

Assessing Per-Sex-Act HIV-1 Risk Reduction Among Women Using the Dapivirine Vaginal Ring

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Background. Confounding introduced by individuals' sexual risk behavior is potentially a significant source of bias in HIV-1 prevention intervention studies. To more completely account for sexual behaviors when assessing the efficacy of the monthly dapivirine ring, a new longer-acting HIV-1 prevention option for women, we estimated per-sex-act risk reduction associated with product use.

Methods. We conducted a secondary analysis of data from MTN-020/ASPIRE, a phase 3, randomized, placebo-controlled efficacy trial of the dapivirine ring that recruited HIV-uninfected, African women aged 18–45 years. With cumulative sex acts as the time scale, we used multivariable Cox regression with inverse probability of censoring weights to estimate HIV-1 risk reduction associated with a rate of dapivirine release indicative of consistent product use.

Results. Women in the dapivirine ring group ($n = 1187$) had an estimated incidence rate of 2.3 (95% confidence interval [CI], 1.8–3.1) HIV-1 acquisition events per 10 000 sex acts versus 3.6 (95% CI, 2.9–4.4) per 10 000 acts in the placebo group ($n = 1187$). Dapivirine release indicative of consistent ring use was associated with a 63% (95% CI, 33%–80%) per-sex-act HIV-1 risk reduction.

Conclusions. These results support the efficacy of the dapivirine vaginal ring for HIV-1 prevention and help to inform decision-making for women, providers, and policymakers regarding product use.

Clinical Trials Registration. NCT01617096.

Keywords. dapivirine; vaginal ring; preexposure prophylaxis; HIV-1; per sex act.

In sub-Saharan Africa, women and girls account for more than half of new human immunodeficiency virus-1 (HIV-1) infections, underscoring the need for acceptable, novel, safe, and effective biomedical prevention interventions in this region [1]. Oral preexposure prophylaxis (PrEP) with tenofovir disoproxil fumarate and emtricitabine (TDF/FTC) is an important antiretroviral-based prevention strategy and is being scaled up rapidly in areas of sub-Saharan Africa with high HIV-1 burden [2]. However, challenges with daily PrEP adherence have been observed in past randomized trials and implementation studies [3–5], indicating that additional options are needed to fit women's diverse needs and preferences. One such option is the vaginal ring containing the nonnucleoside reverse transcriptase inhibitor dapivirine, a product with a demonstrated

favorable safety profile and HIV-1 risk reduction efficacy that women can wear continuously and discreetly for up to 1 month [6, 7]. In 2020, the ring received a positive opinion from the European Medicines Agency and prequalification and a recommendation from the World Health Organization [8–11]. The ring has since received regulatory approval in multiple countries in eastern and southern Africa [12–14].

As the dapivirine ring becomes an available PrEP option for women, it is important that its efficacy (ie, how well the product works when used under ideal conditions) be thoroughly assessed so that women can make well-informed decisions. The dapivirine ring was evaluated in 2 phase 3 efficacy trials in sub-Saharan Africa, MTN-020/ASPIRE and IPM 027/The Ring Study, and was found to be effective, reducing HIV-1 acquisition risk by 27% and 31% in intention-to-treat analyses, respectively [6, 7]. As observed in studies of oral PrEP agents [15, 16], efficacy was highly associated with greater adherence, and thus the intention-to-treat findings likely underestimate the product's efficacy when there are varying levels of adherence in the intention-to-treat population. Further evaluation with objective measures of adherence, however, has helped to further elucidate product efficacy. Within the Partners PrEP Study, a

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case-cohort analysis demonstrated a 91% relative risk reduction (RRR) associated with plasma tenofovir concentrations consistent with daily use among TDF/FTC users [17], in contrast to 75% effectiveness of TDF/FTC estimated in the intention-to-treat analysis [15]. Similarly, a substudy of the iPrEx trial found that intracellular tenofovir metabolite concentrations indicative of 4 or more oral doses per week were predictive of 90% HIV-1 risk reduction relative to placebo versus 44% risk reduction in intention-to-treat analysis [18, 19]. For the dapivirine ring, disproportionately low adherence was seen among certain groups, particularly younger women. A secondary analysis of data from MTN-020/ASPIRE identified reductions in HIV-1 risk of 48% or more among women who demonstrated continuous ring use based on measured rates of dapivirine released from the ring [20].

However, a key factor to consider when further evaluating the efficacy of HIV prevention products in secondary analyses of clinical trial data where exposure groups are no longer randomized is the potential for confounding by sexual behavior. HIV-1 acquisition is dependent on exposure to the virus through unprotected sex, specifically with a partner with HIV-1. However, analytic approaches to evaluating HIV-1 prevention methods that use a standard time scale assume that individuals who are not having sex accumulate risk at the same rate as individuals who are having frequent sex, which can lead to biased efficacy estimates if sexual behaviors systematically differ between product use groups. By assessing risk on a sex act scale, periods of time with no sexual activity, and therefore without any chance of exposure, would not inform relative risk estimates. This work builds on the previous secondary analysis within MTN-020/ASPIRE by more thoroughly accounting for sexual behavior and estimating per-sex-act risk reduction associated with rates of dapivirine release from returned rings indicative of continuous use. We hypothesized that HIV risk reduction estimates will be higher after more thorough accounting of women's sexual behaviors than previously estimated with dapivirine ring use.

METHODS

Study Population and Procedures

We conducted a secondary, longitudinal analysis of data collected from MTN-020/ASPIRE (NCT01617096), a phase 3 randomized placebo-controlled trial to assess the efficacy of the dapivirine vaginal ring for prevention of HIV-1 acquisition. Study procedures have been previously described in detail [6]. Briefly, healthy, HIV-uninfected cisgender women between ages 18 and 45 years, sexually active, and using an effective contraceptive method were recruited from 15 sites in Malawi, South Africa, Uganda, and Zimbabwe. Women were randomized 1:1 to use a silicone vaginal ring formulated with 25 mg of dapivirine or an inactive placebo ring and instructed to use the product continuously for 28 days. At monthly visits, women received a resupply of ring as well as

serologic HIV-1 testing and product adherence counseling. Safety of ring use was evaluated at each follow-up visit and study product was held for one or more of the following criteria: pregnancy or breastfeeding, positive HIV-1 rapid test, experience of adverse event or allergic reaction related to the study product, or report of HIV-1 postexposure prophylaxis. Ethics review committees at each site approved the study protocol and all participants provided written informed consent prior to initiation of study procedures.

Survey and Laboratory Measures

Information on women's sociodemographic characteristics, family planning use, menses/bleeding, and partner HIV-1 status and knowledge of ring use were collected by interviewer-administered surveys.

HIV-1 testing followed a standardized algorithm. At monthly study visits, women received 2 separate serological HIV-1 tests. If either test yielded a positive result, confirmatory western blot and HIV-1 RNA testing were performed on sera collected at the most recent quarterly visit. Archived sera from previous quarterly visits were also analyzed for HIV-1 RNA if needed to identify the earliest evidence of HIV-1 infection.

Testing for STIs and bacterial vaginosis (BV) occurred at the screening visit and then semiannually and when clinically indicated. Urine samples were tested for *Neisseria gonorrhoeae* and *Chlamydia trachomatis* using a nucleic acid amplification test. Rapid tests for *Trichomonas vaginalis* were conducted on vaginal swabs. Vaginal fluid samples were assessed for BV by Gram stain using the Nugent score.

Adherence Measures

Ring adherence was evaluated using multiple objective measures. First, stored plasma specimens from quarterly study visits were analyzed for dapivirine concentrations via a previously described ultraperformance liquid chromatography-tandem mass spectrometry assay [21]. Second, approximately 1 year after study enrollment began, used rings were collected at monthly visits and analyzed for quantities of residual dapivirine to assess cumulative use. Residual dapivirine in rings was collected using acetone extraction and then quantified using high-pressure liquid chromatography. The amount of drug released during intravaginal use was estimated by subtracting the amount of residual dapivirine measured in the ring from the batch-specific dapivirine loading mass. Given that women may not have returned to the clinic exactly 4 weeks after their previous visit, we standardized the amount of dapivirine released to 28 days of use. In previous phase 1 studies, approximately 4–5 mg of drug was released from the ring among women who reported continuous product use over a 28-day period [22, 23]. Because this analysis uses a continuous dapivirine release rate exposure variable, we used the point estimate at the midpoint of these values (4.5 mg/month) to assess risk reduction associated with

consistent ring use. Use of the 4.5 mg/month midpoint allows for adequate comparison to previous evaluations of ring efficacy that use categorical exposure variables. These studies define consistent ring use as >4.0 mg/month, which includes all dapivirine release rates between 4 and 5 mg/month. Given that measurement of residual dapivirine in returned rings began a year into the study, we imputed monthly dapivirine release rates using a joint hidden Markov model ([Supplementary Methods](#)).

Estimating Per-Sex-Act Risk Reduction

Modeling the Sex Act Time Scale

At quarterly visits, women were asked to provide information about their sexual behaviors, including the number of sexual partners they had in the past 3 months, whether they had a consistent primary partner over the past 3 months, the number of vaginal sex acts with their primary partner in the past week, and number of times a condom was used. Women were also asked about numbers of anal sex acts; however, only vaginal sex acts were considered in this analysis given that anal sex may have occurred at the same encounter and receptive anal intercourse was not found to be an independent risk factor for HIV-1 acquisition in prior analyses of ASPIRE participants [24]. To carry out our analysis on the per-sex-act scale, we required complete monthly sexual behavior data for participants across follow-up. To predict numbers of monthly sex acts, we modeled participant-reported sex acts in the past week using a mixed effects Poisson model. The model included participant-specific intercepts and slopes for the study visit month variable to allow for individual variation in sex acts at baseline and change in sex acts over time. We included as fixed effects baseline and time-varying covariates thought to be correlated with sexual behaviors based on our own hypotheses and the published literature, including study site, age, marital status, number of live births, BV status, *N. gonorrhoeae*, *C. trachomatis*, and *T. vaginalis* infection, alcohol use, smoking status, having a primary or new sexual partner, number of sexual partners, number of sex acts at baseline, family planning use, and having any vaginal cleaning practices. A single fitted model was used to predict numbers of sex acts since the last monthly visit.

Regression Models

Models of per-sex-act risk reduction included women who were confirmed to be without HIV-1 at enrollment and had imputed sex acts and dapivirine release data across follow-up. Additionally, we excluded women from 2 study sites with lower product adherence and study retention that were discontinued early, as was done in a post hoc analysis for the ASPIRE trial. To model per-sex-act HIV-1 risk reduction associated with ring release rates of 4.5 mg/month (ie, consistent ring use), we fit Cox regression models using the cumulative number of predicted sex acts between enrollment and first HIV-1 detection as the outcome.

Given that tests for HIV-1 RNA were only conducted quarterly, the month when HIV-1 acquisition occurred was imputed using the same joint hidden Markov model used to impute monthly dapivirine release rates ([Supplementary Methods](#)). Primary exposure variables for study group (dapivirine or placebo ring) and continuous ring release rate were included in the Cox models; the exponentiated linear combination of the coefficients of these 2 terms was used to obtain a parameter estimate analogous to a hazard ratio (HR), comparing the probability of HIV-1 infection at a particular number of cumulative sex acts between women in the dapivirine ring group with 4.5 mg/month drug release and women in the placebo ring group (for whom dapivirine release rate was 0 mg/month), assuming no HIV-1 infection up to that point. Percent RRR was calculated using the following formula: $\%RRR = (1 - HR) \times 100$. Women were censored at first pregnancy, first product hold >3 days, and loss to follow-up given likely changes in HIV-1 risk after these events. All models were stratified by study site. Multivariable models were adjusted for time-varying participant-reported condomless sex in the past 7 days and the following baseline variables: age, education, marital status, BV status, *N. gonorrhoeae*, *C. trachomatis*, and *T. vaginalis* diagnosis, any new sexual partners, multiple sexual partners, number of sex acts in last 3 months, any reported condomless sex in last 7 days, alcohol use, partner HIV-1 status and knowledge of ring use, family planning method, and age-specific local male HIV-1 prevalence (assuming that women had partners who were in their age group). Model parameter estimates were calculated separately for each of the 20 imputed datasets and averaged; standard errors were pooled using Rubin's rules to obtain 95% confidence intervals (CIs) and 2-sided *P* values. Given that women in our analysis were censored for reasons that may be related to the study outcome of HIV-1 acquisition, we further adjusted our models using inverse probability of censoring weights (IPCWs) to account for potential selection bias due to informative censoring ([Supplementary Methods](#)). Subgroup analyses were carried out based on baseline age (<25 and ≥25 years), *N. gonorrhoeae*/*C. trachomatis*/*T. vaginalis* diagnoses, BV status, and whether women reported bleeding in the prior month given that women may have less HIV-1 exposure while experiencing bleeding. All statistical analyses were carried out in R version 3.6.1 (R Core Team).

RESULTS

In ASPIRE, 2629 women were randomized to use of either the dapivirine (*n* = 1313) or placebo ring (*n* = 1316). Of those women, 3 were retrospectively discovered to have had acute HIV-1 infection at baseline and were excluded. An additional 12 women had no follow-up after the enrollment visit. For this analysis, 9 women who had a product hold before their first follow-up visit, 17 with missing data for variables used in prediction and Markov chain Monte Carlo (MCMC) models, and an additional 214 women from 2 sites with lower product

Table 1. Baseline Participant Characteristics

Characteristic	Placebo Group (n = 1187)	Dapivirine Group: No Months With Estimated Release Rate ≥ 4.5 mg/mo (n = 813)	Dapivirine Group: at Least 1 Month With Estimated Release Rate ≥ 4.5 mg/mo (n = 374)
Age, y, median (IQR)	26.0 (23.0–31.0)	27.0 (23.0–32.0)	26.0 (22.0–31.0)
Country			
Malawi	136 (11.5)	103 (12.7)	29 (7.8)
South Africa	592 (49.9)	373 (45.9)	220 (58.8)
Uganda	126 (10.6)	100 (12.3)	27 (7.2)
Zimbabwe	333 (28.1)	237 (29.2)	98 (26.2)
Highest level of education			
None	11 (0.9)	10 (1.2)	1 (0.3)
Primary, not complete	111 (9.4)	98 (12.1)	26 (7.0)
Primary, complete	65 (5.5)	50 (6.2)	18 (4.8)
Secondary, not complete	466 (39.3)	313 (38.5)	136 (36.4)
Secondary, complete	462 (38.9)	311 (38.3)	155 (41.4)
College or university	72 (6.1)	31 (3.8)	38 (10.2)
Married	537 (45.2)	383 (47.1)	135 (36.1)
No. of live births, median (IQR)	2.0 (1.0, 3.0)	2.0 (1.0, 3.0)	2.0 (1.0, 2.0)
Has primary partner	1181 (99.5)	811 (99.8)	373 (99.7)
Partner knows about ring use	779 (66.0)	530 (65.4)	238 (63.8)
Partner is circumcised	508 (43.0)	331 (40.8)	161 (43.2)
Partner HIV-1 status			
Positive	12 (1.0)	15 (1.8)	7 (1.9)
Negative	650 (55.0)	429 (52.9)	198 (53.1)
Don't know	519 (43.9)	367 (45.3)	168 (45.0)
Multiple sexual partners in the past 3 mo	211 (17.8)	135 (16.6)	57 (15.2)
Any new partner in past 3 mo	34 (2.9)	28 (3.4)	16 (4.3)
No. of sex acts in last 3 mo, median (IQR)	21.0 (8.0–36.0)	24.0 (10.0–40.0)	15.0 (6.0–36.0)
No. of sex acts in last 7 d, median (IQR)	2.0 (1.0–3.0)	2.0 (1.0–4.0)	1.0 (0.0–3.0)
Any condomless sex with partner in last 7 d	524 (44.1)	359 (44.2)	125 (33.4)
Intrauterine device	159 (13.4)	128 (15.7)	32 (8.6)
Oral pills	118 (9.9)	88 (10.8)	32 (8.6)
Hormonal implant	238 (20.1)	194 (23.9)	62 (16.6)
Depot medroxyprogesterone acetate	500 (42.1)	279 (34.3)	173 (46.3)
Norethisterone enanthate	149 (12.6)	96 (11.8)	75 (20.1)
Any menstrual bleeding in past 3 mo	926 (78.0)	656 (80.8)	276 (74.0)
Any vaginal practices	771 (65.0)	533 (65.6)	219 (58.6)
Usual bleeding pattern at screening			
Amenorrheic	254 (21.4)	147 (18.1)	99 (26.5)
Irregular	301 (25.4)	194 (23.9)	95 (25.4)
Regular	632 (53.2)	472 (58.1)	180 (48.1)
<i>Neisseria gonorrhoeae</i>	48 (4.0)	39 (4.8)	14 (3.7)
<i>Chlamydia trachomatis</i>	126 (10.6)	96 (11.8)	51 (13.6)
<i>Trichomonas vaginalis</i>	82 (6.9)	60 (7.4)	21 (5.6)
Bacterial vaginosis	472 (39.8)	379 (46.6)	1 (31.8)

Data are No. (%) except where indicated. Abbreviation: IQR, interquartile range.

adherence and study retention were also excluded. Therefore, we included a total of 2374 women (1187 each in the dapivirine and placebo groups) in our analysis.

Women in the placebo and dapivirine ring groups (regardless of estimated release rates) had comparable sociodemographic characteristics (Table 1). Women who were estimated to have demonstrated consistent ring use at least once during the trial (≥ 4.5 mg/month) reported a median of 15.0 sex acts in the last 3 months (interquartile range [IQR], 6.0–36.0) and 33.4%

reported recent condomless sex. Depot medroxyprogesterone acetate was the most used contraceptive method (used by 40.1% of women) at enrollment among all participants, and most women experienced regular bleeding at screening (53.2%).

In our mixed model to predict monthly sex acts, the standard deviation of the study month random effect term was 0.35. The median number of predicted sex acts per 28 days across follow-up was similar between women in the placebo and dapivirine ring groups (median, 8; IQR, 5–13 in both groups) and sexual

Table 2. Per-Sex-Act HIV-1 Relative Risk Reduction Associated With Consistent Dapivirine Ring Use (Estimated Release of 4.5 mg/mo)

	Dapivirine Group: HIV-1 Events Per 10 000 Sex Acts (95% CI) ^a	Placebo Group: HIV-1 Events Per 10 000 Sex Acts (95% CI)	Unadjusted Model		Adjusted Model ^b		Adjusted and Weighted Model ^{b,c}		
			Per-Sex-Act RRR, % (95% CI)	P	Per-Sex-Act RRR, % (95% CI)	P	Per-Sex-Act RRR, % (95% CI)	P	Interaction P
Overall	2.3 (1.8–3.1)	3.6 (2.9–4.4)	57 (24–75)	.004	62 (32–79)	.001	63 (33–80)	.001	...
Baseline age									
<25 y (n = 886)	5.0 (3.5–6.9)	5.3 (3.9–7.3)	53 (0–77)	.049	61 (16–82)	.016	62 (17–82)	.014	.844
≥25 y (n = 1488)	1.1 (.7–1.8)	2.8 (2.0–3.7)	47 (–25 to 77)	.148	54 (–20 to 82)	.114	55 (–16 to 83)	.096	
Baseline STIs/BV									
<i>N. gonorrhoeae</i> not detected (n = 2273)	2.1 (1.6–2.8)	3.3 (2.6–4.2)	56 (19–76)	.009	61 (27–79)	.003	62 (29–80)	.003	.683
<i>N. gonorrhoeae</i> detected (n = 101)	7.2 (2.9–14.9)	11.0 (5.0–20.8)	59 (–104 to 92)	.277	72 (–67 to 95)	.162	72 (–64 to 95)	.159	
<i>C. trachomatis</i> not detected (n = 2101)	1.9 (1.3–2.6)	3.2 (2.5–4.1)	49 (3–73)	.039	53 (9–76)	.026	54 (10–77)	.023	.181
<i>C. trachomatis</i> detected (n = 273)	6.2 (3.5–10.2)	7.3 (4.2–11.9)	75 (2–92)	.020	81 (37–95)	.007	83 (42–95)	.005	
<i>T. vaginalis</i> detected (n = 2211)	2.1 (1.6–2.9)	3.5 (2.7–4.3)	62 (30–80)	.002	66 (35–83)	.001	68 (38–83)	.001	.250
<i>T. vaginalis</i> detected (n = 163)	4.9 (2.1–9.7)	5.2 (2.2–10.2)	–26 (–621 to 78)	.798	16 (–348 to 84)	.839	10 (–363 to 83)	.899	
BV not detected (n = 1404)	2.0 (1.4–2.9)	2.7 (1.9–3.7)	51 (–6 to 77)	.071	53 (–1 to 78)	.052	54 (1 to 78)	.048	.441
BV detected (n = 970)	2.8 (1.8–4.1)	5.0 (3.6–6.7)	63 (10–84)	.028	71 (26–89)	.010	72 (28–89)	.008	
Menses/bleeding in past mo									
No	2.9 (2.0–3.9)	4.0 (3.0–5.2)	47 (0–72)	.049	55 (12–77)	.020	56 (14–78)	.017	.282
Yes	1.6 (.9–2.7)	2.9 (1.9–4.2)	77 (26–93)	.014	78 (28–94)	.013	79 (30–94)	.011	

All models were stratified by study site. RRR calculated as (1 – hazard ratio) × 100.

Abbreviations: BV, bacterial vaginosis; CI, confidence interval; HIV-1, human immunodeficiency virus-1; RRR, relative risk reduction; STI, sexually transmitted infection.

^aCrude incidence rates of HIV-1 events per 10 000 sex acts include all participants in the dapivirine ring group. However, the per-sex-act RRR estimates in the table compare the probability of HIV-1 infection between women in the dapivirine ring group with 4.5 mg/month drug release and women in the placebo ring group (for whom dapivirine release rate was 0 mg/month). Hence, comparison of the crude incidence rates will not equate to the unadjusted RRR estimate.

^bMultivariable Cox regression models were adjusted for report of any condomless sex and the following baseline variables: age, education, marital status, BV status, *Chlamydia trachomatis*, *Neisseria gonorrhoeae*, and *Trichomonas vaginalis* diagnosis, any new sexual partners, multiple sexual partners, number of sex acts in last 3 months, any condomless sex in last 7 days, alcohol use, partner HIV-1 status and knowledge of ring use, family planning method, and local HIV-1 prevalence for men within the same age group.

^cAdjusted using inverse probability of censoring weights to account for selection bias due to participant censoring at first pregnancy, first product hold >3 days, and loss-to-follow-up given likely changes in HIV-1 risk after these events.

behaviors gradually declined in frequency over time (Supplementary Figure 1A). Among women in the dapivirine ring group, the median estimated dapivirine release rate across the 20 posterior samples was 3.3 mg/28 days (IQR, 2.7–3.7) (distribution of dapivirine release rates by study visit shown in Supplementary Figure 1B). The median estimated dapivirine release rate was similar across categories of sex acts (Supplementary Figure 1C).

Overall, women in the placebo group experienced an estimated 83 HIV-1 acquisition events over 231 637 predicted sex acts, for an incidence rate of 3.6 events per 10 000 acts (95% CI, 2.9–4.4) (Table 2). Women in the dapivirine group experienced an estimated 54 HIV-1 acquisition events over 230 517 predicted sex acts, for an incidence rate of 2.3 events per 10 000 acts (95% CI, 1.8–3.1).

Women in the dapivirine ring group demonstrating consistent ring use (estimated dapivirine release of 4.5 mg/28 days) had a 57% reduction in HIV-1 risk (95% CI, 20%–74%) relative to placebo in unadjusted models. In multivariable models adjusting for time-varying report of any condomless sex and baseline variables for age, education, marital status, BV status,

N. gonorrhoeae/*C. trachomatis*/*T. vaginalis* diagnoses, sexual behaviors, alcohol use, partner HIV-1 status and knowledge of ring use, family planning method, and local HIV-1 prevalence for men within the same age group, release rates associated with consistent ring use were associated with a 62% HIV-1 RRR (95% CI, 32%–79%) compared to placebo.

Mean stabilized IPCW values were close to 1 (South Africa sites IPCW mean, 0.997 [range: 0.319–3.514]; non-South Africa sites IPCW mean, 0.995 [range: 0.505–10.550]). However, standardized mean differences of covariates between censored and uncensored women were similar after weighting, indicating possible inadequate model specification (Supplementary Figure 3). In the adjusted model incorporating IPCWs, dapivirine release indicative of consistent ring use was associated with a 63% RRR (95% CI, 33%–80%). There was significant risk reduction with dapivirine release rates as low as 1.3 mg/month, but risk reduction estimates were higher with greater dapivirine release (Figure 1).

In subgroup analyses, estimated per-sex-act HIV-1 incidence was highest among women with detected *N. gonorrhoeae* (11.0 events per 10 000 acts in placebo group and 7.2 events per 10 000 acts in dapivirine group) and *C. trachomatis* (7.3 events

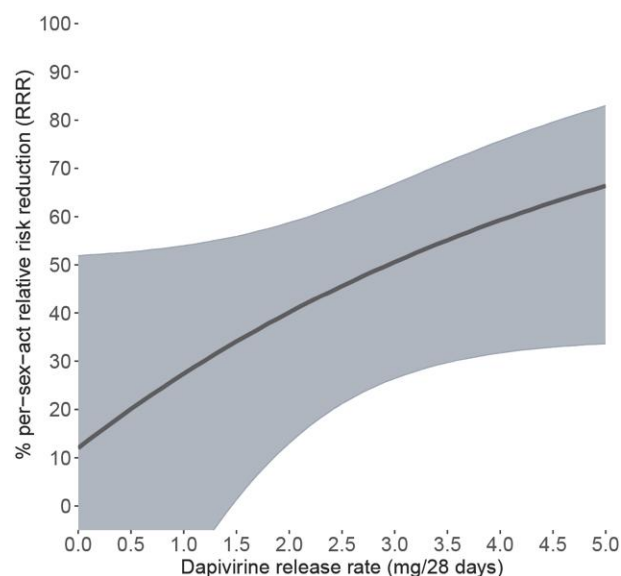


Figure 1. Per-sex-act HIV-1 risk reduction with incremental dapivirine release rates. Solid line indicates percent relative risk reduction estimate. Shaded area indicates 95% confidence interval.

per 10 000 acts in placebo group vs 6.2 events per 10 000 acts in dapivirine group) at enrollment. However, RRR estimates did not differ between women with or without *N. gonorrhoeae* and *C. trachomatis* infection at enrollment according to interaction models ($P = .683$ and $P = .181$, respectively). Additionally, no significant difference in RRR was identified between age, BV status, and *T. vaginalis* detection subgroups. In our interaction model based on time-varying experience of bleeding in the past month, risk reduction estimates were higher for months when bleeding was reported versus when not reported (Supplementary Figure 2), but this difference was not statistically significant ($P = .282$).

DISCUSSION

In this secondary analysis of data from the ASPIRE trial, we conducted a rigorous evaluation of HIV-1 risk reduction associated with use of the dapivirine vaginal ring with a more complete accounting for women's sexual behaviors. Unlike other analyses that evaluate HIV-1 risk reduction on a time scale, assessing risk reduction on the sex act scale may better control for the heterogeneity in women's sexual behaviors and ensure that accumulation of HIV-1 risk is directly tied to potential exposure. We demonstrated a positive relationship between rate of drug release from the dapivirine vaginal ring and HIV-1 risk reduction. Further, dapivirine release rates indicative of consistent ring use were associated with an estimated 63% reduction in per-sex-act HIV-1 risk after adjusting for other potential confounders and accounting for informative censoring using IPCWs.

Dapivirine release rates were imputed in our analysis using MCMC methods to account for missingness and measurement error in observed ring release measures. In our previous analysis using all available measured ring release data, which excluded 46 women with HIV-1 end points that occurred prior to collection of used rings for dapivirine quantification, we demonstrated a 43% risk reduction associated with dapivirine release rates consistent with continuous use [20]. In this previous analysis, the same levels of dapivirine release were associated with a 71% HIV-1 risk reduction when starting follow-up at 12 months to create a common start time; however, as this was a secondary analysis that excluded an additional 54 HIV-1 end points and 1436 woman-years of follow-up, it is possible this estimate was biased due to missing data from early study visits. Here, our ability to use the full range of study follow-up data due to our imputation methods allows us to correct for selection bias due to previous exclusion of these end points.

We observed little change in HIV-1 risk reduction estimates when models were weighted using IPCWs. This was also observed in previous work utilizing IPCWs to evaluate the efficacy of oral PrEP, which censored participants at first missed visit and detection of poor adherence [19]. Estimating unbiased IPCWs is dependent correct model specification [25, 26] and inclusion of all variables associated with time to censoring and time to failure. Therefore, this small change could be due to poor weight model specification and might suggest our models exclude other unmeasured predictors of censoring. This is supported by our IPCW diagnostics that showed similar covariate balance between censored and uncensored before and after weighting.

Other potential limitations of our analysis should be considered. First, predicted numbers of monthly sex acts were based on women's self-reported sex acts in the prior week. Such data are subject to recall bias (particularly among women who have frequent sex) and social desirability bias. Additionally, given that estimated values (ie, predicted sex acts and IPCWs) are treated as fixed quantities in the Cox regression models, the pooled standard errors used to estimate per-sex-act HIV risk reduction may not fully reflect the variability in these data. Second, per-sex-act risk depends largely on the HIV-1 status of their partner. However, <2% of women reported that their primary partner was living with HIV-1 at baseline while nearly half of women did not know their partner's status. We also adjusted for age-specific local HIV-1 prevalence among men to further account for partner HIV-1 status and variation in women's HIV-1 exposure, but this is an imperfect proxy for confirmation of women's partner status. As a result, per-act incidence calculated in our analysis is likely an underestimate of the per-act incidence of women who are HIV-exposed.

Our analysis supports the use of the dapivirine ring as an efficacious HIV-1 prevention product for women. While the efficacy of the ring for preventing male-to-female HIV-1 transmission via vaginal sex is not as high as the efficacy of daily PrEP or consistent

condom use, both of which reduce the risk of HIV transmission via multiple routes and are associated with >90% risk reduction [17, 27], the ring has important characteristics that may suit some women's needs (eg, longer duration, less user-dependent, allows for discreet use, potentially fewer side effects, and high acceptability [28]). Therefore, communication of risk associated with the ring relative to other PrEP options should be part of a person-centered approach, acknowledging that women are in the best position to decide which option is best for their individual situation [29]. As additional HIV-1 prevention products emerge from the development pipeline, including multipurpose prevention technologies, we encourage similar analyses to evaluate risk reduction associated with product exposure as a complement to intention-to-treat analysis as an approach to best understand product efficacy.

Supplementary Data

Supplementary materials are available at *The Journal of Infectious Diseases* online (<http://jid.oxfordjournals.org/>). **Supplementary materials** consist of data provided by the author that are published to benefit the reader. The posted materials are not copyedited. The contents of all **supplementary data** are the sole responsibility of the authors. Questions or messages regarding errors should be addressed to the author.

Notes

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design. R. M. S. led the data analyses. T. Q. D. and E. R. B. contributed to the data analyses and provided statistical guidance. C. W. H., T. P. P., A. v. d. S., S. L. H., F. M. K., N. M. M., M. A. M., L. G. B., L. S. T., J. M. B., and E. R. B. led parts of or directly contributed to implementation and data collection for the parent study (MTN-020/ASPIRE). R. M. S. drafted the initial manuscript, and all authors contributed to results interpretation and manuscript revisions. All authors have read and approved the manuscript.

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