Feasibility and potential acceptability of three cervical barriers among vulnerable young women in Zimbabwe

Ariane van der Straten, Nuriye Sahin-Hodoglugil, Kate Clouse, Sibongile Mietwa, Mike Z Chirenje

Abstract

Background We explored the potential acceptability of three cervical barriers (CB) (Ortho All-Flex® diaphragm, SILCS® diaphragm, FemCap™ cervical cap) among sexually experienced Zimbabwean young women.

Methods Forty-five young women (aged 16–21 years) received an individual CB educational session. Participants were then randomly assigned to one of the three CBs in a 1:1:1 ratio, and practised insertion and removal of their device at the clinic. Next, participants were interviewed on their practice experiences, and their post-practice attitudes towards CB.

Results All 45 young women were willing and able to insert their assigned device. The majority reported “easy” insertion and removal and 93% “liked” the device they tried. All showed interest in participating in future CB studies: when asked which device they would like to try in the future, over half (58%) chose SILCS, regardless of the device they had tried. The majority felt comfortable touching their genitals to insert/remove the CB and most participants favoured methods’ attributes associated with female-control and non-interference with sex. Over half the participants said they would prefer to use a CB continuously compared to episodic use. Two-thirds of them expressed interest in CB for dual protection.

Conclusion The concept of CB, and initial insertion experience, were well accepted in this selected, small group of Zimbabwean young women. Evaluating CB in larger studies seems feasible in this population.

Keywords acceptability, cervical barrier method, contraception, HIV prevention, young women


Introduction

Young women are the most vulnerable population for heterosexual acquisition of HIV. Adolescents are more susceptible than adult women, likely because an immature genital tract increases their susceptibility to physical trauma and sexually transmitted infections (STIs). Furthermore, gender inequalities in sexual relationships often leave young women unable to negotiate condom use. In Zimbabwe, 25% of females aged 15–24 years are infected with HIV compared to 11% of males in that age group. Additionally, with 40% of Zimbabwean women aged 15–49 years being non-current users of modern family planning methods, there remains a significant unmet need for contraception, particularly among young sexually active women (63% non-current users among 15–19-year-olds), as indicated by estimates that up to 60,000–80,000 illegal and unsafe abortions occur each year.

Male condoms are the only currently available method known to provide dual protection by preventing pregnancy and STIs including HIV; although not proven in clinical trials, female condoms are believed to provide similar protection. While female condoms are woman-initiated, they still require male partner co-operation, as they are noticeable during sex. Cervical barriers (CB), including the diaphragm, are also woman-initiated methods, and act by protecting the cervix from exposure to ejaculate. The diaphragm is one of the oldest contraceptives, although it has lost much of its popularity with the advent of hormonal contraceptives. CB might be of interest to women who prefer a non-hormonal method or who have experienced side effects with hormonal methods. CB have the potential to provide dual protection because they cover the cervix, a ‘hot spot’ for some STIs and HIV, leading to a renewed interest in these methods and ongoing research on the diaphragm’s possible role for HIV or STI prevention. This effort continues, despite disappointing results from a recently completed trial of the diaphragm used with a lubricant gel, the MIRA trial, which was unable to demonstrate a protective effect of the intervention against HIV or cervical STIs, and is over and above that of male condoms. Nevertheless, it is still biologically plausible that the diaphragm or other CB can provide partial protection from cervical infections, and function as a reusable delivery mechanism for, or enhance the effectiveness of, a microbicide gel. Indeed, as CB retain gel close to the cervix, they may potentially prolong the duration of action of a microbicide in situ, by slowing the rapid leakage of gel to the lower vagina and the introitus during sexual intercourse.

Recently, new CB devices with improved designs have
been developed, which do not require fitting, come in one or only a few sizes, and harbour special features ... a contraceptive gel. Gel can be loaded on the cervical and vaginal sides of the device prior to insertion.

(a) Ortho All-Flex® (Ortho-McNeil, Inc., Titusville, NJ, USA) is a peach, dome-shaped, latex rubber cup with a flexible rim. Worldwide, it is the most available CB. It comes in nine sizes (ranging from 55 to 95 mm, in 5 mm increments) and must be fitted by a clinician. It is US Food and Drug Administration (FDA) approved for contraception when used with a contraceptive gel, which can be loaded onto the cervical side (within the cup) prior to insertion.

(b) SILCS® (PATH, Seattle, WA, USA) is a purple, dome-shaped silicone diaphragm. It has an anatomically shaped, contoured design for easy placement and removal. It is a “one-size-fits-most” device and requires no fitting. It is not yet FDA approved, but is being evaluated in a Phase II/III contraceptive effectiveness trial.25 Gel can be loaded on the cervical and vaginal sides of the device prior to insertion.

(c) FemCap™ (FemCap Inc., Del Mar, CA, USA) is a dome-shaped, white silicone device with a wide brim. It is designed to conform to the shape of the vaginal fornices and cervix. The brim is designed to hold gel and trap sperm. There is a removal strap over the dome. It comes in three sizes, as determined by obstetric history, so no clinician fitting is required. It is FDA approved for contraception when used with a contraceptive gel. Gel can be loaded on the cervical and vaginal sides of the device prior to insertion.

Figure 1 The three cervical barriers (CB) employed in the study. (a) Ortho All-Flex® (Ortho-McNeil, Inc., Titusville, NJ, USA) is a peach, dome-shaped, latex rubber cup with a flexible rim. (b) SILCS® (PATH, Seattle, WA, USA) is a purple, dome-shaped silicone diaphragm. (c) FemCap™ (FemCap Inc., Del Mar, CA, USA) is a dome-shaped, white silicone device with a wide brim.

Methods

Study design and participants

This was a mixed methods exploratory study with two interrelated components: the first qualitative and the second clinical. For the first component, focus group discussions (FGDs) were conducted with young women and with adult women (who were mothers and aunts of the former) on sociocultural issues around sex, reproductive health, knowledge about HIV prevention, and to discuss CB methods. During the FGDs, participants were presented with three different CB and each was demonstrated using a pelvic model. Attitudes about, and reactions to, the devices were collected. Qualitative data from these FGDs are presented elsewhere.36

For the second (i.e. clinical) study component, young women who participated in the FGDs were invited back 2 weeks later, to practise inserting and removing one of the three randomly assigned CB in the clinic under the supervision of a clinician. They were then interviewed on their practice experiences, along with their post-practice attitudes towards CB. The results of this second component are reported here.

This study was conducted concurrently with the MIRA trial18, between May and September 2006; a total of 93 young women aged 16–21 years were approached and pre-screened by outreach workers in schools, youth centres, sports clubs, youth-friendly clinics, and market places in the city of Chitungwiza, near Harare. Fifty-one (55%) young women came to the study site and were re-screened for eligibility. Of those, 47 were available, and completed a FG; Eligibility criteria included being aged 16–21 years, ever having had vaginal sex, living in the greater Harare region, being able to read, write and speak English or Shona (the local language), and willing and able to give written informed consent. Forty-five (96%) young women returned for one follow-up clinical visit 2 weeks after their FGD and these individuals constitute our analytical sample. Two women withdrew prior to the follow-up visit (one moved to Botswana and one started working full time).

Study procedures

A female study clinician presented the three study CB and provided each participant with a standardised educational overview: briefly, participants were told that CB can prevent pregnancy when used with a contraceptive gel, that they have the potential to provide some protection against STIs, and that the diaphragm was currently being evaluated for HIV/STI prevention. The three devices were handed to the participants, who were encouraged to examine their shape and feel, and were shown how they are worn inside the vagina, as demonstrated on a translucent pelvic model. The clinician then pointed to each CB (in random order) and explained its fitting and sizing requirements (Figure 1).

After the educational session, participants were given the choice to opt out of the practice session but none of them declined. All 45 participants were randomly assigned to one of the three CB, after opening sequentially numbered, sealed, opaque envelopes: 14 were assigned to SILCS, 15 to Ortho All-Flex and 16 to FemCap. The participants assigned to Ortho All-Flex were fitted for the device by the clinician (median size 70 mm, range 60–75 mm). For those assigned to FemCap, seven used a size 22 mm and nine a size 26 mm based on their obstetric history. Each participant was given K-Y® Jelly Personal Lubricant (Personal Products Company, Skillman, NJ, USA) to spread onto the rim of her device to facilitate insertion, and practised insertion and removal under the guidance of the study clinician. The clinician also conducted an assessment of the insertion and removal process and placement of the device in situ, and reported how many attempts were needed before correct insertion of the device.

Measures and analysis

Prior to the FGD, all participants completed a brief self-administered demographic form. At the follow-up visit, each participant completed an additional background and demographic questionnaire administered in private by a trained female interviewer. For the CB practice session, the clinician collected device fitting information, insertion and removal data on a study form. Finally, a trained interviewer conducted a follow-up interview using a structured interview guide with close- and open-ended questions to assess each participant’s experiences with insertion and removal of their device, along with their post-practice
Table 1 Characteristics of the study participants (n = 45)

<table>
<thead>
<tr>
<th>Characteristic</th>
<th>n</th>
<th>%</th>
</tr>
</thead>
<tbody>
<tr>
<td>Median age (range)</td>
<td>19</td>
<td>15–21</td>
</tr>
<tr>
<td>Median years of education (range)</td>
<td>11</td>
<td>4–13</td>
</tr>
<tr>
<td>Median total lifetime sexual partners (range)</td>
<td>1</td>
<td>1–10</td>
</tr>
<tr>
<td>Median age at first sex (range)</td>
<td>17</td>
<td>13–20</td>
</tr>
<tr>
<td>Lifetime frequency of sex</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Rarely or occasionally</td>
<td>24</td>
<td>53.3</td>
</tr>
<tr>
<td>Regularly</td>
<td>21</td>
<td>46.7</td>
</tr>
<tr>
<td>Has a husband or regular partner</td>
<td>33</td>
<td>73.3</td>
</tr>
<tr>
<td>Marital status</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Married</td>
<td>24</td>
<td>53.3</td>
</tr>
<tr>
<td>Single</td>
<td>13</td>
<td>28.9</td>
</tr>
<tr>
<td>Divorced or widowed</td>
<td>8</td>
<td>13.3</td>
</tr>
<tr>
<td>Has children</td>
<td>26</td>
<td>57.8</td>
</tr>
<tr>
<td>Earned income in the past year</td>
<td>17</td>
<td>37.8</td>
</tr>
<tr>
<td>‘Very worried’ about getting pregnant</td>
<td>32</td>
<td>71.1</td>
</tr>
<tr>
<td>‘Very worried’ about getting HIV</td>
<td>39</td>
<td>84.4</td>
</tr>
<tr>
<td>Intravaginally inserted products&lt;sup&gt;a&lt;/sup&gt; in past year</td>
<td>30</td>
<td>66.7</td>
</tr>
<tr>
<td>Washes intravaginally with finger&lt;sup&gt;b&lt;/sup&gt;</td>
<td>38</td>
<td>84.5</td>
</tr>
<tr>
<td>Ever touched her cervix</td>
<td>18</td>
<td>40.0</td>
</tr>
<tr>
<td>Ever heard of the diaphragm</td>
<td>13</td>
<td>28.9</td>
</tr>
<tr>
<td>Ever used the diaphragm</td>
<td>0</td>
<td>0.0</td>
</tr>
<tr>
<td>Current contraceptive method</td>
<td></td>
<td></td>
</tr>
<tr>
<td>No method</td>
<td>13</td>
<td>28.9</td>
</tr>
<tr>
<td>Pills</td>
<td>16</td>
<td>35.6</td>
</tr>
<tr>
<td>Male condoms</td>
<td>8</td>
<td>17.8</td>
</tr>
<tr>
<td>Injectables or Norplant&lt;sup&gt;c&lt;/sup&gt;</td>
<td>5</td>
<td>11.1</td>
</tr>
<tr>
<td>Multiple methods (hormonal + condoms)</td>
<td>2</td>
<td>4.4</td>
</tr>
<tr>
<td>Rhythm</td>
<td>1</td>
<td>2.2</td>
</tr>
<tr>
<td>Frequency of condom use during sex</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Every time</td>
<td>11</td>
<td>24.4</td>
</tr>
<tr>
<td>Sometimes</td>
<td>22</td>
<td>48.9</td>
</tr>
<tr>
<td>Never</td>
<td>12</td>
<td>26.7</td>
</tr>
</tbody>
</table>

<sup>a</sup>Products reported were: tampons, cotton wool, cloth, female condoms, herbs, other.

<sup>b</sup>Reported washing regularly (n = 31) or occasionally (n = 7) with water (n = 31) or water and soap (n = 7).

Feasibility and acceptability of cervical barriers

It is important for you to have a method which you can insert ahead of time before sex

It is important for you to have a disease prevention method that you can decide when to use

You would prefer to use a method which prevents HIV infection but allows you to get pregnant

In your opinion, men in this area will not be happy about using a gel if it makes the vagina wet

You would not use an HIV prevention method if it does not give “skin to skin contact” feeling

You would not be able to use a cervical barrier if you need to have a pelvic exam before getting it

You are concerned with having to leave the cervical barrier inside your vagina for at least six hours after sex

In your opinion, young women in general will not be interested in using cervical barriers

You are afraid that the cervical barrier may get lost inside your vagina

It feels awkward to touch your genitals to insert a cervical barrier

You will not have a private place to store the cervical barrier when you are not using it

It would be difficult for you to find a place to insert and remove a cervical barrier privately

It is difficult to learn how to insert cervical barriers

You think that cervical barriers are messy

Figure 2 Study respondents’ attitudes towards using a cervical barrier (n = 45). The figure shows the percentage of study respondents who “agreed” or “strongly agreed” with the statements in a 15-item cervical barrier attitude questionnaire.
Cervical barrier (CB) groups [% (n)]

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Total (n = 45)</th>
<th>SILCS (n = 14)</th>
<th>Ortho All-Flex (n = 15)</th>
<th>FemCap (n = 16)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Number of attempts for successful device insertion</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>First attempt</td>
<td>71.1 (32)</td>
<td>50.0 (7)</td>
<td>80.0 (12)</td>
<td>81.3 (13)</td>
</tr>
<tr>
<td>Second attempt</td>
<td>22.2 (10)</td>
<td>42.9 (6)</td>
<td>20.0 (3)</td>
<td>6.3 (1)</td>
</tr>
<tr>
<td>Third to fifth attempt</td>
<td>6.6 (3)</td>
<td>7.1 (1)</td>
<td>6.0</td>
<td>12.6 (2)</td>
</tr>
<tr>
<td>Perceived ease of the insertion and removal process</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Insertion was</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Easy</td>
<td>73.3 (33)</td>
<td>78.6 (11)</td>
<td>73.3 (11)</td>
<td>68.8 (11)</td>
</tr>
<tr>
<td>Somewhat difficult</td>
<td>28.7 (12)</td>
<td>21.4 (3)</td>
<td>26.7 (4)</td>
<td>31.3 (5)</td>
</tr>
<tr>
<td>Very difficult</td>
<td>0.0</td>
<td>0.0</td>
<td>0.0</td>
<td>0.0</td>
</tr>
<tr>
<td>Removal was</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Easy</td>
<td>84.4 (38)</td>
<td>100.0 (14)</td>
<td>86.7 (13)</td>
<td>68.8 (11)</td>
</tr>
<tr>
<td>Somewhat difficult</td>
<td>15.6 (7)</td>
<td>0.0</td>
<td>13.3 (2)</td>
<td>31.3 (5)</td>
</tr>
<tr>
<td>Very difficult</td>
<td>0.0</td>
<td>0.0</td>
<td>0.0</td>
<td>0.0</td>
</tr>
<tr>
<td>Liked the device she tried</td>
<td>93.3 (42)</td>
<td>100.0 (14)</td>
<td>93.3 (14)</td>
<td>87.5 (14)</td>
</tr>
</tbody>
</table>

Clinical practice of CB insertion and removal

All the participants practised CB insertion and removal. Overall, 71% were able to correctly insert their assigned device at their first attempt, although more in the SILCS group required a second attempt (Table 2). All the participants but one (assigned to FemCap, and unable to remove her device herself) removed their device properly. The majority inserted (n = 31) and removed (n = 30) their device while squatting; the next most common position was with one foot on a chair or stool (data not shown). Overall, the majority thought that insertion or removal of their device was easy (73% and 84%, respectively). All those in the SILCS group said device removal was “easy”, while a few participants in the other two groups reported some difficulty with removal. This was highlighted during participants’ narratives, with four women in the FemCap group and one in the Ortho All-Flex group, reporting problems with insertion or removal of their device, because it was slippery, caused discomfort or could not be dislodged.

Staff specifically probed about participants’ comfort level touching their genitals to insert the CB, and the majority explained they were comfortable. Several women even referred to the familiar behavior of intravaginal washing during bathing. “[I] did not feel any different from when I put my fingers during bathing. I did not feel pain or discomfort” (20 years old, divorced, Ortho All-Flex group). Women who reported intravaginal washing seemed more likely to find that insertion was easy (79% vs 43%). Conversely, in narratives about insertion experience, five women mentioned some discomfort or pain with touching their genitals, and six (13%) agreed with the attitudinal statement: “It feels awkward to touch your genitals to insert a cervical barrier” (Figure 2).

Device and use preferences, and attitudes towards using a CB

Some 93% of participants said they liked the device they tried (Table 2) and all participants said they were interested in participating in a CB study in the future. When asked which device they would like to try in the future, over half in the SILCS group and about a third in the Ortho All-Flex and FemCap groups chose the device they had tried. Over half the participants in the Ortho All-Flex and FemCap groups also chose SILCS (Table 2). Reasons given for this choice included finding SILCS (from the practice or from its appearance) easy to insert and remove due to its contoured design and finger groove, or finding it aesthetically pleasing.

Over half (56%) the participants said they would prefer to use a CB continuously (removing it only once a day for washing) compared to episodic use (during sex only). Those who preferred continuous use mentioned several reasons: (a) they liked to “be prepared” if sex happens (“Wearing it all the time means you are always ready when your husband comes and he wants sex” (21 years old, married, FemCap group)); (b) they wanted to be “protected” at all times (“Because sometimes you do not expect to have sex with someone, so that way you prevent many things” (16 years old, single, FemCap group)); (c) they found continuous use easier/more convenient (“It is easier that way. Wearing it and then taking it off is not convenient” (19 years old, married, Ortho All-Flex group)) and (d) they could wear it “quietly” without their partner’s
knowledge “[I can prevent infection and pregnancy, and if my husband does not want I can use it until I decide to get pregnant. I liked the fact that my husband cannot tell it’s there if he does not like it. Also it does not hurt me if I wear it all the time” (21 years old, married, SILCS group)]. Conversely, those who preferred episodic use invoked disliking “wearing it all the time” or concerns about the effect on the body of keeping the CB for a long time in situ “[I would also want to find out how my body will tolerate/take it before I can use it continuously” (17 years old, married, FemCap group)]. They also mentioned not being used to the device, worries about possible discomfort or that it may move out of place. Finally, many just said they liked to use the device only when needed “[Because it’s only when you are having sex that you are at risk of infection” (20 years old, married, Ortho All-Flex group)]. One participant mentioned that the device would “last longer” if it is worn episodically.

Participants only experienced using gel on the rim of their devices to ease insertion. They were asked their hypothetical preference regarding use of gel with CB after it was explained to them that to afford maximum protection CB are normally used in conjunction with a gel (i.e. a spermicide for contraception), though the precise mode of application varies between device manufacturers. As shown in Table 2, the majority (82%) said they could use gel with a CB, even if it required using an applicator to insert gel in the vagina having it all the time.

Overall, the participants’ attitudes towards CB were favourable (Figure 2). Most participants favoured attributes associated with female control of the methods (such as providing the woman with decision-making power and allowing use without the partner noticing) as well as attributes associated with non-interference with sex (insertion ahead of time, and – to a lesser extent – desire for a method that allows skin-to-skin contact). Of note, over half the participants agreed with the statements “men in this area will not be happy about using a gel if it makes the vagina wet” and “you would prefer to use a method which prevents HIV infection but allows you to get pregnant”. Other attributes of CB that may represent a challenge for their acceptance and use included the requirement to have a pelvic examination prior to use (for the Ortho All-Flex diaphragm only), the current recommendation to leave the CB for at least 6 hours postcoitally before removal, and concerns that the device may get lost inside the vagina; these were perceived as unfavourable by a fifth to a quarter of the participants. Fewer than 7% of the participants had concerns about privacy for insertion or storage, or learning the skills required to use CB. Only one participant agreed with the statement that CB were messy.

**Dual protection**

As shown in Table 1, just over 50% (n = 23) of the participants currently used effective hormonal contraception, while almost a third used no method. Only a quarter of the women reported using condoms every time they had sex. This highlighted substantial vulnerability for both pregnancy and HIV acquisition. Indeed, over two-thirds of the women said they were “very worried” about getting pregnant, and even more (84%) were “very worried” about getting infected with HIV. When asked what would be their main reason for using a CB, the majority (67%) said it was to protect both from pregnancy and disease, while almost a third said it would be mainly for disease prevention and two participants said for pregnancy prevention only (Table 2).

**Discussion**

The main objectives of this study were to assess the feasibility of conducting a larger CB study among vulnerable young women for contraception and/or for disease prevention (when used in combination with a microbicide) and to identify a promising candidate among three existing CB devices for such future study. The concept of CB, and initial insertion experience, were well accepted in this selected, small group of diaphragm-naïve Zimbabwean young women. Further evaluation of these methods clearly appears feasible in this setting: all the participants were willing to try their assigned CB at the clinic, most liked their device and were comfortable with insertion and removal, finding the process easy. All said they were interested in participating in a future CB study, with just over half in the SILCS group and more than a third in each of the other two CB groups expressing interest in trying their assigned device again. A majority of women in all three groups said they wanted to try SILCS in the future. A (hypothetical) preference for SILCS was also revealed in the FGDs that were conducted prior to the device practice visit. This preference may have been carried over from the FGDs, although not all FGDs had a majority of participants preferring SICLS. In another acceptability study among sexually active monogamous couples, SILCS was preferred over Ortho All-Flex. It was noteworthy that only half of all SILCS users in the present study were able to successfully insert the device on the first attempt. Nevertheless, over three-quarters reported that SILCS insertion was “easy” and the participants’ narratives did not reveal any information on this issue. Still, this matter is of concern and should be investigated further, as outside of a clinical study, those who fail at the first attempt may not be sufficiently motivated to try again. A previous study of the FemCap also noted difficulties with removal of the device, despite the removal strap, as was reported by those who tried this device here. A large proportion of the study participants had experience with intravaginal finger or product insertion, and intravaginal washing experience was associated with finding the CB insertion process easy. Other populations where intravaginal practice is not so prevalent may find the insertion process of CB more awkward and daunting.

While the manufacturer of the Ortho All-Flex diaphragm recommends application of gel only to the cervical side, other diaphragm variants and newer CB allow for the delivery of gel on both the cervical and vaginal side in one step, a feature that may be particularly relevant if CB are used with microbicides for disease prevention. Participants were comfortable with the idea of using gel on the cervical and vaginal side of the CB, even if this required use of an applicator. However, this question was asked hypothetically, as volunteers had only used a limited amount of gel on the rim of their devices to ease insertion. In this study we chose to focus our assessment of the three devices separately from gel to avoid confounding the acceptability of one with the other. Ease of gel application with each device, insertion and removal of the devices when loaded with gel, and willingness to use gel, especially when having sex, will have to be empirically assessed with the use of a combination product. While here, more than half the participants thought men wouldn’t like to have sex if gel made the vagina “wet”. In user-based microbicide and diaphragm studies, gel in moderate amount was generally well accepted and liked by women and their male partners.

Most participants favoured CB methods’ attributes associated with female-control and non-interference with
Several limitations to this study. First, the sample size was small. While our findings would have been more robust if all subjects had tried all three devices, funding limitations precluded multiple visits, and we were concerned that participants would be physically uncomfortable if we asked them to insert three different devices in one single session. Also, post-practice evaluations were conducted at the clinic by study staff; this may have led to socially desirable responses and overly positive reports about the devices tried.

The potential of CB as a dual-purpose method was important for most participants. However, simultaneously, the majority of participants agreed they wanted a HIV-prevention method that also allowed conception. This was a sample of young women, many were nulliparous, and few were likely to have achieved their desired family size. Clearly both options are needed for women, as some will want dual protection while others will only want to be protected from disease. CB do not offer a disease prevention option only, but may satisfy unmet needs for dual-protection methods, when combined with a contraceptive gel that is also microbicidal. For those individuals wanting to conceive, other disease prevention methods should be developed, as condoms are the only proven methods currently available, and they will prevent both pregnancy and disease.

In summary, based on the present results, it should be feasible to conduct a user acceptability study of CB among vulnerable young women in Zimbabwe, to explore the dual-purpose potential of CB by assessing them in combination with a contraceptive and potential microbicidal gel agent. In view of the diminished power in sexual relationships experienced by adolescent women in Sub-Saharan Africa, dual-purpose women-initiated methods could be particularly relevant to this group.

**Statements on funding and competing interests**

**Funding** The study was funded by the University of California San Francisco (UCSF) AIDS Research Institute (ARI) Pilot Awards Program.

**Competing interests** None identified. Most of the work for this study was conducted while Dr van der Straten, Dr Sahin-Hodoglugil and Ms Clouse were based at the Department of OBGYN and Reproductive Sciences, University of California, San Francisco, CA, USA.

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Feasibility and acceptability of cervical barriers/Fiction review


31 Hardy E, de Padua KS, Jimenez AL, Zaneveld LJ. Women’s preferences for vaginal antimicrobial contraceptive. Il. Preferred characteristics according to women’s age and socioeconomic status. Contraception 1998; 58: 239–244.


43 Montgomery CM. The role of partnership dynamics in determining the acceptability of condoms and microbicides. AIDS Care 2008; 20: 733–740.


This is a love story that is narrated as if it takes place spanning a day from morning till nightfall. Yet the actual story is spread out over months. The narrative gives the reader glimpses into the lives of the characters, starting with a scene in the morning and ending with nightfall.

The main characters, Laura and Ben, meet after a long time apart in a chance meeting at the hospital where Ben works. He is a genitourinary medicine (GUM) doctor while Laura has returned home to care for her ailing mother, Professor Jellico, an erstwhile expert virologist. Ben’s brother, Bobby, is a young gay man with Down’s syndrome. Ben’s wife, Chloe, Aerosoles on the fringe of the story yet seems to affect Ben, Bobby and Laura in different ways.

The storyline flows well and is an easy, light read. The passage of time from morning to nightfall wasn’t obvious to me till I read the interview with the author at the end of the book, which is very interesting indeed. The story covers important themes relating to sexual and reproductive health care. Some of the more important are the attitudes to sex and sexuality in people with learning disabilities and their right to sexual expression. At the other end of the spectrum the book looks at the ageing Professor Jellico and the impact of the physical effects of ageing on her quality of life – this in turn affects Laura who cares for her – thus illustrating children who care for their parents and how it can affect their lives, privacy, sexuality and relationships. Seemingly they are free to do as they please but cannot do the position they find themselves in.

I found the book easy to read with simple themes and a believable story line. The author gave Ben the profession of a GUM doctor but I felt disappointed that this aspect of his role did not receive the weighting it could have. Perhaps the themes of homosexuality, HIV and GUM are intertwined in the story for the purpose of setting the platform for Ben’s and Bobby’s characters. In all, an enjoyable read!