6-1-2017


Kerry A. Thomson  
*University of Washington*

Jessica Haberer  
*Massachusetts General Hospital*

Mark A. Marzinke  
*Johns Hopkins University*

Andrew Mujugira  
*Makerere University*

Craig Hendrix  
*Johns Hopkins University*

*See next page for additional authors*

Let us know how access to this document benefits you.

Follow this and additional works at: [https://pdxscholar.library.pdx.edu/sph_facpub](https://pdxscholar.library.pdx.edu/sph_facpub)

Part of the [Community Health and Preventive Medicine Commons](https://pdxscholar.library.pdx.edu/sph_facpub)

Citation Details


This Article is brought to you for free and open access. It has been accepted for inclusion in OHSU-PSU School of Public Health Faculty Publications and Presentations by an authorized administrator of PDXScholar. For more information, please contact [pdxscholar@pdx.edu](mailto:pdxscholar@pdx.edu).
Medication Sharing Is Rare Among African HIV-1 Serodiscordant Couples Enrolled in an Efficacy Trial of Oral Pre-exposure Prophylaxis (PrEP) for HIV-1 Prevention

Kerry A. Thomson, MPH,* Jessica E. Haberer, MD, MS,† Mark A. Marzinke, PhD,‡ Andrew Mujugira, MBChB, PhD, MSc,§ Craig W. Hendrix, MD,∥ Connie Celum, MD, MPH,¶ Patrick Ndase, MBChB, MPH,∥ Allan Ronald, MD,,** David R. Bangsberg, MD, MPH,†† and Jared M. Baeten, MD, PhD,‡‡ for the Partners PrEP Study Team

Abstract: Sharing of pre-exposure prophylaxis (PrEP) medications is a concern for PrEP implementation. For HIV-1 serodiscordant couples, sharing may undermine the HIV-1 prevention benefit and also cause antiretroviral resistance if taken by HIV-1 infected partners. Within a PrEP efficacy trial among HIV-1 serodiscordant couples, we assessed the occurrence of PrEP sharing by self-report and plasma tenofovir concentrations in HIV-1 infected partners. PrEP sharing was self-reported at <0.01% of visits, and 0%–1.6% of randomly selected and 0% of purposively selected specimens from HIV-1 infected participants had detectable tenofovir concentrations (median: 66.5 ng/mL, range: 1.3–292 ng/mL). PrEP sharing within HIV-1 serodiscordant couples was extremely rare.

Key Words: HIV-1, pre-exposure prophylaxis, HIV-1 serodiscordant couples, adherence, prescription drug diversion

INTRODUCTION

Pre-exposure prophylaxis (PrEP), in which an HIV-1 uninfected individual takes oral antiretrovirals to prevent HIV-1 acquisition, is recommended by the World Health Organization for populations at high risk of HIV-1, including HIV-1 serodiscordant couples. Open-label studies of PrEP are ongoing, and have shown high adherence and reductions in HIV-1 acquisition of 90% or more. “Drug diversion,” where HIV-1 uninfected individuals share or sell their prescribed PrEP medications, is a potential challenge for PrEP implementation. To sufficiently disrupt HIV-1 replication, antiretroviral therapy (ART) for HIV infected adults should include at least 3 drugs from 2 classes of HIV antiretrovirals. Use of mono or dual agent antiretrovirals prescribed as PrEP by HIV-1 infected persons could result in antiretroviral resistance due to suboptimal viral suppression and undermine the level of protection for the intended HIV-1 uninfected user due to insufficient drug concentrations. PrEP sharing may occur more frequently by HIV-1 uninfected persons with a known HIV-1 infected partner, particularly if their partner is not yet eligible for or experiences barriers accessing ART for their own treatment. We assessed the extent to which PrEP sharing occurred within a large cohort of HIV-1 serodiscordant couples enrolled in an efficacy trial in Africa.

METHODS

Study Population

The Partners PrEP Study was a phase III, randomized, double-blind, placebo-controlled, 3-arm clinical trial of daily oral PrEP (ClinicalTrials.gov NCT00557245). Beginning in 2008, the HIV-1 uninfected partner in 4747 HIV-1 serodiscordant couples was randomized to receive once-daily tenofovir (TDF), combination tenofovir–emtricitabine (TDF/FTC), or matching placebo and followed for a maximum of 48 months for safety and HIV-1 seroconversion. The Partners PrEP Study was conducted at 9 clinical research sites in Kenya and Uganda; study details have been described previously. In July 2011, the Data and Safety Monitoring Board recommended discontinuation of the placebo arm due to the demonstrated 67% efficacy for HIV-1 prevention with
TDF and 75% efficacy with TDF/FTC. Participants were offered the option to continue in the study and receive open-label PrEP, with participants in the placebo arm re-randomized to TDF or TDF/FTC.

Study Procedures

This analysis includes data from the duration of study follow-up (2008–2013). HIV-1 uninfected participants attended monthly study visits. At each visit, participants were counselled on the importance of adherence to study drug and the potential danger of HIV-1 infected partners taking study drug. Plasma from HIV-1 uninfected participants was collected at enrollment, months 1, 3, and quarterly thereafter, and study exit. At enrollment, all HIV-1 infected partners were ineligible for ART based on national guidelines and self-reported not taking ART. HIV-1 infected participants attended quarterly study visits and were referred to initiate ART once they met national eligibility criteria. Plasma from HIV-1 infected participants was collected at enrollment, at 6 months intervals thereafter, and study exit. Plasma was also collected at any visit where an HIV-1 uninfected participant first tested positive for HIV, and as soon as possible from their study partner. Self-reported data on sharing study drug, including the number of tablets shared, were obtained from HIV-1 uninfected partners at monthly study visits, re-randomization, study exit, and annually from HIV-1 infected partners.

Assessment of PrEP Medication Sharing

For the current analysis, we assessed PrEP medication sharing using self-report and objective assessment through plasma tenofovir testing in 4 different groups that were selected based on potential or previously documented non-adherence to study drug (Fig. 1). Tenofovir concentrations in plasma were measured using previously described validated liquid chromatography-mass spectrometry/mass spectrometry methods. Detectable tenofovir for all analyses was defined as ≥0.31 ng/mL; this threshold was the lower limit of quantification and has 92% specificity for at least one dose in the past week. Results from 2 prior analyses informed the sampling scheme for the current study, and additional testing of plasma from HIV-1 infected subjects was performed. The Ancillary Adherence Study (AAS) assessed use of study drug during the blinded phase of the trial at 3 sites in Uganda. In this substudy, plasma was collected from both HIV-1 uninfected and HIV-1 infected members of 1147 couples at unannounced home visits during follow-up. Group 1 was a random sample of plasma specimens collected at AAS unannounced home visits from HIV-1 infected participants (n = 100), with the rationale that participants sharing medication may modify this behavior before clinic visits and thus an unannounced visit would be the best opportunity to identify sharing. Group 2 included specimens belonging to HIV-1 infected participants enrolled in the AAS whose HIV-1 uninfected study partner’s specimen was included in a previous analysis testing 268 randomly selected home specimens and did not have detectable tenofovir concentrations (n = 29). Group 3 was derived from a case-cohort study of plasma tenofovir concentrations nested within the Partners PrEP Study, including 298 randomly-selected HIV-1 uninfected participants from the active arms, which assessed the association between PrEP use and HIV-1 protection. Group 3 included specimens belonging to HIV-1 infected participants whose HIV-1 uninfected study partner’s specimen did not have detectable tenofovir concentrations among the 1802 randomly selected specimens in the cohort analysis (n = 406). The rationale for both Groups 2 and 3 was that the absence of tenofovir in the HIV-1 uninfected partner may have been because PrEP medication was shared with the HIV-1 infected partner. Group 4 included HIV-1 infected participants whose study partner acquired HIV-1 during study follow-up (n = 52), with the rationale that these participants were not taking sufficient PrEP to achieve

FIGURE 1. Assessment of potential sharing of PrEP medication within HIV-1 serodiscordant couples, frequency, (%), [95% CI].
the HIV-1 prevention benefit and may have diverted PrEP to their partner.

The underlying sampling scheme for Groups 1, 2, and 3 was a random selection across the study population; within Groups 2 and 3, and for Group 4, purposeful testing was done to investigate drug sharing because uninfected partners had no tenofovir concentrations in plasma and/or acquired HIV-1 during follow-up, despite having access to active study drug. Specimens from couples who had been randomized to the trial’s active arms and time periods when PrEP was dispensed (ie, excluding study drug holds or missed visits) and ART was not reported by the HIV-1 infected partner were eligible for testing. We tested the specimen from the HIV-1 infected participant that was closest to the date of undetectable tenofovir concentrations (or HIV-1 seroconversion) in their HIV-uninfected partner. For any HIV-1 infected participant with detectable tenofovir concentrations we also tested the following: an additional specimen from the same visit to rule out inadvertent specimen mix-up, plasma archived at enrollment/randomization (to assess undisclosed ART use), plasma HIV-1 RNA viral load (undetectable defined as <80 copies/mL), and plasma for emtricitabine.

Statistical Methods

We used SAS 9.4 (SAS Institute, Cary, NC) for descriptive analysis; 95% confidence intervals were generated with Stata 13.1 (StataCorp, College Station, TX), and one-sided exact 95% confidence intervals were generated for proportions in which the numerator was zero.

Ethics

The protocols for the parent study and AAS sub-study were approved by ethics review committees at the University of Washington and each study site. All participants provided written informed consent.

RESULTS

Self-Report of Medication Sharing

Eight instances of HIV-1 infected partners using study medication were self-reported from 7 couples across 155,875 study visits (8/155,875, <0.01%). Five of these instances were reported during follow-up and 3 were reported at study exit, including one partner who reported medication sharing once during follow-up and again at study exit (Fig. 1). No seroconverter reported that their study partner had used their study medication. All self-reported instances of HIV-1 infected participants taking study medication were described as a small number of pills (maximum 4) during a single study month, and there was no increase in reported drug sharing once the placebo arm was discontinued.

Plasma Drug Concentrations

Of 290 specimens tested for this analysis, a total of 11 specimens from HIV-1 infected participants had detectable tenofovir concentrations, though only 3 specimens are potential instances of drug sharing (Fig. 1). These 11 specimens came from 9 participants, none of whom self-reported drug sharing, and all were investigated further for potential drug sharing and undisclosed ART use (Table 1). Two of the 100 specimens in Group 1 had detectable tenofovir concentrations. Follow-up testing determined that Case 1 had a tenofovir concentration of 71.5 ng/mL and an undetectable HIV-1 viral load at enrollment, suggesting unreported tenofovir-containing ART use before randomization and not drug sharing thereafter. Case 2 had undetectable viral load and a very low tenofovir concentrations at month 11 (1.3 ng/mL), but no tenofovir present at enrollment. Thus, PrEP sharing may have occurred in this case and overall, potential drug sharing in Group 1 was 1.0%. In Group 2, 2 specimens from the same participant had detectable tenofovir concentrations (Case 3). This couple was randomized to the TDF arm; however, the HIV-1 infected partner had both tenofovir and emtricitabine detected in a specimen collected at enrollment. Case 3 is suggestive of undisclosed ART use before study entry and unlikely PrEP sharing thereafter; therefore potential drug sharing in Group 2 was 0%. In Group 3, 7 specimens from 6 participants had detectable tenofovir concentrations (Cases 4–9). Three participants randomized to the TDF/ FTC arm had detectable tenofovir without emtricitabine (Case 4–6) and Case 7 had detectable tenofovir before randomization, all circumstances suggesting undisclosed ART use. Case 8 had detectable tenofovir concentrations at month 24 of follow-up and drug concentrations persisted until study exit at month 36, and Case 9 had detectable tenofovir concentrations and undetectable HIV-1 viral load at month 33. In the absence of detectable tenofovir concentrations at randomization for these 2 cases, potential drug sharing in Group 3 was 1.6%. No specimens in Group 4 had detectable tenofovir and therefore potential drug sharing in this group was 0%.

DISCUSSION

We found strong evidence that PrEP medication sharing was extremely infrequent among East African HIV-1 serodiscordant couples enrolled in a PrEP efficacy trial. No extended drug sharing was self-reported, and random testing and purposive sampling among groups most likely to share study drug identified only 3 couples with evidence of potential sharing of PrEP. Only one prior study has explored drug sharing within the context of a PrEP efficacy trial and also found that it was rare. Among a subset of HIV-1 uninfected women enrolled in the FEM-PrEP study, 10/224 (4%) of women reported that they gave their study pills to someone else and described instances of selling or sharing study medication to HIV-1 infected and uninfected individuals. While outside the scope of the current analysis, sharing PrEP with other HIV-1 uninfected individuals who are not receiving adherence counselling and frequent HIV-1 testing is also a concern for PrEP delivery and should be explored in future studies.

For 6 HIV-1 infected partners, we found detectable drug at a time point prior to their study partner receiving PrEP and/or concentrations that were not consistent with the arm to
which the study couple had been randomized. Although it was not feasible to test these specimens for all possible antiretrovirals, the timing detectable tenofovir or emtricitabine strongly suggests undisclosed ART use. This finding is consistent with prior studies that have described undisclosed ART use occurring among 2.8%–5% of HIV-1 infected partners participating in clinical trials of HIV-1 serodiscordant couples. 21,22 Undisclosed ART use could also explain the

Table 1. Characteristics and Plasma Drug Concentrations of HIV-1 Infected Participants With Detectable Tenofovir

<table>
<thead>
<tr>
<th>Case</th>
<th>Testing Group</th>
<th>Gender</th>
<th>Total Study Months</th>
<th>Drug Levels by Study Month (ng/mL)*</th>
<th>HIV-1 Viral Load (copies/mL)</th>
<th>Conclusion on Drug Sharing within Study Partnership</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>1</td>
<td>Male</td>
<td>36</td>
<td>Month 0: FTC ND, TFV 71.5. Month 12: FTC NT, TFV 79.2. Month 15: FTC NT, TFV 72.7.†</td>
<td>Month 0: UD. Month 12: UD.</td>
<td>Detectable TFV only and no FTC at randomization, indicates drug could not come from study partner. Undisclosed ART at enrollment</td>
</tr>
<tr>
<td>2</td>
<td>1</td>
<td>Male</td>
<td>36</td>
<td>Month 0: FTC ND, TFV ND. Month 11: FTC NT, TFV 1.32.†</td>
<td>Month 0: 217,738. Month 11: UD.†</td>
<td>Potential PrEP sharing. Alternative explanation may be ART initiation without reporting to the study team</td>
</tr>
<tr>
<td>3</td>
<td>2</td>
<td>Female</td>
<td>36</td>
<td>Month 0: FTC 73.1, TFV 27.4. Month 6: FTC ND, TFV 56.1. Month 18: FTC ND, TFV 41.3.</td>
<td>Month 0: UD. Month 12: UD.</td>
<td>Detectable TFV and FTC, starting at randomization, indicates drug could not come from study partner. Undisclosed ART at enrollment</td>
</tr>
<tr>
<td>4</td>
<td>3</td>
<td>Female</td>
<td>36</td>
<td>Month 0: FTC ND, TFV 68.6. Month 6: FTC ND, TFV 75.2.</td>
<td>Month 0: UD. Month 12: UD.</td>
<td>Detectable TFV only and no FTC, starting at randomization, indicates drug could not come from study partner. Undisclosed ART use during follow-up</td>
</tr>
<tr>
<td>5</td>
<td>3</td>
<td>Female</td>
<td>36</td>
<td>Placebo TDF/FTC Month 0: FTC ND, TFV ND. Month 24: FTC ND, TFV 46.6 (Rerandomization).</td>
<td>Month 0: 682. Month 12: 212. Month 24: 436.</td>
<td>Detectable TFV only and no FTC indicates drug could not come from study partner. Undisclosed ART use during follow-up</td>
</tr>
<tr>
<td>6</td>
<td>3</td>
<td>Female</td>
<td>33</td>
<td>Month 0: FTC ND; TFV ND. Month 33: FTC ND, TFV 58.1.</td>
<td>Month 0: UD. Month 12: UD.</td>
<td>Detectable TFV only and no FTC, starting at randomization, indicates drug could not come from study partner. Undisclosed ART at enrollment</td>
</tr>
<tr>
<td>7</td>
<td>3</td>
<td>Female</td>
<td>24</td>
<td>Month 0: FTC ND, TFV 23.8. Month 6: FTC ND, TFV 68. Month 12: FTC ND, TFV 52.7.</td>
<td>Month 0: UD. Month 12: UD.</td>
<td>Detectable TFV, starting at randomization, indicates drug could not come from study partner. Undisclosed ART use during follow-up</td>
</tr>
<tr>
<td>9</td>
<td>3</td>
<td>Female</td>
<td>33</td>
<td>Month 0: FTC ND, TFV ND. Month 27: FTC ND, TFV ND. Month 30: FTC ND, TFV ND. Month 33: FTC ND, TFV ND, TFV 74.3.</td>
<td>Month 0: 41,350; Month 33: UD</td>
<td>Potential PrEP sharing. Alternative explanation may be ART initiation without reporting to the study team</td>
</tr>
</tbody>
</table>

*Plasma tenofovir >0.31 ng/mL is consistent with dosing within the last week, >10 ng/mL is consistent with dosing in the previous 2-3 days, and > 40 ng/mL is the lower bound of the 95% confidence interval for directly observed dosing at steady state. 19
†Indicates plasma collected at an unannounced home visit.
ART, antiretroviral therapy; FTC, emtricitabine; TDF or TNF, tenofovir; ND, drug levels not detected (< 0.31 ng/mL); NT, not tested; UD, HIV-1 RNA viral load undetectable (< 80 copies/mL).
3 cases of potential PrEP sharing we observed in the current study.

Plasma was not always available at the exact same time point for both members of the partnership because HIV-1 uninfected participants were seen monthly and HIV-1 infected partners were seen quarterly. It is possible that instances of time-limited drug sharing could have been undetected. Our study population of mutually disclosed HIV-1 serodiscordant couples enrolled in a clinical trial places some limitations on the generalizability of our findings. All participants in the Partners PrEP Study received frequent adherence counselling, including the potential for ART drug resistance if the HIV-1 infected study partner took study medication, counselling which is more intensive than in an implementation setting. Nevertheless, medication sharing was feasible in our study population.

HIV-1 serodiscordant couples have been identified as a priority for PrEP implementation. Using prospectively collected data, including self-report and objective testing of drug levels in 4 groups that represent randomly selected specimens and subpopulations with low adherence to study medication, we comprehensively assessed PrEP sharing within this population. Re-randomization of placebo arm participants created a unique opportunity to assess drug sharing when all HIV-1 uninfected partners received active PrEP, conditions that could have incentivized drug sharing. Importantly, we did not find an increase in drug sharing post-unblinding.

**CONCLUSIONS**

Self-reported drug sharing in the Partners PrEP Study was extremely rare. Objective plasma tenofovir testing supports the self-reported data and indicates that sharing PrEP within HIV-1 serodiscordant couples was uncommon. These results suggest that it is unlikely for medication sharing to be a serious limitation for PrEP delivery to HIV-1 uninfected members of HIV-1 serodiscordant couples.

**ACKNOWLEDGMENTS**

The authors thank the couples who participated in this study and the teams at the study sites for work on data collection and management.

**Partners PrEP Study Team.**

University of Washington Coordinating Center and Central Laboratories: Connie Celum (principal investigator, protocol co-chair), J.M.B (medical director, protocol co-chair), Deborah Donnell (protocol statistician), Robert W. Coombs, Lisa Frenkel, Craig W. Hendrix, Jairam Lingappa, M. Juliana McElrath.


Data management was provided by DF/Net Research, Inc. (Seattle, USA) and site laboratory oversight was provided by Contract Laboratory Services (University of the Witwatersrand, Johannesburg, South Africa). Study medication donated by Gilead Sciences, Inc.

**REFERENCES**


