuation was also higher with age at 12 and 24 months when nulliparous TCu 380A IUD users aged <20 and 20–25 were compared (table 3). 19

Studies of IUD types currently available in the UK involving nulliparous women of all ages

Five studies reporting data pertaining to seven population subgroups were amenable to meta-analysis examining the proportion of women continuing to use the TCu 380A IUD at 12 months post insertion. 19 30 34 44 45 The pooled

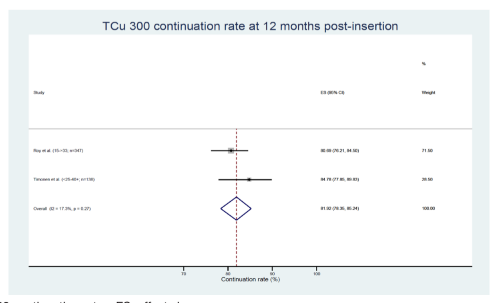


Figure 5 TCu 300 continuation rates. ES, effect size.

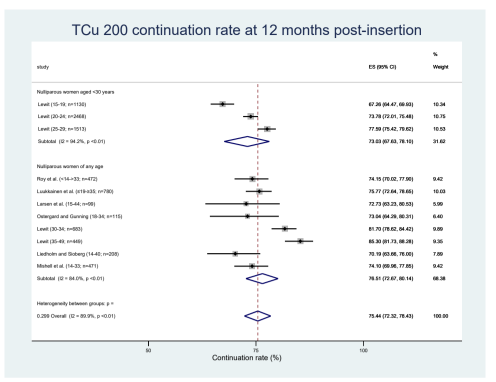


Figure 6 TCu 200 continuation rates. ES, effect size.

estimated continuation rate of the Copper T380A IUD type in nulliparous women of all ages from four studies was 81.93% (95% CI 79.66% to 84.09%). 19303445 Additionally, statistical heterogeneity was found to be low/absent but was not statistically significant (tau 2 =0.0, I 2 =0.0%, p=0.62). Sensitivity analysis confirmed that the overall ES was largely robust to the exclusion of individual studies (-1.01% to +0.21% change in ES; see online supplemental material 4).

The estimated TCu 380A continuation rate in nulliparous women of all ages remained good at 71.65% (95% CI 51.15% to 88.44%; tau²=0.299, I²=98.4%, p=<0.01) when

the Otero-Flores *et al* data were included⁴⁴ (figure 3). An LFK index value of 6.77 identified major Doi plot asymmetry consistent with publication bias (see online supplemental material 8).

Individual studies showed the TCu 380A had higher discontinuation related to bleeding/pain and expulsion^{34 44 46} when compared with IUDs of smaller size or those with flexible arms^{30 44} (table 3).

The highest continuation rates at 12 months were reported with smaller sized IUDs—the Copper 380A Nul (TCu 380A Nul: 91.3%), Multiload Copper 375 sl (ML Cu 375 sl: 89%) and Mini TT380 slimline (86.8%) (table 3).

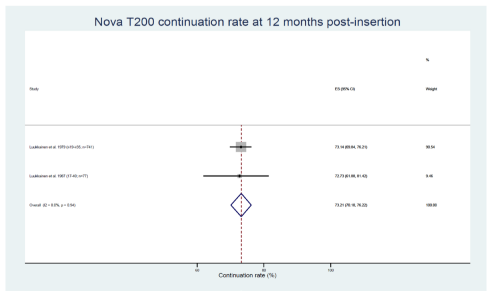


Figure 7 Nova T200 continuation rates. ES, effect size.



These data were obtained from only two studies whose participants were aged $15\text{--}37.^{30.44}$ Meta-analysis of continuation rate data on the TCu 380A Nul/Mini TT380 slimline IUD type gave a weighted average of 91.02% (95% CI 88.01% to 93.64%) (figure 4). These smaller IUDs were also associated with the lowest rates of removals for bleeding/pain (3.80%–6.68%) and expulsion (1.87%–3.77%) reported in nulliparous women at 12 months (table 3).

Studies of IUD types comparable to those in the UK involving nulliparous women of all ages

Two studies reporting data pertaining to two population subgroups were amenable to meta-analysis examining the proportion of women continuing to use the Copper T300 IUD (TCu 300) at 12months post insertion, with an overall ES of 81.92% (95% CI 78.35% to 85.24%, see figure 5). 45 47

Seven studies reporting data pertaining to 11 population subgroups were amenable to meta-analysis examining the proportion of women continuing to use the Copper T200 IUD (TCu 200 or Cu T200) at 12 months post insertion, with a weighted average of 75.44% (95% CI 72.32% to 78.43%, see figure 6). 36-38 40 41 43 45 These studies were also amenable to meta-analysis examining the proportion of women discontinuing the TCu 200 at 12 months post insertion due to bleeding and/or pain, expulsion and pregnancy (see online supplemental material 9). For these meta-analyses, nulliparous women aged <30 years compared with nulliparous women of any age were less likely to continue to use the TCu 200 at 12 months (73.03% (95% CI 67.63% to 78.10%) vs 76.51% (95% CI 72.67% to 80.14%)), and less likely to discontinue the TCu 200 due to bleeding and/or pain (7.05% (95% CI 5.59% to 8.65%) vs 12.77% (95% CI 8.48 to 17.78%)). Nulliparous women aged <30 years compared with nulliparous women of any age were however more likely to discontinue the TCu 200 due to expulsion (10.52% (95% CI 7.17% to 14.41%) vs 4.93% (95% CI 2.93% to 7.39%)) and pregnancy (2.19% (95% CI 1.47% to 3.05%) vs 1.15% (95% CI 0.54% to 1.95%)). The overlapping confidence intervals for these two ESs suggest the difference in effect is not statistically significant, and therefore may or may not be clinically significant. Statistical heterogeneity values for overall TCu 200 continuation rates as well as discontinuation rates for bleeding/ pain and expulsion were $tau^2=0.012$, $I^2=89.9\%$, p=<0.01; $\tan^2 = 0.025 \text{ I}^2 = 93.2\%$, p=<0.01; and $\tan^2 = 0.018$, $\ln^2 = 96.3\%$, p=<0.01 respectively (see figure 6 and online supplemental material 9). Sensitivity analyses confirmed that the overall ESs were largely robust due to the exclusion of individual studies (see online supplemental material 4). In all cases, their LFK index values identified major Doi plot asymmetry consistent with publication bias (see online supplemental material 8).

Continuation rates were seen to progressively improve with age where Lewit³⁷ reported rates in nulliparous TCu

200 users by age groups 15–19, 20–24, 25–29, 30–34 and $35-49^{37}$ (table 3).

Two studies reporting data pertaining to two population subgroups were amenable to meta-analysis examining the proportion of women continuing to use the Nova T200 at 12 months post insertion, with a weighted average of 73.21% (95% CI 70.10% to 76.22%, see figure 7). 39 40

Studies also showed that IUDs with flexible arms (Nova T, Multiload) were associated with higher continuation and lower removal rates for bleeding/pain, expulsion and pregnancy when compared with IUDs with rigid arms (Cu T or TCu)^{31 39 44} (table 3).

DISCUSSION

Findings and interpretation

Evidence on IUDs currently used in nulliparous women aged under 30 is limited. These findings estimate the continuation rate for the recommended TCu 380A IUD 11 to be 81% at 12 months post insertion based on four studies involving young nulliparous women. $^{19\,30\,34\,45}$ This was the same estimate for the TCu 300 based on two studies. $^{45\,47}$ Smaller sized and flexible IUDs had higher continuation rates of 86%–91% in this group of women, based on two studies, as well as fewer removals for bleeding/pain and expulsion compared with the TCu 380A or IUDs of the same rigid design or size. $^{30\,44}$ Lower continuation rates of 75% and 73% were obtained for the Cu T200 and Nova T200 based on eight studies. $^{36-41\,43\,45}$

The study by Otero-Flores et al was the only reported RCT solely involving IUDs currently used in the UK with nulliparous women aged ≤30.⁴⁴ Over a thousand nulliparous women aged 15-30 were randomised to receive three different IUDs: TCu 380A (width 32mm), TCu 380A Nul (width 23 mm) and ML Cu 375 sl (width≤20 mm), the latter two being primarily designed for nulliparous women. The TCu 380A overall rate of discontinuation (69.3%) and bleeding/pain as a reason for discontinuation (61.6%) were significantly higher than for TCu 380A Nul (8.7% and 3.81%) and ML Cu 375 sl (11.0% and 6.68%), as well as significantly different from rates reported by other included studies involving the TCu 380A. This could be because the TCu 380A considerably differs in size from the TCu 380A Nul and ML Cu 375 sl IUDs, and Otero-Flores et al also exclusively involved nulligravid participants (as opposed to nulliparous).

Sivin and Stern⁴⁶ was the only other RCT involving a TCu 380A that reported separately on nulliparous users.⁴⁶ However, their TCu 380A discontinuation and bleeding/pain rates, 44.3% and 21.9%, respectively, were obtained at 2 years and their participants were aged <20–35+ years.

The disparity in discontinuation rates reported by Otero-Flores *et al*¹⁴ and Sivin and Stern⁴⁶ suggests that the findings by Otero-Flores *et al* may be unreliable. But it may in fact be inappropriate to directly compare other studies' TCu 380A data, including that of Sivin and Stern, to Otero-Flores *et al*'s data. Their studies' designs as well as participants' ages, gravidity/parity, environments and



reported durations of use were not the same. Otero-Flores et al's participants were younger (≤30 years), exclusively nulligravid, 'highly educated' and based in a Mexico city with free access to healthcare in the millenial era, with the study being single-(patient) blinded. This contrasts with most studies involving the TCu 380A or similar IUDs where participants were more likely to be aged 30 years or older and parous with unspecified educational attainment. The Sivin and Stern study population were living and accessing healthcare (which was not stated to have been free) across the USA, in the late 1970s (over two decades earlier than the Otero-Flores et al's study, and not long after the Dalkon Shield era), with the study being double-blinded. Other explanations for the disparity could be that the modern younger nulligravid cohort may be less tolerant of unwanted IUD effects, and that some contraceptive research may be less likely to acknowledge participants' reasons and wishes for early IUD discontinuation.⁴⁹

The TCu 200 IUD was $\geq 33\,\mathrm{mm}$ in width and/or height so perhaps larger than a standard-sized TCu 380A. ⁵⁰ IUD size may contribute to pain, which may explain TCu 200's lower continuation rates compared with the TCu 380A. However the TCu 300, of the same design and size as the TCu 200, ⁴⁷ unexpectedly had a higher continuation rate than the TCu 200. This is because higher copper content has been associated with more bleeding which contributes to early discontinuation. ⁵¹ The TCu 300 data were limited to two studies that both had total MMAT scores of 7, ⁴⁵ ⁴⁷ whereas the TCu 200 data had been obtained from seven studies with MMAT scores of 7, ³⁷ ³⁸ ⁴¹ ⁴⁵ 6³⁹ and 5, ⁴³ respectively.

Strengths and limitations

This is the first systematic review to explore IUD types in younger aged nulliparous women. It has included all observational studies that provided information on IUD continuation or reasons for discontinuation in this user group. Non-restriction to RCTs may be considered a limitation, but a realist approach of expanding the inclusion criteria where RCT evidence is lacking could be commendable and more representative of routine practice. Using the MMAT, the quality of reviewed and included studies in this systematic review was good overall.

Articles for inclusion were unfortunately limited to publications in the English language. There was an absence of studies on IUDs currently available in the UK and solely involving women aged under 30. This warranted including all ages if women under 30 years were involved, and up to (≤) 30 years for the TCu 380A data and meta-analysis because of the ages of the Hall and Kutler study participants (18–30 years). Many studies did not report all the required information, hence some included studies had missing information (table 3). Most studies did not differentiate between nulligravid and nulliparous participants, many age ranges were not specific (eg, ≤19–≥35), while some reports, for example, Sivin and Stern, ⁴⁶ were a combination of individual studies. Similarly, it appeared

common for older studies to only state numbers (rather than rates or percentages), or only graphically depict data on continuation rates or unwanted effects. It is also not unusual for a systematic review to include such studies, for example, Hubacher⁷, and to calculate or measure rates accordingly, as has been done in this review. These are potential limitations which are not considered to impact the validity of the review. All mitigating actions that were taken have also been appropriately stated.

Relevance of findings

IUD use in young nulliparous women has been established to be safe, effective and acceptable. ^{52–54} It is recommended that women are provided with the most appropriate IUD types for their uterine cavity size. Uterine cavity width (measurable using a cavimeter or ultrasonography, not routinely practised) in addition to uterine length (routinely measured using a hysterome) should be recognised as influencing IUD type choice. ^{29 55–57} This systematic review suggests which IUD types may be more suitable for younger aged nulliparous women and emphasises the need for further research.

Recommendations

Strengthening the evidence for contraceptive choice and continuation is needed to improve sexual health in younger aged women. Prospective observational studies that include various IUD designs and types, and detailed reporting of users' experiences could facilitate a better understanding of early IUD discontinuation and reasons for discontinuation based on IUD types. Studies designed to overcome the challenges of recruiting large numbers from varied demographic backgrounds, significant loss to follow-up, and time or funding constraints are also likely to yield data widely applicable to IUC provision in and outside the UK.

CONCLUSION

Research is lacking on outcomes with the IUD types currently in use by young nulliparous women in the UK. Available evidence estimates a continuation rate of 81% at 12 months for the recommended standard-sized TCu 380A IUD in these women. More studies are needed to better estimate continuation rates for smaller sized and flexible IUDs in this user group.

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writing—review and editing. MM: Meta-analysis, writing—original draft, review and editing. JR: Contributed to research idea, study design, protocol, funding applications, and project administration, as well as supervision and writing—review and editing. All authors approved the final version.

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PRISMA 2020 Checklist

| Section and Topic | Item # | Checklist item | Location where item is reported |
|-------------------------------|-----------|--|-----------------------------------|
| TITLE | | | |
| Title | 1 | Identify the report as a systematic review. | Page 1 |
| ABSTRACT | | | |
| Abstract | 2 | See the PRISMA 2020 for Abstracts checklist. | Page 1 Supplemental material 1 |
| INTRODUCTION | | | |
| Rationale | 3 | Describe the rationale for the review in the context of existing knowledge. | Pages 1-2 |
| Objectives | 4 | Provide an explicit statement of the objective(s) or question(s) the review addresses. | Page 2 |
| METHODS | • | | |
| Eligibility criteria | 5 | Specify the inclusion and exclusion criteria for the review and how studies were grouped for the syntheses. | Pages 2-3 |
| Information sources | 6 | Specify all databases, registers, websites, organisations, reference lists and other sources searched or consulted to identify studies. Specify the date when each source was last searched or consulted. | Page 2 Supplemental material 3 |
| Search strategy | 7 | Present the full search strategies for all databases, registers and websites, including any filters and limits used. | Page 2 Supplemental material 3 |
| Selection process | 8 | Specify the methods used to decide whether a study met the inclusion criteria of the review, including how many reviewers screened each record and each report retrieved, whether they worked independently, and if applicable, details of automation tools used in the process. | Pages 2-3 |
| Data collection process | 9 | Specify the methods used to collect data from reports, including how many reviewers collected data from each report, whether they worked independently, any processes for obtaining or confirming data from study investigators, and if applicable, details of automation tools used in the process. | Pages 2-3 |
| Data items | 10a | List and define all outcomes for which data were sought. Specify whether all results that were compatible with each outcome domain in each study were sought (e.g. for all measures, time points, analyses), and if not, the methods used to decide which results to collect. | Page 3 |
| | 10b | List and define all other variables for which data were sought (e.g. participant and intervention characteristics, funding sources). Describe any assumptions made about any missing or unclear information. | Page 3 |
| Study risk of bias assessment | 11 | Specify the methods used to assess risk of bias in the included studies, including details of the tool(s) used, how many reviewers assessed each study and whether they worked independently, and if applicable, details of automation tools used in the process. | Pages 2-3 Supplemental material 6 |
| Effect measures | 12 | Specify for each outcome the effect measure(s) (e.g. risk ratio, mean difference) used in the synthesis or presentation of results. | Page 3 |
| Synthesis methods | 13a | Describe the processes used to decide which studies were eligible for each synthesis (e.g. tabulating the study intervention characteristics and comparing against the planned groups for each synthesis (item #5)). | Page 3 |
| | 13b | Describe any methods required to prepare the data for presentation or synthesis, such as handling of missing summary statistics, or data conversions. | Page 3 |
| | 13c | Describe any methods used to tabulate or visually display results of individual studies and syntheses. | Page 3 |
| | 13d | Describe any methods used to synthesize results and provide a rationale for the choice(s). If meta-analysis was performed, describe the model(s), method(s) to identify the presence and extent of statistical heterogeneity, and software package(s) used. | Page 3 |
| | 13e | Describe any methods used to explore possible causes of heterogeneity among study results (e.g. subgroup analysis, meta- | Page 3 |



PRISMA 2020 Checklist

| Section and Topic | Item # | Checklist item | Location where item is reported |
|-------------------------------|-----------|--|--|
| | | regression). | |
| | 13f | Describe any sensitivity analyses conducted to assess robustness of the synthesized results. | Page 3 Supplemental material 4 |
| Reporting bias assessment | 14 | Describe any methods used to assess risk of bias due to missing results in a synthesis (arising from reporting biases). | Page 3 |
| Certainty assessment | 15 | Describe any methods used to assess certainty (or confidence) in the body of evidence for an outcome. | Page 3 |
| RESULTS | | | |
| Study selection | 16a | Describe the results of the search and selection process, from the number of records identified in the search to the number of studies included in the review, ideally using a flow diagram. | Page 5 Figure 1 |
| | 16b | Cite studies that might appear to meet the inclusion criteria, but which were excluded, and explain why they were excluded. | Page 5 Supplemental material 5 |
| Study characteristics | 17 | Cite each included study and present its characteristics. | Table 2 |
| Risk of bias in studies | 18 | Present assessments of risk of bias for each included study. | Table 2 |
| Results of individual studies | 19 | For all outcomes, present, for each study: (a) summary statistics for each group (where appropriate) and (b) an effect estimate and its precision (e.g. confidence/credible interval), ideally using structured tables or plots. | Pages 5-11 Tables 3-4 Figures 2–7 Supplemental material 9 |
| Results of | 20a | For each synthesis, briefly summarise the characteristics and risk of bias among contributing studies. | Supplemental material 4,7,8 |
| syntheses | 20b | Present results of all statistical syntheses conducted. If meta-analysis was done, present for each the summary estimate and its precision (e.g. confidence/credible interval) and measures of statistical heterogeneity. If comparing groups, describe the direction of the effect. | Pages 7-11 Table 4 Figures 2–7 |
| | 20c | Present results of all investigations of possible causes of heterogeneity among study results. | Pages 7-11 Table 4 Figures 2–7 Supplemental material 7,8 |
| | 20d | Present results of all sensitivity analyses conducted to assess the robustness of the synthesized results. | Supplementary material 4 |
| Reporting biases | 21 | Present assessments of risk of bias due to missing results (arising from reporting biases) for each synthesis assessed. | Pages 7-11 Table 4 Figures 2–7 Supplemental material 4,7,8 |
| Certainty of evidence | 22 | Present assessments of certainty (or confidence) in the body of evidence for each outcome assessed. | Pages 7-11 Table 4 |



PRISMA 2020 Checklist

| Section and Topic | Item # | Checklist item | Location where item is reported |
|--|-----------|--|---------------------------------|
| | | | Figures 2–7 |
| | | | Supplemental material 4,7,8 |
| DISCUSSION | | | |
| Discussion | 23a | Provide a general interpretation of the results in the context of other evidence. | Pages 11-12 |
| | 23b | Discuss any limitations of the evidence included in the review. | Page 12 |
| | 23c | Discuss any limitations of the review processes used. | Page 12 |
| | 23d | Discuss implications of the results for practice, policy, and future research. | Page 12 |
| OTHER INFORMA | TION | | |
| Registration and | 24a | Provide registration information for the review, including register name and registration number, or state that the review was not | Page 2 |
| protocol | | registered. | Supplemental material 2 |
| | 24b | Indicate where the review protocol can be accessed, or state that a protocol was not prepared. | Page 2 |
| | | | Supplemental material 2 |
| | 24c | Describe and explain any amendments to information provided at registration or in the protocol. | Pages 2 and 5 |
| Support | 25 | Describe sources of financial or non-financial support for the review, and the role of the funders or sponsors in the review. | Page 13 |
| Competing interests | 26 | Declare any competing interests of review authors. | Page 13 |
| Availability of data, code and other materials | 27 | Report which of the following are publicly available and where they can be found: template data collection forms; data extracted from included studies; data used for all analyses; analytic code; any other materials used in the review. | Not applicable |

From: Page MJ, McKenzie JE, Bossuyt PM, Boutron I, Hoffmann TC, Mulrow CD, et al. The PRISMA 2020 statement: an updated guideline for reporting systematic reviews. BMJ 2021;372:n71. doi: 10.1136/bmj.n71

For more information, visit: http://www.prisma-statement.org/

PROSPERO International prospective register of systematic reviews



Copper intrauterine contraception discontinuation in nulliparous and young women Hannat Akintomide, Pam Barnes, Nataliya Brima, Judith Rankin

Citation

Hannat Akintomide, Pam Barnes, Nataliya Brima, Judith Rankin. Copper intrauterine contraception discontinuation in nulliparous and young women. PROSPERO 2019 CRD42019120969 Available from: http://www.crd.york.ac.uk/PROSPERO/display record.php?ID=CRD42019120969

Review question

Which copper intrauterine devices are associated with higher discontinuation rates in young and nulliparous women?

Searches

Databases [including the Cochrane Library, the Database of Abstracts and Reviews of Effects (DARE), MEDLINE (Ovid), Excerpta Medica Database (EMBASE), Turning Research into Practice (TRIP) database and National Electronic Library of Health] and relevant websites [including Bandolier, Medicines and Healthcare products Regulatory Agency, Faculty of Sexual and Reproductive Healthcare, Royal College of Obstetricians and Gynaecologists, Department of Health, Medical Defence Unions, National Institute for Health and Care Excellence, Scottish Intercollegiate Guidelines, World Health Organisation and Google Scholar] will be searched using MeSH terms combined with key words for relevant articles published from 1966 to date. Reference lists of relevant articles will also be searched to identify more articles. The full texts of relevant articles will be screened, duplicates excluded and then data from selected articles included in the review.

Randomised controlled trials (RCTs) involving copper intrauterine devices (IUDs) available or comparable to those in the UK published in English will be included. Other studies that report on the main outcome (observational and qualitative studies) will be included and/or summarised if the number of RCTs eligible for inclusion are too few to answer the review question.

Key words

Copper intrauterine device related: copper intrauterine device, copper intrauterine contraceptive device, copper intrauterine contraception, copper coil, IUD

Nulliparous related: nulliparous, nulligravid, never pregnant, never delivered Young women related: young women, adolescent, aged under, teenage

Types of study to be included

Inclusion criteria: Articles published in English on studies in women who are nulliparous and aged under 30 that involved copper intrauterine devices available, or of the same design and size to those available, in the UK

Exclusion criteria: Articles not published in English, studies solely in parous women aged 30 or over, or that involved copper intrauterine devices not available, or not of the same design and size to those available, in the UK.

Condition or domain being studied

Copper intrauterine contraception in nulliparous and young women

Participants/population

Women who are nulliparous and aged under 30

Intervention(s), exposure(s)

Copper intrauterine devices available or comparable to those in the UK

Comparator(s)/control

Any IUD, other contraceptive or no contraception where applicable

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Context

Copper intrauterine devices (IUDs) are of various shapes, sizes, copper surface area and copper distribution on the frame of the device. There are many types of IUDs available in the UK but none shown to be associated with better outcomes in nulliparous and young women. The identification and use of those IUDs associated with less discontinuation could improve outcomes including satisfaction and continuation rates of intrauterine contraception in nulliparous and younger women.

Main outcome(s)

Copper intrauterine contraception discontinuation rates in nulliparous and young women based on type of IUD

Timing and effect measures

Additional outcome(s)

Reasons for IUD discontinuation

Timing and effect measures

Data extraction (selection and coding)

The abstracts of published articles obtained from the literature and websites searches will be reviewed by two authors to assess eligibility for inclusion in the systematic review based on the inclusion/exclusion criteria. All retrieved full texts of published articles will be reviewed to agree which studies to include in the systematic review, with disagreements resolved by the third author. All retrieved articles to be included in the systematic review will undergo a quality assessment using a risk of bias tool applicable to the type of study.

Main data to be extracted:

type of copper intrauterine device (IUD)

age of women

gravidity/parity of women

place/time of IUD insertion

IUD discontinuation rate(s)

reason(s) for IUD discontinuation

Risk of bias (quality) assessment

All retrieved articles to be included in the systematic review will undergo a quality assessment. One author will complete the inclusion criteria checklist while the second author will review the checklist, with disagreements resolved by the third author/consensus. Retrieved articles with a high risk of bias will be excluded from the systematic review.

Strategy for data synthesis

Data from the included studies will be extracted using a standardised form by one author while the second author will check these. Disagreements will be resolved by a further review of the study with the third author and consensus. One author will enter the extracted data into Review Manager (RevMan®) Software while the second author will again check these for accuracy. It is planned that aggregate data will be used. However, individual data on the intervention and population of interest (IUDs in nulliparous and young women aged under 30) will be extracted where studies have reported on this subgroup their outcomes in conjunction with other population subgroups or study outcomes.

A quantitative synthesis is planned based on the expected homogeneity of the data to be obtained for the main outcome to be studied. This homogeneous data will be combined for meta-analysis. Heterogeneous

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data, some of which is expected to be obtained on the additional outcome, will be narratively synthesised.

Analysis of subgroups or subsets

IUDs of same size and design will be grouped and discontinuation rates presented based on IUD type.

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Anticipated or actual start date 28 January 2019

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Language English

Country England

Stage of review Review_Ongoing

Subject index terms status Subject indexing assigned by CRD

Subject index terms

Contraception; Copper; Female; Humans; Intrauterine Devices; Parity; Pregnancy

Date of registration in PROSPERO 07 February 2019

Date of publication of this version 07 February 2019

Details of any existing review of the same topic by the same authors

Stage of review at time of this submission

Versions 07 February 2019

| PROSPERO International prospective register of systematic reviews | | NHS al Institute for alth Research |
|---|---------|--|
| Stage | Started | Completed |
| Preliminary searches | Yes | No |
| Piloting of the study selection process | Yes | No |
| Formal screening of search results against eligibility criteria | No | No |
| Data extraction | No | No |
| Risk of bias (quality) assessment | No | No |
| Data analysis | No | No |

PROSPERO

This information has been provided by the named contact for this review. CRD has accepted this information in good faith and registered the review in PROSPERO. CRD bears no responsibility or liability for the content of this registration record, any associated files or external websites.

Table – Search Strategies

| Databases and additional sources search | Search term(s) used | Limits | Records identified |
|---|---|------------------|--------------------|
| Allied and Complementary Medicine (AMED) | (copper intrauterine).ti,ab OR (copper | Title, Abstract | |
| British Nursing Index (BNI) | intrauterine device).ti,ab OR (copper coil).ti,ab | English language | |
| Cumulative Index to Nursing and Allied Health Literature (CINAHL) | OR (copper IUD).ti,ab OR (copper T).ti,ab | | |
| Excerpta Medica Database (EMBASE) | | | |
| Nursing and Allied Health Professionals Database (EMCARE) | | | 725 |
| Health Management Information Consortium (HMIC) | | | |
| General Medical Database (MEDLINE) | | | |
| Psychology and Allied Fields (PsychINFO) | | | |
| PubMed | | | |
| The Cochrane Library | 'copper intrauterine' | - | |
| Database of Abstracts and Reviews of Effects (DARE) | | | |
| Turning Research into Practice (TRIP) | | | |
| Bandolier | | | |
| National Electronic Library of Health | | | |
| Medicines and Healthcare products Regulatory Agency (MHRA) | | | |
| Faculty of Sexual and Reproductive Healthcare (FSRH) | | | 22 |
| Royal College of Obstetricians and Gynaecologists (RCOG) | | | |
| Department of Health | | | |
| National Institute for Health and Care Excellence (NICE) | | | |
| Scottish Intercollegiate Guidelines, | | | |
| World Health Organisation (WHO) | | | |
| Google Scholar | 'copper intrauterine device young nulliparous' | - | |

TCu 380A continuation at 12 months post-insertion – sensitivity analysis

| Subgroup 1 (Nulliparous women aged <30 years) | (81.60% (95% CI 76.52-86.21%)) | | |
|---|--------------------------------|--|--|
| Excluding Abraham et al. (<20) | 82.04% (95% CI 76.48-87.04%) | | |
| Excluding Abraham et al. (20-25) | 78.01% (95% CI 66.60-87.74%) | | |
| Excluding Hall and Kutler (18-30) | 81.83% (95% CI 76.66-86.49%) | | |
| | | | |
| Subgroup 2 (Nulliparous women of any age) | (80.97% (95% CI 76.04-85.48%)) | | |
| Excluding Abraham et al. (>25) | 81.99% (95% CI 79.19-84.63%) | | |
| Excluding Akintomide et al. (15-37) | 81.94% (95% CI 79.41-84.34%) | | |
| Excluding Roy et al. (14-33) | 80.12% (95% CI 73.92-85.70%) | | |
| | | | |
| Overall effect size (all studies) | (81.93% (95% CI 79.66-84.09%)) | | |
| Excluding Abraham et al. (<20) | 81.84% (95% CI 79.13-84.40%) | | |
| Excluding Abraham et al. (20-25) | 81.44% (95% CI 78.16-84.53%) | | |
| Excluding Hall and Kutler (18-30) | 81.87% (95% CI 79.60-84.03%) | | |
| Excluding Abraham et al. (>25) | 81.57% (95% CI 78.38-84.58%) | | |
| Excluding Akintomide et al. (15-37) | 82.14% (95% CI 79.87-84.31%) | | |
| Excluding Roy et al. (14-33) | 80.92% (95% CI 76.93-84.64%) | | |

TCu 200 continuation at 12 months post-insertion – sensitivity analysis

| Subgroup 1 (Nulliparous women aged <30 years) | (73.03% (95% CI 67.63-78.10%)) | | |
|---|--------------------------------|--|--|
| Excluding Lewit (15-19) | 75.26% (95% CI 73.90-76.59%) | | |
| Excluding Lewit (20-24) | 73.33% (95% CI 71.62-75.00%) | | |
| Excluding Lewit (25-29) | 71.78% (95% CI 70.30-73.24%) | | |
| | | | |
| Subgroup 2 (Nulliparous women of any age) | (76.51% (95% CI 72.67-80.14%)) | | |
| Excluding Roy et al. (14-33) | 76.83% (95% CI 72.49-80.91%) | | |
| Excluding Luukkainen et al. (19-35) | 76.53% (95% CI 71.86-80.91%) | | |
| Excluding Larsen et al. (15-44) | 76.85% (95% CI 72.79-80.67%) | | |
| Excluding Ostergard and Gunning (18-34) | 76.84% (95% CI 72.76-80.69%) | | |
| Excluding Lewit (30-34) | 75.59% (95% CI 71.42-79.54%) | | |
| Excluding Lewit (35-49) | 75.20% (95% CI 71.98-78.29%) | | |
| Excluding Liedholm and Sioberg (14-40) | 77.32% (95% CI 73.40-81.01%) | | |
| Excluding Mishell et al. (14-33) | 76.84% (95% CI 72.51-80.91%) | | |
| | | | |
| Overall effect size (all studies) | (75.44% (95% CI 72.32-78.43%)) | | |
| Excluding Lewit (15-19) | 76.43% (95% CI 73.71-79.04%) | | |
| Excluding Lewit (20-24) | 75.59% (95% CI 71.81-79.17%) | | |
| Excluding Lewit (25-29) | 76.16% (95% CI 71-60-78.56%) | | |
| Excluding Roy et al. (14-33) | 75.56% (95% CI 72.16-78.81%) | | |
| Excluding Luukkainen et al. (19-35) | 75.38% (95% CI 71.89-78.72%) | | |
| Excluding Larsen et al. (15-44) | 75.60% (95% CI 72.34-78.70%) | | |
| Excluding Ostergard and Gunning (18-34) | 75.59% (95% CI 72.33-78.71%) | | |
| Excluding Lewit (30-34) | 74.72% (95% CI 71.59-77.73%) | | |

| Excluding Lewit (35-49) | 74.37% (95% CI 71.53-77.10%) |
|--|------------------------------|
| Excluding Liedholm and Sioberg (14-40) | 75.87% (95% CI 72.61-78.98%) |
| Excluding Mishell et al. (14-33) | 75.56% (95% CI 72.16-78.81%) |

TCu 200 discontinuation at 12 months due to pain/bleeding – sensitivity analysis

| Subgroup 1 (Nulliparous women aged <30 years) | (7.05% (95% CI 5.59-8.65%)) | | |
|---|-------------------------------|--|--|
| Excluding Lewit (15-19) | 7.31% (95% CI 6.52-8.14%) | | |
| Excluding Lewit (20-24) | 6.31% (95% CI 5.41-7.27%) | | |
| Excluding Lewit (25-29) | 7.88% (95% CI 7.02-8.78%) | | |
| | | | |
| Subgroup 2 (Nulliparous women of any age) | (12.77% (95% CI 8.48-17.78%)) | | |
| Excluding Roy et al. (14-33) | 13.10% (95% CI 8.10-19.06%) | | |
| Excluding Luukkainen et al. (19-35) | 11.02% (95% CI 8.41-13.92%) | | |
| Excluding Larsen et al. (15-44) | 12.40% (95% CI 7.87-17.76%) | | |
| Excluding Ostergard and Gunning (18-34) | 12.86% (95% CI 8.20-18.35%) | | |
| Excluding Lewit (30-34) | 13.61% (95% CI 8.83-19.22%) | | |
| Excluding Lewit (35-49) | 13.79% (95% CI 9.10-19.25%) | | |
| Excluding Liedholm and Sioberg (14-40) | 12.08% (95% CI 7.56-17.45%) | | |
| Excluding Mishell et al. (14-33) | 13.13% (95% CI 8.13-19.08%) | | |
| | | | |
| Overall effect size (all studies) | (10.87% (95% CI 7.98-14.15%)) | | |
| Excluding Lewit (15-19) | 11.37% (95% CI 8.08-15.12%) | | |
| Excluding Lewit (20-24) | 11.23% (95% CI 7.70-15.32%) | | |
| Excluding Lewit (25-29) | 11.52% (95% CI 8.34-15.14%) | | |
| Excluding Roy et al. (14-33) | 10.90% (95% CI 7.77-14.47%) | | |
| Excluding Luukkainen et al. (19-35) | 9.32% (95% CI 7.62-11.17%) | | |
| Excluding Larsen et al. (15-44) | 10.51% (95% CI 7.58-13.86%) | | |
| Excluding Ostergard and Gunning (18-34) | 10.78% (95% CI 7.77-14.20%) | | |
| Excluding Lewit (30-34) | 11.23% (95% CI 8.01-14.92%) | | |
| Excluding Lewit (35-49) | 11.34% (95% CI 8.17-14.94%) | | |
| Excluding Liedholm and Sioberg (14-40) | 10.26% (95% CI 7.40-13.53%) | | |
| Excluding Mishell et al. (14-33) | 10.92% (95% CI 7.78-14.50%) | | |

TCu 200 discontinuation at 12 months due to expulsion – sensitivity analysis

| Subgroup 1 (Nulliparous women aged <30 years) | (10.52% (95% CI 7.17-14.41%)) | | |
|---|-------------------------------|--|--|
| Excluding Lewit (15-19) | 8.59% (95% CI 7.74-9.48%) | | |
| Excluding Lewit (20-24) | 11.21% (95% CI 10.03-12.44%) | | |
| Excluding Lewit (25-29) | 10.36% (95% CI 9.38-11.38%) | | |
| | | | |
| Subgroup 2 (Nulliparous women of any age) | (4.93% (95% CI 2.93-7.39%)) | | |
| Excluding Roy et al. (14-33) | 4.85% (95% CI 2.57-7.78%) | | |
| Excluding Luukkainen et al. (19-35) | 4.17% (95% CI 2.68-5.96%) | | |
| Excluding Larsen et al. (15-44) | 4.92% (95% CI 2.79-7.58%) | | |
| Excluding Ostergard and Gunning (18-34) | 4.80% (95% CI 2.69-7.46%) | | |
| Excluding Lewit (30-34) | 4.74% (95% CI 2.41-7.76%) | | |
| Excluding Lewit (35-49) | 5.24% (95% CI 3.03-7.99%) | | |
| Excluding Liedholm and Sioberg (14-40) | 5.84% (95% CI 3.95-8.07%) | | |

| Excluding Mishell et al. (14-33) | 4.85% (95% CI 2.57-7.77%) |
|---|-----------------------------|
| | |
| Overall effect size (all studies) | (6.44% (95% CI 4.49-8.69%)) |
| Excluding Lewit (15-19) | 5.76% (95% CI 4.14-7.61%) |
| Excluding Lewit (20-24) | 6.16% (95% CI 3.87-8.93%) |
| Excluding Lewit (25-29) | 6.16% (95% CI 3.96-8.79%) |
| Excluding Roy et al. (14-33) | 6.55% (95% CI 4.47-8.99%) |
| Excluding Luukkainen et al. (19-35) | 6.01% (95% CI 3.98-8.42%) |
| Excluding Larsen et al. (15-44) | 6.54% (95% CI 4.51-8.91%) |
| Excluding Ostergard and Gunning (18-34) | 6.46% (95% CI 4.43-8.83%) |
| Excluding Lewit (30-34) | 6.47% (95% CI 4.36-8.95%) |
| Excluding Lewit (35-49) | 6.87% (95% CI 4.87-9.18%) |
| Excluding Liedholm and Sioberg (14-40) | 7.29% (95% CI 5.39-9.45%) |
| Excluding Mishell et al. (14-33) | 6.55% (95% CI 4.47-8.99%) |

TCu 200 discontinuation at 12 months due to pregnancy – sensitivity analysis

| Subgroup 1 (Nulliparous women aged <30 years) | (2.19% (95% CI 1.47-3.05%)) | | |
|---|-----------------------------|--|--|
| Excluding Lewit (15-19) | 2.27% (95% CI 1.82-2.75%) | | |
| Excluding Lewit (20-24) | 1.83% (95% CI 1.35-2.39%) | | |
| Excluding Lewit (25-29) | 2.63% (95% CI 2.13-3.18%) | | |
| | | | |
| Subgroup 2 (Nulliparous women of any age) | (1.15% (95% CI 0.54-1.95%)) | | |
| Excluding Roy et al. (14-33) | 1.07% (95% CI 0.40-1.99%) | | |
| Excluding Luukkainen et al. (19-35) | 0.96% (95% CI 0.38-1.75%) | | |
| Excluding Larsen et al. (15-44) | 1.18% (95% CI 0.53-2.05%) | | |
| Excluding Ostergard and Gunning (18-34) | 1.31% (95% CI 0.65-2.16%) | | |
| Excluding Lewit (30-34) | 1.35% (95% CI 0.70-2.18%) | | |
| Excluding Lewit (35-49) | 1.31% (95% CI 0.62-2.20%) | | |
| Excluding Liedholm and Sioberg (14-40) | 1.00% (95% CI 0.42-1.78%) | | |
| Excluding Mishell et al. (14-33) | 1.07% (95% CI 0.40-1.99%) | | |
| | | | |
| Overall effect size (all studies) | (1.49% (95% CI 0.96-2.13%)) | | |
| Excluding Lewit (15-19) | 1.39% (95% CI 0.81-2.09%) | | |
| Excluding Lewit (20-24) | 1.34% (95% CI 0.83-1.94%) | | |
| Excluding Lewit (25-29) | 1.48% (95% CI 0.87-2.22%) | | |
| Excluding Roy et al. (14-33) | 1.46% (95% CI 0.89-2.16%) | | |
| Excluding Luukkainen et al. (19-35) | 1.40% (95% CI 0.83-2.09%) | | |
| Excluding Larsen et al. (15-44) | 1.53% (95% CI 0.98-2.19%) | | |
| Excluding Ostergard and Gunning (18-34) | 1.62% (95% CI 1.07-2.26%) | | |
| Excluding Lewit (30-34) | 1.69% (95% CI 1.18-2.29%) | | |
| Excluding Lewit (35-49) | 1.64% (95% CI 1.10-2.28%) | | |
| Excluding Liedholm and Sioberg (14-40) | 1.41% (95% CI 0.88-2.06%) | | |
| Excluding Mishell et al. (14-33) | 1.46% (95% CI 0.89-2.16%) | | |

Table – Characteristics of studies excluded following full text assessment

| Study / Authors | Year | Country | Study Design | Study Objectives | Reasons for Exclusion |
|---------------------|------|-------------------|-------------------|---|---|
| Akintomide et al[5] | 2021 | Austria, Finland, | Prospective | Secondary analysis of continuation, unwanted effects and | Undifferentiable results - IUD type |
| | | Germany, Poland, | cohort | cost consequences at 1 year in IUD users ≤30 in the | categories based on IUD characteristics |
| | | Sweden, UK | | European Active Surveillance Study for Intrauterine Devices | rather than brand or name of IUD |
| Garbers et al[20] | 2013 | USA | Retrospective | Prevalence and predictors of IUD discontinuation at 6 | Undifferentiable results; varied duration; |
| | | | records review | months in 306 Cu T380A users | 23 excluded from continuation analysis |
| Goldstuck[21] | 1980 | UK | Prospective | Clinical evaluation of the combined multiload copper 250- | Undifferentiable results; disparity |
| | | | cohort (selected) | mini IUD in selected nulliparous women | between data in tables and text |
| Hindle[27] | 1978 | Unable to confirm | | Clinical evaluation and follow-up on 3,829 IUD procedures | Full text unobtainable |
| Lete et al[22] | 1998 | Spain | Prospective | Evaluation of IUD use in nulliparous women compared to | Data reported as incidence of events |
| | | | cross-sectional | parous women over a 12-year period | rather than rates |
| Ogedengbe et | 1991 | Nigeria | Prospective | A comparison efficacy and discontinuation at 1 year of | Parity of participants not detailed (mean |
| al[23] | | | cohort | multiload and copper-T IUDs sequentially assigned to users | parity 4); only one nulliparous participant |
| Patnaik[28] | 2003 | India | Unable to confirm | Uptake, satisfaction, retention and reasons for | Full text unobtainable |
| | | | | discontinuation of the copper T IUD | |
| Petersen et al[29] | 1991 | Unable to confirm | RCT – | Significance of endometrial cavity length in the clinical | Full text unobtainable |
| | | | double blind | performance of IUDs in nulligravidae | |
| Phillips et al[24] | 2017 | USA | Retrospective | Comparison of continuation and performance of | Undifferentiable results |
| | | | records review | levonorgestrel and copper intrauterine devices over 5 years | |
| Sivin and | 1981 | USA | Prospective | Clinical performance of the TCu 380A IUD over 4 years | Undifferentiable results |
| Tatum[25] | | | cohort | | |
| Teal et al[26] | 2015 | USA | Retrospective | Evaluation of the success and safety of intrauterine device | Undifferentiable results |
| | | | records review | (IUD) placement in adolescents based on age and parity | |
| | | | | | |

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Table – Quality Assessment of Included Studies Using the Mixed Methods Appraisal Tool (MMAT) version 2018

| Study / Authors | Design Category | | Responses to MMAT Questions (and Scores): Yes (1) / No (0) / Can't Tell (0) | | | | | | |
|----------------------------|------------------------------|-------------|---|-------------|-------------|-------------|-------------|-------------|-------|
| | | Screening 1 | Screening 2 | Appraisal 1 | Appraisal 2 | Appraisal 3 | Appraisal 4 | Appraisal 5 | Total |
| Abraham et al 2015 | Quantitative, non-randomised | yes | yes | yes | yes | yes | yes | yes | 7 |
| Akintomide et al 2019 | Quantitative, non-randomised | yes | yes | yes | yes | no | yes | yes | 6 |
| Allonen et al 1980 | Quantitative, randomised | yes | yes | can't tell | yes | yes | yes | yes | 6 |
| Elkhateeb et al 2020 | Quantitative, non-randomised | yes | yes | yes | yes | yes | yes | yes | 7 |
| Fugere 1990 | Quantitative, non-randomised | yes | yes | yes | yes | yes | yes | yes | 7 |
| Hall and Kutler 2015 | Quantitative, non-randomised | yes | yes | yes | yes | yes | yes | yes | 7 |
| Kaislasuo et al 2015 | Quantitative, non-randomised | yes | yes | yes | yes | yes | yes | yes | 7 |
| Larsen et al 1981 | Quantitative, randomised | yes | yes | can't tell | yes | yes | no | yes | 5 |
| Lewit 1973 | Quantitative, non-randomised | yes | yes | yes | yes | yes | yes | yes | 7 |
| Liedholm and Sjoberg 1974 | Quantitative, non-randomised | yes | yes | yes | yes | yes | yes | yes | 7 |
| Luukkainen et al 1979 | Quantitative, randomised | yes | yes | can't tell | yes | yes | yes | yes | 6 |
| Luukkainen et al 1987 | Quantitative, randomised | yes | yes | yes | yes | yes | no | yes | 6 |
| Mishell et al 1973 | Quantitative, non-randomised | yes | yes | yes | yes | yes | yes | yes | 7 |
| Nygren et al 1981 | Quantitative, randomised | yes | yes | yes | yes | yes | yes | yes | 7 |
| Ostergard and Gunning 1979 | Quantitative, randomised | yes | yes | yes | can't tell | yes | no | yes | 5 |
| Otero-Flores et al 2003 | Quantitative, randomised | yes | yes | yes | yes | yes | no | yes | 6 |
| Roy et al 1974 | Quantitative, non-randomised | yes | yes | yes | yes | yes | yes | yes | 7 |
| Sivin and Stern 1979 | Quantitative, randomised | yes | yes | can't tell | can't tell | yes | yes | yes | 5 |
| Timonen et al 1974 | Quantitative, non-randomised | yes | yes | yes | yes | yes | yes | yes | 7 |
| | | | | | | | | | |

Tau² Values for Heterogeneity of Included Studies

| IUD type | Tau ² Values for Heterogeneity of Included Studies for Continuation Rates | | | | | | |
|--|---|------------------------------|-----------------------------------|--|--|--|--|
| | Nulliparous women aged <30 | Nulliparous women of any age | Overall effect size (all studies) | | | | |
| TCu 380A excluding Otero- Flores data | 0.0ª [19, 34] | 0.005 [19, 30, 45] | 0.0 [19, 30, 34, 45] | | | | |
| TCu 380A including Otero- Flores data | 0.487 [19, 34, 44] | 0.005 [19, 30, 44, 45] | 0.299 [19, 30, 34, 44, 45] | | | | |
| Smaller TCu 380A ^b | not applicable – only one study group | 0.0 [30, 44] | 0.0 [30, 44] | | | | |
| TCu 300 | not applicable – no study | 0.0 [45, 47] | 0.0 [45, 47] | | | | |
| TCu 200 | 0.010 [37] | 0.012 [37-39, 41, 43, 45] | 0.012 [37-39, 41, 43, 45] | | | | |
| Nova T200 | not applicable – no study | 0.0 [39, 40] | 0.0 [39, 40] | | | | |
| | Tau ² Values for Heterogeneity of Included Studies for Discontinuation Rates | | | | | | |
| TCu 200 discontinuation due to bleeding/pain | 0.001 [37] | 0.036 [36-39, 41, 43, 45] | 0.025 [36-39, 41, 43, 45] | | | | |
| TCu 200 discontinuation due to expulsion | 0.010 [37] | 0.018 [36-39, 41, 43, 45] | 0.018 [36-39, 41, 43, 45] | | | | |
| TCu 200 discontinuation due to pregnancy | 0.002 [37] | 0.005 [36-39, 41, 43, 45] | 0.004 [36-39, 41, 43, 45] | | | | |

a – includes women aged 30 from Hall and Kutler study data; b – TCu 380A Nul/Mini TT380 Slimline IUDs

Supplementary material – Doi plots

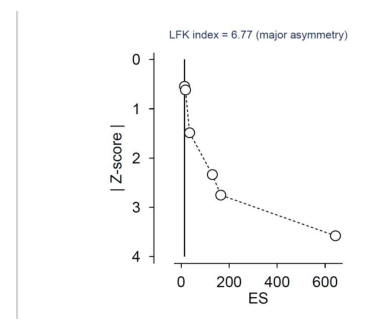


Figure 1 - Doi plot for TCu 380A continuation at 12 months

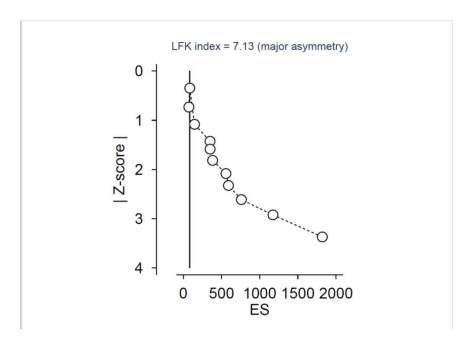


Figure 2 – Doi plot for TCu 200 continuation at 12 months

Supplementary material – Doi plots

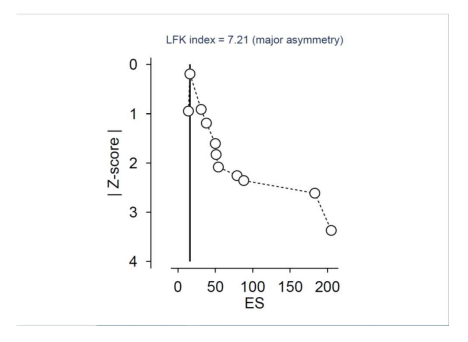


Figure 3 – Doi plot for TCu 200 discontinuation at 12 months due to bleeding/pain

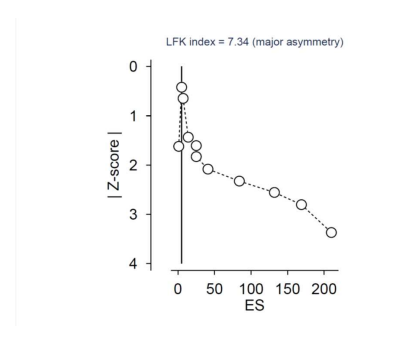


Figure 4 – Doi plot for TCu 200 discontinuation at 12 months due to expulsion

Supplementary material – Doi plots

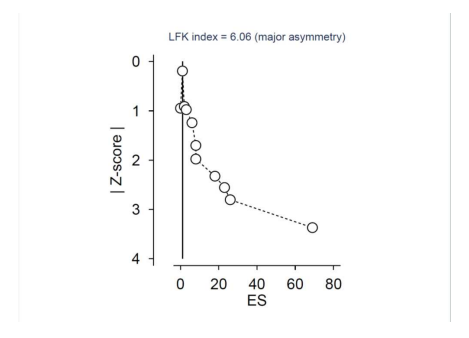


Figure 5 – Doi plot for TCu 200 discontinuation due to pregnancy

Supplementary material - TCu 200 discontinuation rates due to pain/bleeding, expulsion and pregnancy

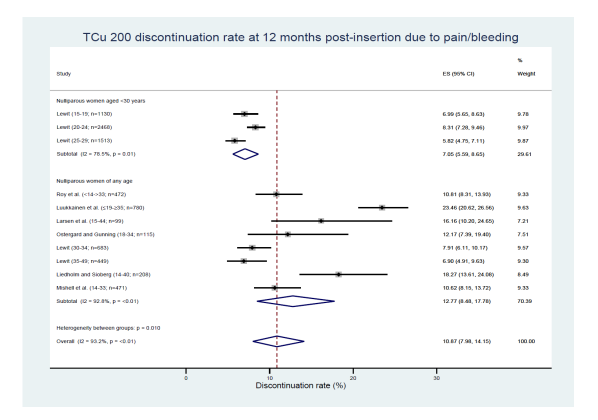


Figure 1 - TCu 200 discontinuation at 12 months due to pain/bleeding

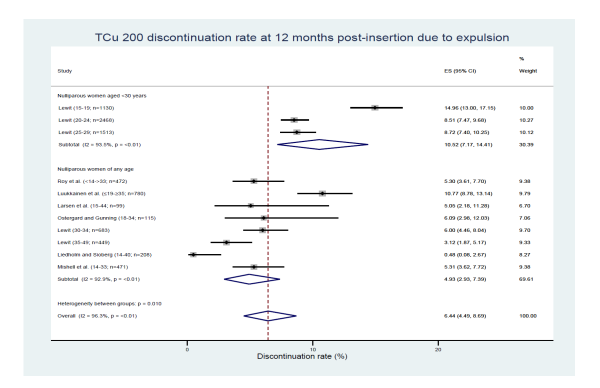


Figure 2 – TCu 200 discontinuation at 12 months due to expulsion

Supplementary material – TCu 200 discontinuation rates due to pain/bleeding, expulsion and pregnancy

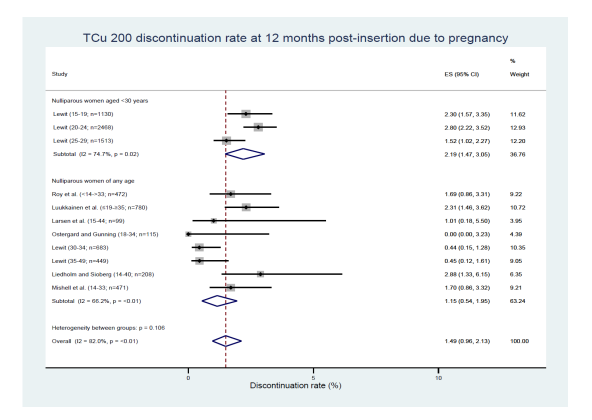


Figure 3 – TCu 200 discontinuation at 12 months due to pregnancy