Review article

Does insertion and use of an intrauterine device increase the risk of pelvic inflammatory disease among women with sexually transmitted infection?

A systematic review

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Abstract

Concerns exist as to whether the insertion of copper and levonorgestrel-releasing intrauterine devices (IUDs) increases the risk of pelvic inflammatory disease (PID) among women with sexually transmitted infection (STI). We searched the MEDLINE database for all articles published between January 1966 and March 2005 that included evidence relevant to IUDs and STIs and PID. None of the studies that examined women with STIs compared the risk of PID between those with insertion or use of an IUD and those who had not received an IUD. We reviewed indirect evidence from six prospective studies that examined women with insertion of a copper IUD and compared risk of PID between those with STIs at the time of insertion with those with no STIs. These studies suggested that women with chlamydial infection or gonorrhea at the time of IUD insertion were at an increased risk of PID relative to women without infection. The absolute risk of PID was low for both groups (0–5% for those with STIs and 0–2% for those without).

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1. Introduction

Approximately 160 million women worldwide use intrauterine devices (IUDs), making the IUD the most popular contraceptive method after sterilization [1]. IUDs are safe and highly effective, long-acting methods of contraception. Theoretical concerns exist, however, as to whether women who have sexually transmitted infections (STIs) and who are therefore at risk of developing pelvic inflammatory disease (PID) further increase their risk of PID with the insertion and use of IUDs. The majority of cases of PID are caused by sexually transmitted microorganisms, with endogenous flora of the lower genital tract playing a lesser role [2]. It is hypothesized that when an IUD is inserted, sexually transmitted microorganisms that may be present in the endocervical canal could be transported to the uterine cavity [3].

We conducted this systematic review in preparation for an Expert Working Group of international family planning experts convened by the World Health Organization (WHO) in October 2003 to develop and revise medical eligibility criteria for contraceptive use. In this report, we describe the evidence obtained through our systematic review regarding whether insertion and use of a copper or levonorgestrel-releasing IUD increases the risk of PID among women with STIs; we also provide the WHO recommendations that were derived in part from this evidence.

2. Materials and methods

We searched the MEDLINE database for all articles (in all languages) published in peer-reviewed journals between January 1966 and March 2005, for evidence relevant to the insertion and use of copper and levonorgestrel-releasing intrauterine devices and STIs and PID. The following search strategy was used: [mirena.mp. or levonorgestrel.mp. and (exp intrauterine devices/ or (iud or iucd or ius).mp. or (intrauterine adj3 system).mp. or (intra-uterine adj3 sys-
We searched reference lists from articles identified by the search, as well as key review articles, to identify additional articles. We did not try to identify unpublished articles or abstracts from scientific conferences. In an attempt to locate additional articles, we contacted an expert in the field but did not learn of any other published literature.

2.1. Selection of studies

The search strategy identified a total of 365 articles that considered IUDs as well as STIs and PID. Our primary goal was to identify studies that examined whether, among women with STIs, the insertion and use of copper or levonorgestrel-releasing IUDs increased the risk of PID over that of non-IUD users. After reviewing the titles and abstracts of these articles as well as the full article when necessary, we did not identify any studies that met these criteria. When we looked for indirect evidence that could help us assess our study question, we identified six studies that examined women who received a copper IUD and assessed whether women with STIs at the time the device was inserted had a greater risk of PID than women without STIs at the time of insertion [4–9]. We contacted one author to clarify whether screening for STI occurred before or after insertion in the author’s study. We did not identify any direct or indirect evidence that examined the use of levonorgestrel-releasing IUDs among women with STIs and risk of PID.

2.2. Assessment of the study quality and synthesis of the data

We summarized and systematically assessed the evidence through the use of standard abstract forms [10]. We assessed the quality of each individual piece of evidence using a preliminary draft of the Grades of Recommendation Assessment, Development and Evaluation (GRADE) system (Appendix A) [11].

Because six studies reported percentages of women with STIs and PID, crude relative risks were calculated from the study reports for this systematic review (Table 1). We assessed heterogeneity by examining the characteristics of the participants included in each study. We did not calculate a summary statistic for PID because of the heterogeneity of the studies, but summary graphs of the relative risks are included in this review (Figs. 1 and 2). We have also included evidence that summarizes the studies reviewed (Table 2).

3. Results

We did not identify any studies that examined whether copper or levonorgestrel-releasing IUD insertion or use modified the risk of PID among women with STIs, i.e., studies that examined a group of women with current STIs and compared the risk of PID in women who had an IUD inserted with women who did not undergo IUD insertion. We did not identify any studies regarding levonorgestrel-releasing IUDs either.

In the absence of direct evidence, we identified six studies that provided indirect evidence, i.e., they examined whether women who had an STI at the time of copper IUD insertion were at a greater risk of developing PID than women without an STI at the time of IUD insertion (Table 1) [4–9]. These studies varied substantially in their design and methodology. Three were prospective studies whose primary goal was to evaluate screening for STIs prior to IUD

<table>
<thead>
<tr>
<th>Author, year</th>
<th>Country</th>
<th>Follow-up</th>
<th>STI present at insertion</th>
<th>No STI present at insertion</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sinei et al., 1990 [6]</td>
<td>Kenya</td>
<td>1 month</td>
<td>4.3% (5) of 117 women(a)</td>
<td>1.3% (9) of 670 women</td>
</tr>
<tr>
<td>Sinei et al., 1990 [6]</td>
<td>Kenya</td>
<td>1 month</td>
<td>3.0% (7) of 232 women(b)</td>
<td>1.1% (15) of 1339 women</td>
</tr>
<tr>
<td>Pap-Akeson et al., 1992 [8]</td>
<td>Sweden</td>
<td>2 years</td>
<td>0% (0) of 13 women</td>
<td>2.1% (9) of 432 women</td>
</tr>
<tr>
<td>Walsh et al., 1994 [9]</td>
<td>USA (LA)</td>
<td>3 months</td>
<td>0% (0) of 7 women</td>
<td>0.5% (2) of 435 women</td>
</tr>
<tr>
<td>Skjeldestad et al., 1996 [7]</td>
<td>Norway</td>
<td>3 months</td>
<td>0% (0) of 5 women</td>
<td>0% (0) of 952 women</td>
</tr>
<tr>
<td>Fauñdes et al., 1998 [4]</td>
<td>Brazil</td>
<td>1 month</td>
<td>5.2% (1) of 19 women</td>
<td>0% (0) of 308 women</td>
</tr>
<tr>
<td>Fauñdes et al., 1998 [4]</td>
<td>Brazil</td>
<td>1 month</td>
<td>10.5% (2) of 19 women(c)</td>
<td>0% (0) of 308 women</td>
</tr>
<tr>
<td>Morrison et al., 1999 [5]</td>
<td>Kenya</td>
<td>4 months</td>
<td>3.1% (1) of 32 women</td>
<td>0.4% (2) of 548 women</td>
</tr>
</tbody>
</table>

Adapted from: Best K. IUD not recommended for increased STD risk. Network 2000;20:12–5.

\(a\) Placebo group only.
\(b\) Combining the placebo and antibiotic groups (there were no treatment effects).
\(c\) Includes one woman suspected of having PID (not diagnosed).
\(d\) Cervical swabs taken 1 month after insertion of IUD.
insertion [4,5,7]. The three others were randomized controlled trials evaluating either the use of antibiotics or the placement of strings of the IUD as approaches to decrease PID among women receiving the device; for our purposes, however, we considered these to be prospective observational studies, as we used the presence of an STI at time of IUD insertion as the “exposure” and development of PID as the outcome [6,8,9]. All six studies reported rates of PID following IUD insertion; three of the studies did not provide or describe diagnostic criteria for PID [4,7,8]. The degree of screening for STI risk to determine eligibility for inserting an IUD differed among studies, with four studies using some type of syndromic screening [4–6,8]; two of these studies also used behavioral risk assessment [4,5]. Two of the six studies did not report details of their criteria for screening and exclusion [7,9]. Three studies tested for cervical chlamydial infection only [4,7,8], and the rest for both chlamydial infection and gonorrhea [5,6,9]. In four of the studies, results of STI testing were obtained after insertion [4,6–8]; in one, laboratory results were obtained

Fig. 1. Crude relative risk of PID among women with and without STI at insertion [1,2].

Fig. 2. Crude relative risk of PID among women with and without chlamydial infection or gonorrhea at insertion [1,2].
<table>
<thead>
<tr>
<th>Author, year, support</th>
<th>Objective</th>
<th>Study Design</th>
<th>Population</th>
<th>Outcome (and assessment)</th>
<th>Screening</th>
<th>Results</th>
<th>Weaknesses</th>
<th>Quality</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sinei et al., 1990 [6], FHI, USAID</td>
<td>To determine the effectiveness of 200 mg of doxycycline given orally at the time of insertion in reducing the occurrence of PID</td>
<td>Double-blind RCT for 1 month; Kenya 1984–1986</td>
<td>1813 prospective IUD users; 904 in doxycycline group and 909 in placebo group</td>
<td>PID: defined by the Infectious Disease Society of Obstetrics and Gynecology in the United States</td>
<td>– Women with an active PID were excluded from the study. There is no other information about screening based on signs and symptoms of chlamydial infection or gonorrhea – Samples were taken before insertion, but results were obtained after insertion</td>
<td>Women with cervical infections at insertion: RR for PID among women with cervical infections vs. women with no cervical infections (not including women who had both chlamydial infection and gonorrhea): – Placebo only: 3/27 women with gonorrhea had PID; 2/90 women with chlamydial infection had PID – Combined (placebo and doxycycline): 3/46 women with gonorrhea had PID; 4/186 women with chlamydial infection had PID Women with no cervical infection at insertion: – Placebo only: 9/670 had PID – Combined (placebo and doxycycline): 15/1339 had PID</td>
<td>– Short follow-up (1 month)</td>
<td>Low</td>
</tr>
<tr>
<td>Pap-Akeson, et al., 1992 [8], support not stated</td>
<td>To study the influence of the position of the threads of an intrauterine contraceptive device on the development of genital tract infection</td>
<td>RCT — 2 years; Sweden – IUD was either inserted with threads up or threads down</td>
<td>445 prospective IUD users: 208 threads up, 237 in the threads-down position</td>
<td>Salpingitis: diagnostic criteria not stated Endometritis: diagnostic criteria not stated</td>
<td>– Women with signs and symptoms of genital infection and irregular bleeding and history of salpingitis were excluded – Samples were taken before insertion, but results were obtained after insertion</td>
<td>Women with cervical infection at insertion: – All 13 women were treated with tetracyclines with the IUD still in place and had no complications or symptoms for 2 years (although not all women were followed up for that time)</td>
<td>– Did not specify the diagnostic criteria for salpingitis or endometritis</td>
<td>Very low</td>
</tr>
</tbody>
</table>
– 13/445 (2.9%) had asymptomatic chlamydial infection at insertion (no difference between the groups)

Women with cervical infection at insertion:
– 6 women with chlamydial infection had no signs or symptoms during the study
– Woman with gonorrhea had IUD removed

Women with no cervical infection at insertion:
– 1/201 in the antibiotic group and 1/200 in the placebo group were found to have PID (combining both groups PID 2/394, 0.05%)
– Unclear if women with STIs at insertion were treated

STI screening information is only available for 272 women (61%) of the total IUD prospective users

Very Low (continued on next page)
<table>
<thead>
<tr>
<th>Author, year, support</th>
<th>Objective</th>
<th>Study Design</th>
<th>Population</th>
<th>Outcome (and assessment)</th>
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<th>Results</th>
<th>Weaknesses</th>
<th>Quality</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fau`ñdes et al., 1998 [4], Population Council, MacArthur Foundation and UNDP/UNFPA/WHO/WB</td>
<td>To objectively evaluate how effective the use of sociodemographic factors, sexual behavior and signs and symptoms are in the identification of women at high risk for STIs among new contraceptive acceptors</td>
<td>Prospective study for 1 month; Brazil 1991–1992; Copper T 380A</td>
<td>407 prospective IUD users; 327 actually given IUD</td>
<td>PID: criteria for diagnosis not stated; removal due to PID</td>
<td>– Women with history of multiple partners, purulent cervical secretion, hyperemia and bleeding of the cervix at touch or pelvic pain during bimanual vaginal exam were given a clinical diagnosis of chlamydial infection or gonorrhea and were not given IUDs</td>
<td>Women with cervical infections at insertion:</td>
<td>– No information about PID diagnostic criteria</td>
<td>Very low</td>
</tr>
<tr>
<td>Morrison et al., 1999 [5], support not stated</td>
<td>To evaluate the use of risk assessment algorithms to predict STI and subsequent IUD-related complications among IUD candidates</td>
<td>Prospective study (4 months with a 1-month follow-up visit); Kenya 1994–1995; Copper T 380A</td>
<td>649 prospective IUD users, 615 in analysis (including 144 HIV-positive women and 471 HIV-negative women)</td>
<td>PID: defined by the United States Infectious Disease Society of Obstetrics and Gynecology</td>
<td>– Women with active PID, mucopurulent cervical discharge or high risk for STIs were excluded from the study</td>
<td>Women with cervical infections at 1 month:</td>
<td>– Do not know if women had chlamydial infection or gonorrhea at the time of insertion; available information is for the first month follow-up</td>
<td>Very low</td>
</tr>
</tbody>
</table>

either before or after insertion [9]; and in one, testing for STIs was done at the first month follow-up visit [5]. In that study, it was assumed that because study participants were at low risk of developing STIs, the majority of women who tested positive (32 women) most likely acquired infection before the IUD was inserted (Morrison CS, FHI, June 4, 2003). Follow-up from the time of IUD insertion ranged from 1 month to 2 years.

As shown in Table 1 and Fig. 1, the prevalence of STIs and incidence of PID were low in all studies, producing wide variability in the confidence intervals. Rates of diagnosed PID ranged from 0% to 5% among women with STIs at IUD insertion and 0% to 2% for those without STIs at IUD insertion. One of the studies, however, also identified a possible case in a woman with STI at insertion (the woman had lower abdominal pain but no fever); if this case is included, the upper boundary of PID among those with STI would be 10.5% [4]. With the exception of the study of Skjeldestad et al. [7] that observed no cases of PID in the study population, all of the studies observed a greater risk of PID among women with STIs (combining gonorrhea and chlamydial infection) at IUD insertion than women with no STI at insertion, with crude relative risks ranging from 1.63 to 46.35. The Sinei et al. [6] study, which had the largest sample size of the studies and a high follow-up rate, yielded a relative risk of 2.69 (95% CI 1.11–6.53). Fig. 2 shows results for chlamydial infection and gonorrhea separately and also suggests a trend of greater risk of PID with infection, although the number of cases was quite low, leading to imprecise estimates.

4. Discussion

There are theoretical concerns that, among women with current STIs, the process of inserting an IUD and perhaps the presence of the device may facilitate the ascendance of sexually transmitted organisms from the lower to the upper genital tract. This review found no studies that were able to examine appropriate case and comparison groups to assess this question and, correspondingly, to determine whether among women with STIs the insertion and use of an IUD increases the risk of PID. Of six studies that provided indirect evidence, current STI at the time of IUD insertion increased the risk of PID, albeit only two of the increases were statistically significant [4–9]. This finding is expected, as gonorrhea and chlamydial infection are the major etiologic agents for PID in women who do not use the IUD. We do not know, however, whether this increase in risk is the same or greater than the risk for PID among women who do not undergo IUD insertion. We do know that the absolute risk of PID remained low among women who had an STI when their IUD was inserted, in the range of 0–5% (or possibly as high as 11%, if the “possible” case of PID was included).

Unfortunately, the overall quality of the indirect studies was “very low.” The studies varied in the level of initial screening for STIs and their diagnostic criteria for PID, thereby possibly introducing selection and misclassification bias. Because of the limited number of women with STIs and PID in these studies, assessing possible confounders was not possible. The main limitation of this body of indirect evidence was the small number of cases, which is explained by the current practice of screening for STIs before inserting an IUD and the low prevalence of STIs and incidence of PID in populations desiring IUD insertion.

Ideally, we would like to have evidence from studies that examined the following question: among women with STIs, does insertion of an IUD increase the risk of PID compared with no IUD insertion? Ethically, this study would be difficult to conduct, as women could not be followed, untreated, to assess their risk of developing PID. There is some evidence, however, regarding the risk of PID among women with STIs in the general population who are not IUD users. A study of 129 women with both cervical chlamydial infection and gonorrhea who were treated with penicillin (which does not affect chlamydial infection) found that 9% (11/129) subsequently developed PID [12]. Another study that also treated women having both gonorrhea and chlamydial infection with penicillin found that 30% (6/20) developed PID [13]. Finally, a study of 19 women with gonorrhea found that nine of the women developed PID (four had adnexal tenderness), a median of 11 days after diagnosis of cervical infection [14]. When the risk of developing clinical PID attributable to an IUD was calculated using a model derived from existing studies, the risk was 0.15% when the prevalence of the gonorrhea and chlamydial infection in the population was approximately 10%, and there was screening for genital infections [15].

Some still argue that the use of an IUD increases the risk of PID regardless of whether STIs are present. Indeed, a meta-analysis that identified 36 papers published between 1974 and 1990 concluded that there was a positive association between IUD use and PID, even in subgroups examining IUDs that were not Dalkon Shields, and when separating symptomatic and asymptomatic PID [16]. There are several biases inherent in observational studies of this question [3,17], however, which cannot be resolved through meta-analysis. First, it is difficult to identify a valid control group with which to compare IUD users—the use of hormonal methods, barrier methods or sterilization decreases risk of PID, and women who do not use contraceptives may be very different from users with regard to age, parity and sexual risk behaviors. Second, a detection bias may be at work, as IUD users may be more likely to be diagnosed (or overdiagnosed) with PID. Third, many studies have not adequately controlled for potential confounding factors, primarily sexual behavior. Therefore, it remains difficult to answer this question with observational study designs.

Even without valid relative risks, absolute rates of PID among IUD users are reported to be low. In a 5-year follow-
up study in eight developing countries, the rate of acute PID among users of the copper IUD was 0.6 per 1000 woman-years [18]. In randomized controlled trials of prophylactic use of antibiotics at IUD insertion, PID among the placebo group with IUD insertion has been rare, even in populations with high background prevalence of STIs—1.11% in Nigeria at 30 days after insertion [19] and 1.9% in Kenya at 1 month after insertion [6].

In summary, the study findings consistently showed that women with STIs have a greater risk of PID than women with no STIs when an IUD is inserted, but the absolute risk of PID among women with STIs at the time of IUD insertion is low. We have no information to determine whether IUDs increase the risk of PID in women with STI at the time of IUD insertion is of the same magnitude. Therefore, whether IUDs increase the risk of PID in women with STIs who do not receive an IUD is of the same magnitude. We have no information to determine whether the risk of PID among women with STIs who do not receive an IUD is of the same magnitude. Therefore, whether IUDs increase the risk of PID in women with STI at the time of insertion is not known.

In 2003, the WHO Expert Working Group reviewed this evidence to evaluate current medical eligibility criteria for use of copper and levonorgestrel-releasing IUDs [20]. The Expert Working Group recommended that women with current purulent cervicitis or cervical chlamydial infection or gonorrhea should not have an IUD inserted (WHO Category 4), but women who already have an IUD in place and are found to have a cervical infection can generally continue use of the copper or levonorgestrel-releasing IUD (WHO Category 2). Women with other STIs (excluding women with hepatitis desiring the levonorgestrel-releasing IUD, because of concerns regarding hormonal effects on the course of disease) and women with vaginitis can generally initiate and continue use of copper or levonorgestrel-releasing IUDs (WHO Category 2).

Acknowledgments

This review was supported by resources from the World Health Organization, the US Centers for Disease Control and Prevention (CDC), US Agency for International Development (USAID) and the US National Institute of Child Health and Human Development (NICHD). We would also like to acknowledge the assistance of William Thomas, MLIS, Technical Information Specialist at CDC, in developing the literature search strategies.

The findings and conclusions in this report are those of the author(s) and do not necessarily represent the views of the funding agencies.

Appendix A. Study quality assessment

A.1. Individual study

Each study was given a rating of very low, low, intermediate or high based on the interval validity of the study. If the study was indirect, the quality of the individual study was lowered by one level. If the study was direct, the quality of evidence was kept the same. Similarly, if there was sparseness of the data, the quality of the individual study was lowered by one level.

A.2. Body of evidence

The quality of the body of evidence was the highest rating given to an individual study. If the results were inconsistent, the quality of the body of the evidence was lowered by one level. If results were consistent, then the quality of the body of the evidence was left at the original level. Similarly, if there was reporting bias (publication bias), then the quality of the body of evidence would be lowered by one level.

<table>
<thead>
<tr>
<th>Quality of evidence across studies for each main outcome</th>
<th>RCT</th>
<th>Observational studies</th>
</tr>
</thead>
<tbody>
<tr>
<td>No serious flaws in design or execution</td>
<td>High</td>
<td>Extremely strong association and validity</td>
</tr>
<tr>
<td>Serious flaws in design or execution</td>
<td>Intermediate</td>
<td>Strong, consistent association and no threats to validity</td>
</tr>
<tr>
<td>Very serious flaws in design or execution</td>
<td>Low</td>
<td>No serious flaws in study quality</td>
</tr>
<tr>
<td>Very serious flaws and at least one other serious threat to validity</td>
<td>Very low</td>
<td>Serious flaws in design and execution</td>
</tr>
</tbody>
</table>

Additional factors that lower study quality are as follows: important inconsistency of results; some uncertainty about directness; high probability of reporting bias; and sparseness of data. Major uncertainty about directness can lower the quality by two levels.

Additional factors that may increase quality of observational studies are as follows: all plausible residual confounding, if present, would reduce the observed effect, and evidence of a dose–response gradient.


References


