Acceptability of COL-1492, a vaginal gel, among sex workers in one Asian and three African cities

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Objectives: To evaluate the acceptability of COL-1492, a vaginal gel containing 52.5 mg nonoxynol-9, in an HIV prevention trial.

Methods: Sex workers participating in a phase II/III triple blind, randomised trial in Benin, Côte d’Ivoire, South Africa, and Thailand were interviewed on the gel’s acceptability at monthly scheduled clinic visits. Saver sex counselling, male condoms, and study gels were given at each monthly visit; a gynaecological examination and HIV test were performed. Phase III interviews considered the participants’ appreciation of the gel. On the first, second, and fifth follow up visits, the study volunteers completed more extensive questionnaires.

Results: Responses were similar between treatment arms. Women indicated not liking their gel in 1.8% of the visits; 98.1% of the women found the gel easy to apply; 30.1% said that it affected sexual intercourse. These effects were mostly improvements (92.6%) by facilitating intercourse (73.6%). Intercourse was more often affected in women reporting painful sexual intercourse [OR: 2.59 (95% CI 1.63 to 4.12)] and in older women. The latter effect differed among centres. Conclusion: Most participants found their assigned gel acceptable and the vast majority of reported effects on intercourse were favourable. The type of gel had no significant impact on the findings.

While male condoms provide high levels of protection against HIV and other sexually transmitted infections (STIs), negotiating their use can be difficult for women; hence an urgent need for female controlled methods for HIV prevention. Most research in this area has studied the potential effect of vaginal gels. To be effective in real life these gels must be acceptable. Evidence of acceptability in long term use was therefore collected in a randomised blinded phase II/III trial assessing the effect on HIV transmission of a nonoxynol-9 containing gel, COL-1492, compared with a placebo gel. This paper reports acceptability findings in phase III.

METHODS

Data came from the phase III portion of a phase II/III multicentre, placebo controlled, triple blind study of COL-1492 (Advantage S, Columbia Laboratories, New York, NY, USA), a marketed vaginal spermicide containing 52.5 mg nonoxynol-9. The placebo was Replens (polyacarbophil, Columbia Laboratories, Paris, France), a marketed vaginal moisturiser. Both gels were similar, except COL-1492 contained 3.5% nonoxynol-9 and less carboram. They were packaged in identical single use, disposable applicators, delivering 1.5 ml. Study participants were healthy, HIV-1 negative female sex workers in Abidjan, Côte d’Ivoire; Cotonou, Benin; Durban, South Africa; and Hat Yai, Thailand. The study was approved by all relevant ethics review committees.

Phase III was initiated in August 1997. At monthly scheduled visits women received safer sex counselling, male condoms, and prefilled applicators. They had a gynaecological examination, HIV and STI tests, and free treatment for any curable STI. At each visit, participants answered standardised questions regarding acceptability of their assigned product. They rated the gel (very unpleasant, unpleasant, acceptable, pleasant) and indicated whether their (paying) clients and (non-paying) partners complained (never, rarely, often, always). On the first, second, and fifth follow up visits women answered a more detailed questionnaire. Answers were dichotomised and explained by treatment group or other covariates, correcting for centre, using marginal logistic regression with an independent working correlation matrix. Covariate selection followed a two step procedure. First backward elimination was performed in each centre separately, with p value for removal 5%. To facilitate direct comparisons across centres, a common GEE model retained all covariates significant in at least three centres.

The study ended in June 2000 with a significantly higher HIV incidence of 14.7 per 100 person years among COL-1492 users v 10.3 per 100 person years among placebo users.

RESULTS

In all, 764 women were randomised in centres which continued into phase III; 97 had no follow up data and nine in Durban dropped out (three) or seroconverted (six) before the start of phase III, leaving 658 women in this analysis (table 1). Women lost to follow up were significantly younger (with a shorter history as sex worker). In phase III, participants reported 469 439 vaginal coital acts with clients, of which 70.9% involved use of a condom plus study gel, 17.4% condom only, 9.0% gel only, and 2.7% neither gel nor condom. Participants reported 61 180 vaginal coital acts with non-paying partners, 38.2% with condom plus study gel, 6.8% condom only, 32.0% gel only, and 23.0% neither gel nor condom. In total, 1694 visits were considered (table 1). Acceptability answers were similar between treatment arms except for the proportion of visits at which women reported informing their partners of their gel. As this p value still exceeded 0.83%, the Bonferroni corrected boundary, treatments were pooled in further analyses (table 2).

The percentage of visits where women reported not liking the gel differed significantly between centres (exact p value<0.01), with the highest percentage in Abidjan and the lowest in Durban (table 2).

Approximately one third of responses indicated that the assigned gel affected sexual intercourse favourably (table 2), mostly because it involved “less pain during intercourse, good lubrication, and facilitated intercourse,” except in Durban, where this accounted for just 13.4% of the answers.
(table 2). When asked what one liked most about the gel, 40–50% of answers involved good lubrication, less pain, and facilitated intercourse. Again, Durban differed, with approximately half of the responses indicating that prevention of STIs, including HIV and/or illness, was what they liked most.

In a model explaining any reported effect on sexual intercourse by age at entry, painful sexual intercourse, calendar time, and the method of collecting compliance data, we found a significant interaction with centre for all covariates except for painful sexual intercourse. The adjusted odds of a perceived effect on sexual intercourse was higher for older women in Abidjan (OR = 1.06 per yearly increase in age, 95% CI 1.01 to 1.11) and in Durban (OR = 1.07, 95% CI 1.02 to 1.12). Furthermore, we saw a significantly higher reported effect on intercourse when women reported painful intercourse in Cotonou (OR = 3.55, 95% CI 1.55 to 8.13), Durban (OR = 1.90, 95% CI 1.00 to 3.60), Hat Yai (OR = 4.82, 95% CI 1.16 to 20.02), but not in Abidjan (OR = 1.96, 95% CI 0.15 to 25.51). Reports of effects on intercourse increased significantly with the study in Durban (p = 0.002), but not in the other centres.

Participants reported in 38.8% (41.9%) of the visits would continue their use if available after the trial. This is consistent with the conclusion drawn by Pool et al.1 that women may use any particular product if the choice is limited. Since this was a phase III trial, there was no other formulation available for comparison. As in other trials,18 improved intercourse, mainly through increased lubrication, was a main reason for liking the product. This suggests that use will increase with increased sexual comfort. This is important since the acceptability of gels is often questioned in cultures where a dry vagina is the norm for sexual intercourse—for example, in Durban.7

In Durban, half of the time HIV/STI prevention was named as the best liked feature of the gel, despite monthly intensive safer sex counselling and regular reminders of study information. Similarly, Coggins et al.8 found that 40% of women stated using a product to prevent STIs and another 44% for dual protection (against both STIs and pregnancy).

In our trial, participants reported informing their sex partners about gel use in approximately 40% of visits, less often than in a multinational preference study,9 where 75% of women informed their regular partners. This difference may stem from a different study population, women from the general population versus sex workers.

We found substantially fewer acts with reported gel and condom use with partners than with clients. On the other hand, the proportion of acts with gel alone was higher among partners. This points to the importance of a female controlled method for sex with (non-paying) partners which would also apply to the general population where women often risk infection as a result of their partner’s behaviour.

As in all behavioural research, our results need cautious interpretation because women may wish to please, especially in direct interviews.

Ninety seven women did not return for any follow up. We ignore their reasons, which may have been product related. Thus, the reported results may overestimate the true

<table>
<thead>
<tr>
<th>Table 1</th>
<th>Number of women attending the first, second, and fifth follow up visits during phase III</th>
</tr>
</thead>
<tbody>
<tr>
<td>No of women admitted</td>
<td>Dropped out before visit 1</td>
</tr>
<tr>
<td>Abidjan</td>
<td>188</td>
</tr>
<tr>
<td>Cotonou</td>
<td>259</td>
</tr>
<tr>
<td>Durban</td>
<td>192</td>
</tr>
<tr>
<td>Hat Yai</td>
<td>125</td>
</tr>
<tr>
<td>Total</td>
<td>764</td>
</tr>
</tbody>
</table>

*In Durban 5 women dropped out at onset of the trial, an additional 9 dropped out (3) or seroconverted (6) before the phase III part.

**Table 2** Key acceptability items on the first, second, and fifth follow up visits during phase III, by centre

<table>
<thead>
<tr>
<th></th>
<th>Abidjan</th>
<th>Cotonou</th>
<th>Durban</th>
<th>Hat Yai</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td>Gel not liked % of visits*</td>
<td>4.5</td>
<td>1.3</td>
<td>0.2</td>
<td>1.7</td>
<td>1.8</td>
</tr>
<tr>
<td>Gel easy to apply % (n) easy to apply†</td>
<td>98.2 (379)</td>
<td>99.8 (490)</td>
<td>98.8 (406)</td>
<td>94.4 (288)</td>
<td>98.1 (1563)</td>
</tr>
<tr>
<td>Would like to continue to use the gel % of visits</td>
<td>87.7</td>
<td>91.5</td>
<td>81.8</td>
<td>52.6</td>
<td>81.1</td>
</tr>
<tr>
<td>Gel affected intercourse % (n) affecting‡</td>
<td>24.3 (379)</td>
<td>41.4 (486)</td>
<td>18.2 (406)</td>
<td>35.8 (288)</td>
<td>30.1 (1559)</td>
</tr>
<tr>
<td>% (n) of affecting reporting improvement§</td>
<td>96.7 (92)</td>
<td>95.5 (201)</td>
<td>90.5 (74)</td>
<td>84.5 (103)</td>
<td>92.6 (470)</td>
</tr>
<tr>
<td>Why improved intercourse % (n) facilitating intercourse, causing no pain, and lubricating well</td>
<td>69.7 (89)</td>
<td>93.8 (192)</td>
<td>13.4 (67)</td>
<td>80.0 (80)</td>
<td>73.6 (428)</td>
</tr>
<tr>
<td>What liked most about gel % (n) facilitated intercourse, no pain, and lubrication*</td>
<td>41.6 (375)</td>
<td>49.0 (490)</td>
<td>2.2 (405)</td>
<td>40.0 (270)</td>
<td>33.3 (1540)</td>
</tr>
<tr>
<td>% prevention of STIs and other illness, including HIV†</td>
<td>5.1</td>
<td>22.5</td>
<td>53.6</td>
<td>7.8</td>
<td>23.8</td>
</tr>
<tr>
<td>Informed clients % of visits</td>
<td>56.9</td>
<td>53.5</td>
<td>23.1</td>
<td>21.8</td>
<td>38.8</td>
</tr>
<tr>
<td>Informed partners % of visits</td>
<td>53.9</td>
<td>45.2</td>
<td>40.7</td>
<td>21.8</td>
<td>41.9</td>
</tr>
</tbody>
</table>

*Numbers are expressed as percentages of visits, including those where no answer is recorded or when the answer is deemed not applicable by women. The rationale is that strong opinions are voiced when they are present. No answer means no strong opinion.
†Numbers are expressed as percentages of answers—that is, those (n) visits where an answer to the specific question is actually recorded. The missing data are thus treated as missing at random for this purpose.
‡The denominator consists of all (n) responses indicating that the gel affects intercourse.
acceptability since women staying in the study were more likely to have liked the gel. The difference between acceptability results obtained in trials v real life has been raised in a previous trial.5

In conclusion, our study showed that a small volume of gel was acceptable to most participants. Understanding the factors determining a product’s acceptability and thus its use is critically important since a product can only be effective when used regularly. However, not until an effective microbicide for HIV/STI prevention enters the market, can real life acceptability be assessed.

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CONTRIBUTORS
AV was involved in data analysis, writing of the manuscript, and performed statistical computing; EG was involved in data analysis and writing of the manuscript; GR, VC, MA, and VET were involved in the data collection and revision of the manuscript; LVD was involved in data analysis, writing of the manuscript, and was the overall study coordinator of the COL-1492 trial.

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