Effectiveness of female condom in preventing HIV and sexually transmitted infections: a systematic review protocol

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ABSTRACT

Introduction The HIV pandemic continues to evolve with young women being the most vulnerable group to acquire infection. The presence of sexually transmitted infections (STIs) further enhances HIV susceptibility and also leads to long-term complications such as infertility and cervical cancer. The female condom is a self-initiated method for STI and HIV prevention but there are controversies on its effects. We aim to assess the effectiveness, safety and acceptability of the use of female condoms for prevention of STI and HIV acquisition among women.

Methods and analysis We will search Cochrane Central Register of Controlled Trials, PubMed, EMBASE, Scopus, WHO International Clinical Trials Registry and reference lists of relevant publications for potentially eligible studies. We will screen search outputs, select eligible studies, extract data and assess risk of bias in duplicate; resolving discrepancies through discussion and consensus or arbitration. We will combine data from clinically homogenous studies in a fixed effect meta-analysis and assess the certainty of the evidence using the method for Grading of Recommendations Assessment, Development and Evaluation. We registered the planned systematic review with the International Prospective Register of Systematic Reviews (PROSPERO) in March 2018 and will finalise the search strategy in August 2018; conduct the searches and select eligible studies between August and October 2018; and collect data, conduct statistical analyses and prepare and submit the manuscript for consideration by a peer-reviewed journal between November 2018 and April 2019.

Ethics and dissemination We will use publicly available data; hence no formal ethical approval is required for this review. We will disseminate the findings of this review through conference presentations and publication in an open-access peer-reviewed journal. PROSPERO registration number CRD42018090710.

INTRODUCTION

The disease burden resulting from unsafe sex, including HIV infection and other sexually transmitted infections (STIs), has profoundly impacted low-income and middle-income regions, especially Sub-Saharan Africa. The HIV pandemic continues to evolve in both magnitude and diversity, with over 40 million infections worldwide, with young women aged 15–24 being 2.5 times more likely to be infected than young men. In many cases, STIs go undiagnosed and eventually lead to long-term complications such as infertility and cervical cancer. In addition, the presence of an STI enhances HIV susceptibility. Several interventions exist for the prevention of HIV and STIs such as the male condom. Although male condoms are effective in reducing HIV and STIs transmission, the subordinate status of women in many countries, especially in Sub-Saharan Africa, makes negotiating male condom use with partners especially difficult. Hence, women remain particularly vulnerable to HIV infection and other STIs like gonorrhoea, chlamydia, syphilis, human papilloma virus (HPV) and herpes simplex virus (HSV) infections. There is evidence that increasing the availability of multiple contraceptive methods for women is associated with increased contraceptive uptake, lower pregnancy rates and fewer STIs. Furthermore, the contraceptive needs and preferences of women have been found to change over the course of their reproductive life, and it...
is imperative that women have a wide variety of options available to encourage them to use their contraceptive of choice. There are several methods of contraception that exist for women such as female sterilisation, long-acting hormonal contraceptives, short-acting hormonal contraceptives, copper intrauterine devices, barrier methods and natural method. However, the female condom, which is a barrier method of contraception, is the only female-initiated contraceptive method that offers dual protection against both pregnancy and STIs. In fact, there is evidence suggesting that it may be as effective as the male condom though this conclusion has not been demonstrated.

Introduced over two decades ago, the female condom offers the possibility of an alternative to male condoms. Several types of materials can be used to make female condoms, including polyurethanes, synthetic nitrile rubber latex, natural rubber latex and silicon. Generally, the structure of the female condom consists of a sheath that lines the vagina and may extend to cover the external genitalia. At the closed end of the sheath, a flexible ring of foam sponge is inserted into the vagina to hold the female condom in place. These internal retention features also help to facilitate insertion of the female condom into the vagina. At the other open end of the sheath, there is a ring or frame that stays outside the vulva at the entrance to the vagina. This ring or frame prevents the sheath bunching up inside the vagina and also facilitates removal of the condom. Some female condoms such as the Phoenurse are prelubricated and others like the Cupid are scented. The first-generation female condom, available since 1993, was made out of polyurethane. However, it has been progressively replaced by newer female condoms, designed to lower unit cost and/or increase acceptability. Clinical studies evaluating the efficacy, safety and acceptability of these new designs are ongoing.

In comparison with the male condom, the female condom is said to offer additional coverage to both partners and is not weakened by the use of oil-based lubricants. Furthermore, no serious local side effects or allergies have been reported. However, this non-systemic contraceptive method is not without limitations. It is known to be relatively more expensive, with mechanical problems which could include breakage, slippage, invagination and misdirection among others. These limitations are increasingly being addressed by the designing and manufacture of newer forms of the female condom, with emphasis on proper and frequent use of existing ones.

Research has been undertaken to determine the feasibility of reusing the female condom. A consultation convened by the WHO in January 2002 addressed certain considerations regarding the reuse of female condom. They concluded that although the use of a new female condom during each act of sexual intercourse should be recommended, the female condom can be reused in couples not at risk of pregnancy, STIs or HIV infection, but with careful attention to a disinfection, washing, drying and relubrication procedure. Research on the structural integrity of the female condom shows that it is maintained after five uses. However, additional research on the effectiveness of female condom reuse is still ongoing with newer designs of the female condom.

There is evidence that condom use results in up to 80% reduction in the incidence of HIV. However, these estimates generally refer to the efficacy of the male condom. With the advent of newer forms of the female condom, many randomised controlled studies have examined the effectiveness and acceptability of female condoms in preventing HIV and other STIs. Additionally, stereotypes and strong opinions that tend to hamper the acceptance of female condoms exist. These in turn may hinder their correct and consistent use, an aspect that determines the effectiveness of this barrier method in preventing HIV. In this review, we seek to examine the evidence from both randomised and non-randomised trials, on the effect of female condom use on the incidence of HIV and other STIs among women. We also plan to explore the side effects and acceptability of female condoms.

**METHODS AND ANALYSIS**

**Patient and public involvement**

Patients were not involved in the design of this study. However, the development of the research question and outcome measures were informed by patient’s priorities, experience and preferences as reported in the literature supporting this review. The findings of this review will provide patients and policy-makers with the evidence on the efficacy and safety of existing and newer types of female condoms.

**Criteria for considering studies for this review**

We will include randomised and non-randomised trials that enrolled HIV negative and/or HIV positive women, engaged in heterosexual activity in any setting, with no clinical or laboratory-confirmed signs of STIs.

In addition, eligible trials would be those that compared the female condom to no treatment or other barrier methods for HIV prevention, for example, male condom, microbicides, diaphragm, vaginal sponges and cervical caps.

Finally, eligible studies need to report at least one of our primary or secondary outcomes of interest. Our primary outcomes for this review include acquisition of HIV (determined by a serological test) or STIs (including, but not limited to chlamydia, gonorrhoea, syphilis, HSV, trichomoniasis, candidiasis, lymphogranuloma venereum, HPV and bacterial vaginosis). Eligible studies need to determine STI status by microscopy and/or culture of urogenital specimens and vesicle fluid (when possible) for the causal agents. We will also consider nucleic acid amplification tests, if reported. Cytological pap testing used to determine human papillomavirus infection, and microscopy of Gram stained genital smear used to
detect bacterial vaginosis will be acceptable methods of determining status. We will also include studies in which STIs were diagnosed clinically, and subgroup analyse by method of diagnosis (clinical vs laboratory).

Our secondary outcomes will include acceptability and adverse events of female condom use. Measures of acceptability may include scales to grade acceptability and where possible, these will be standardised to allow for quantitative comparison across trials. If this is not possible, then we will provide a narrative synthesis. Adverse events may include difficulties in insertion and removal of the condom leading to inconsistent use, breakage and slippage of condom, decreased pleasure and penetration difficulties during intercourse, genital ulcerations during intercourse and any other adverse events reported in the trials.

**Search methods for identification of studies**

We will use keywords to build a comprehensive search strategy that will be used to search the Cochrane Central Register of Controlled Trials, PubMed, EMBASE and Scopus for publications indexed from 1980 to July 2018. We have chosen to limit our search to this timeline as it corresponds to the identification of the first case of HIV. We have provided the proposed search strategy for one database, PubMed, in **table 1**. We will also search the WHO International Clinical Trials Registry Platform for ongoing studies and the reference lists of included studies and related reviews for other relevant studies. We will include trial reports available in English or French.

**Study selection**

We will develop the search strategy and conduct the electronic searches with the help of an information specialist. The search output from the various databases will be combined and deduplicated using a reference management software (EndNote). Two authors will independently screen the titles and abstracts obtained from the electronic searches to create a pool of potentially eligible studies. Disagreement between the two authors will be resolved by discussion and consensus, and a third author will arbitrate if discussions fail. We will obtain the full articles of the potentially eligible studies which two authors will independently scrutinise for relevance using a standardised eligibility form with predefined inclusion criteria. The criteria for relevance will be based on the study design, interventions, participants and outcomes. If some of the information needed to classify the study is missing, we will attempt to contact the study authors for clarification. In the event where the authors do not have the missing information, or fail to respond, the study will be classified as ‘awaiting assessment’. Disagreements between the two authors will be resolved by discussion and consensus. Should the disagreement persist, a third author will arbitrate. Following the eligibility assessment, each study will be classified as included, excluded, ongoing or awaiting classification. A study that meets the design, intervention and participant criteria for which relevant outcomes are not yet available will be classified as ongoing (if the study is not yet completed) or awaiting classification (if already completed). We will prepare a table of the excluded studies, with reasons for exclusion. All four authors will take part in study selection.

**Data extraction**

Two authors will independently extract data using a standard data extraction form. Extracted information will include study details such as location and setting, study design, population size and attrition rate; intervention details such as time period for the intervention and length of follow-up; comparator details including the type of comparator, time period for the comparator and length of follow-up; and outcome details such as HIV and STI incidence (with types of laboratory tests used to confirm HIV and STI diagnosis), degree of compliance with female condom use, acceptability and adverse effects. Disagreement between the two authors will be resolved by discussion and consensus. Should the disagreement persist, a third author will arbitrate. Where information in the study report(s) is unclear or missing, we will contact the authors and request for the missing information. If the

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authors fail to provide the missing information, the study will be included in the review; however, the findings that are unavailable will not be synthesised with findings from other included studies addressing the relevant outcome.

Assessing risk of bias
Two authors will independently assess the risk of bias in each included trial using the Cochrane risk of bias tool.26 This will include information on the adequacy of the generation of the allocation sequence and allocation concealment (for assessment of the risk of selection bias), blinding care providers (for performance bias), blinding of outcome assessors (for detection bias), completeness of outcome data (for attrition bias), completeness of outcome reporting (for reporting bias). Given the nature of the interventions considered in this review, the study participants cannot be blinded. The risk of performance bias will therefore be assessed based on whether the care providers were aware of the intervention or not.

Data synthesis
We will use the Cochrane Review Manager for data analyses. We will express study results as risk ratios (for dichotomous variables such as HIV incidence) or mean differences (for continuous outcomes such as acceptability); with their 95% CIs. We will combine study results in a meta-analysis if included trials found are similar in terms of design, participants, interventions and outcomes.

We will assess heterogeneity between trial results by visually inspecting the forest plots to assess whether the CIs overlap, followed by a more formal test, that is, the $\chi^2$ test of homogeneity (with significance defined as an alpha level of 10%). We will also use the I² test to quantify the degree of heterogeneity.

In the absence of significant statistical heterogeneity, we will pool the study results using the Mantel-Haenszel fixed-effect method. If we detect significant heterogeneity and consider it clinically meaningful to combine the trials, we will use the random-effects meta-analysis. We will explore the cause of observed heterogeneity using subgroup analyses, with subgroups defined by study design (randomised vs non-randomised trials), HIV status (for outcomes other than HIV acquisition), method of STI diagnosis (clinical vs laboratory), type of comparison intervention, trial duration, degree of compliance with female condom use and sample size. When a significant statistical association is found, we shall calculate the absolute risk reduction (or increase) with the number needed to treat or number needed to harm, as appropriate. Data obtained from studies that are not similar enough to be meta-analysed will be combined using narrative synthesis. We will use a funnel plot to assess for publication bias if we have more than 10 included studies in a meta-analysis. Finally, we will assess the strength or certainty of the evidence using the Grading of Recommendations Assessment, Development and Evaluation approach which rates the certainty of evidence for each outcome by taking into consideration the risk of bias, directness of evidence, heterogeneity, precision and risk of publication bias.27

Timeline for the systematic review
We registered the planned systematic review with the International Prospective Register of Systematic Reviews (PROSPERO) in March 2018.28 We plan to finalise the search strategy in August 2018, conduct the searches and select eligible studies between August and October 2018; and collect data, conduct statistical analyses, and prepare and submit the manuscript for consideration by a peer-reviewed journal between November 2018 and April 2019.

Ethics and dissemination
We will use data that are readily available in the public domain, hence no formal ethical approval is required for this review. The findings of this review will be presented at relevant conferences and published in a peer-reviewed journal. This protocol has been written following the Preferred Reporting Items for Systematic Reviews and Meta-Analyses Protocols guidelines,29 and the findings of this review and any amendments will be reported according to the Preferred Reporting Items for Systematic Reviews and Meta-Analyses statement.30

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Contributors RKB conceived the study and wrote the first draft of the protocol. ABW, EJK and CSW provided content and methodological expertise. All authors read, amended and approved the final version of the protocol before submission. CSW is the guarantor for this review.

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