The use of preexposure treatments for HIV prophylaxis

Adrian Majid
Robert R Redfield
Bruce L Gilliam
Institute of Human Virology,
University of Maryland School of Medicine, Baltimore, MD, USA

Abstract: Infection with human immunodeficiency virus remains a global concern with a significant number of incident infections still reported worldwide. The use of prophylaxis prior to exposure to the virus to prevent infection has been a growing area of recent research. Results in nonhuman primates and clinical trials in high-risk patient populations using preexposure prophylaxis have shown promising results in terms of efficacy and safety, especially relating to oral preexposure prophylaxis. The potential use of oral antiretroviral agents traditionally used for human immunodeficiency virus treatment as prophylaxis raises interesting considerations, such as the best agents available for such a role, long-term safety in healthy individuals, and the potential development of resistance to these agents should infection occur. From a public health perspective, the cost-effectiveness of implementing this preventive strategy has not been fully defined at this point in time.

Keywords: preexposure prophylaxis, human immunodeficiency virus, tenofovir, maraviroc

Introduction
Human immunodeficiency virus (HIV) remains a significant public health challenge. There are currently more than 33 million people living with acquired immune deficiency syndrome (AIDS) worldwide. Despite increasing access to antiretroviral drugs, the Joint United Nations Program on HIV/AIDS reported a total of 2.7 million new infections worldwide in 2008. Efforts to curb transmission by treating HIV-infected patients in serodiscordant couples and prophylaxis to prevent mother-to-child transmission have likely played a significant role in decreasing incident infections. But these measures, along with longstanding public health measures, have not been able to prevent incident infections. In the United States, for example, there are an estimated 56,000 cases of incident HIV infections per year. Men who have sex with men (MSM) account for 53% of these cases.

In order to decrease these incident infections, new strategies for preventing HIV transmission, especially among high-risk groups, have emerged. Preexposure prophylaxis (PreP), one of these approaches, involves the use of topical or oral agents in HIV-uninfected individuals prior to exposure to the virus in order to prevent HIV acquisition. In addition, PreP may potentially play a role in attenuating the natural history of HIV disease progression in patients who become infected, reducing morbidity and decreasing infectiousness to others.

In this review, the current knowledge regarding the use of topical and oral agents for PreP will be presented. The ideal pharmacokinetics for efficacy of PreP agents,
and the results of preclinical animal models and human clinical trials of PreP will be discussed. Additionally, the debate regarding the long-term safety of PreP, the implications of developing resistant HIV on PreP, and the overall cost-effectiveness of implementing PreP as a prevention strategy will be explored.

Microbicides

Microbicides are topical products that can be applied to either the vaginal or rectal mucosa to prevent HIV transmission. The idea for microbicides developed as a means by which women could have control over their risk of HIV infection and potentially prevent other sexually transmitted infections.\(^\text{5,9}\) Research in the field grew as the percentage of women infected with HIV increased, especially in Sub-Saharan Africa, where 67% of the HIV-infected individuals worldwide live.\(^\text{1}\)

Unfortunately, over the past decade, research into various microbicides has not yielded compelling data for their efficacy. Clinical trials of surfactants, agents that disrupt the cell membranes of viruses and bacteria, and polyanions (eg, Carraguard\(^\text{®}\) cellulose sulfate, PRO 2000) showed promise in in vitro and in animal studies,\(^\text{10–13}\) but did not show statistically significant differences between rates of HIV acquisition in clinical trials.\(^\text{14–19}\) In fact, nonoxynol-9 and cellulose sulfate were associated with an increased risk of HIV acquisition in women, likely due to a rapid and sustained reduction in transepithelial resistance.\(^\text{20,21}\)

Due to the lack of efficacy of these microbicides, focus has shifted toward agents that might be more specific for HIV. Tenofovir (TDF), a nucleotide analog that blocks the reverse transcription of HIV in the host cell, has been studied as a microbicide in a 1% gel form, both by itself and in combination with emtricitabine (FTC), another nucleoside analog that comes in a 5% gel.\(^\text{20}\) These agents have been chosen for their long half-lives (TDF: \(\sim 40\) hours, FTC: \(\sim 40\) hours), allowing them to persist in tissues long after extracellular concentrations decline.

Studies of macaques exposed to simian immunodeficiency virus (SIV) and simian-human immunodeficiency virus (SHIV) vaginally and rectally have been used as animal models for studying HIV acquisition. One study revealed that a 1% TDF gel applied rectally to rhesus macaques offered a statistically significant partial efficacy against SIV.\(^\text{22}\) In one study, the vaginal application of 1% TDF alone or in combination with 5% FTC two times per week fully protected macaques from a total of 20 exposures to homologous SHIV.\(^\text{23}\) These studies are further outlined in Table 1.

Building on these animal models, the Center for the AIDS Programme of Research in South Africa 004 clinical study,\(^\text{24}\) a double-blind randomized controlled trial of about 900 South African HIV-uninfected higher-risk women ages 18–40, compared rates of HIV acquisition with TDF gel versus placebo gel. The HIV incidence in the TDF gel arm was 5.6 per 100 women years compared with 9.1 per 100 in the placebo arm, which was statistically significant. In those with \(>80\)% adherence, the HIV incidence was 54% lower in the TDF gel arm. Although there was documented efficacy, low rates of adverse reactions, and no documentation of resistance to TDF developing in women who acquired HIV, rates of HIV transmission were still high. Therefore, this strategy needs to be used in conjunction with other preventive strategies, such as condoms, in order for it to be fully beneficial.

Other antiretroviral microbicides currently in multiple phase I and II clinical trials are dapivirine and UC 781, two non-nucleoside reverse-transcriptase inhibitors (NNRTIs).\(^\text{25,26}\) From data available, both seem to attain high levels in cervicovaginal fluid, although their efficacy remains to be evaluated in further clinical trials. The safety of these agents in the more delicate epithelium of the rectum is now being studied. A phase I trial of UC781 revealed no significant adverse events and no significant plasma levels in sexually abstinent men and women applying the microbicide rectally during single and 7-day exposures.\(^\text{27}\) Phase III clinical trials evaluating the efficacy of the dapivirine ring, IPM 009A and IPM 009B, will commence shortly.\(^\text{28}\)

Antagonists to C–C chemokine receptor type 5 (CCR5), the chemokine co-receptor for HIV entry into the cell, have also been effective as microbicides in macaque studies,\(^\text{29,30}\) as well as in humanized monoclonal mouse models.\(^\text{31}\) See Table 1 for further details of the macaque studies.

From the data available, the lack of systemic absorption of antiretroviral microbicides appears to prevent problems with resistance developing in patients that are using them while unknowingly infected, and safety trials have not revealed any significant adverse reactions.\(^\text{14}\) However, microbicides do not protect against other sexually transmitted infections, which can also facilitate HIV acquisition.\(^\text{32}\) Therefore, antiretroviral microbicides likely will be most effective when used with barrier protection. The use of such microbicides to prevent rectal transmission in both men and women is still under investigation.

Microbicides by themselves are not likely to have a significant impact on decreasing the incidence of HIV globally. Poor adherence, lower efficacy rates and increased cost
Table 1 Summary of macaque trials of preexposure prophylaxis

<table>
<thead>
<tr>
<th>Preexposure prophylaxis regimen</th>
<th>Dosing regimen</th>
<th>Virus</th>
<th>Mode of viral challenge</th>
<th>Efficacy at preventing infection</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Microbicides</strong></td>
<td></td>
<td></td>
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</tr>
<tr>
<td>TDF 1% gel 2 hours before exposure$^{22}$</td>
<td>1 preexposure dose</td>
<td>SIV</td>
<td>Rectal (high dose)</td>
<td>89%*</td>
</tr>
<tr>
<td>TDF 1% gel with FTC 5% gel 30 minutes before exposure$^{23}$</td>
<td>20 gel applications prior to exposure</td>
<td>SHIV</td>
<td>Vaginal (2 times per week for 20 challenges)</td>
<td>100%*</td>
</tr>
<tr>
<td>Topical CCR5 antagonist 30 minutes before exposure</td>
<td>1 preexposure dose</td>
<td>SHIV</td>
<td>Vaginal single challenge</td>
<td>66%–100%* with CCR5 antagonist and other entry inhibitors</td>
</tr>
<tr>
<td>Topical maraviroc (CCR5 antagonist) topical to vaginal 30 minutes before exposure$^{30}$</td>
<td>1 preexposure dose</td>
<td>SHIV</td>
<td>Vaginal single challenge</td>
<td>75%–86%*</td>
</tr>
<tr>
<td><strong>Systemic agents</strong></td>
<td></td>
<td></td>
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<td></td>
</tr>
<tr>
<td>TDF 20 mg/kg subcutaneously 48 hours before exposure$^{44}$</td>
<td>Daily × 4 weeks</td>
<td>SIV</td>
<td>Intravenous × 1</td>
<td>100%*</td>
</tr>
<tr>
<td>Oral TDF 22 mg/kg 2 hours prior to exposure$^{45}$</td>
<td>Daily TDF vs weekly TDF × 36 weeks</td>
<td>SHIV</td>
<td>Rectal (weekly × 14 weeks)</td>
<td>No statistically significant difference between the groups. Infection appeared to be delayed in treated macaques</td>
</tr>
<tr>
<td>Oral TDF 22 mg/kg 7 days prior to exposure$^{46}$</td>
<td>Daily until 28 days after inoculation</td>
<td>SHIV</td>
<td>Rectal (single high-dose exposure)</td>
<td>40%</td>
</tr>
<tr>
<td>(1) 20 mg/kg of FTC subcutaneously</td>
<td>Daily regimens given 7–9 days prior to exposure up 28 days after exposure</td>
<td>SHIV</td>
<td>Rectal (weekly × 14 weeks)</td>
<td>33% and 67%* in groups 1 and 2. 100%* in groups 3 and 4</td>
</tr>
<tr>
<td>(2) Oral FTC/TDF (20 and 22 mg/kg, respectively)</td>
<td>Weekly regimen with weekly dosing with second dose</td>
<td>SHIV</td>
<td>Rectal (weekly × 14 weeks)</td>
<td>33% and 67%* in groups 1 and 2. 100%* in groups 3 and 4</td>
</tr>
<tr>
<td>(3) FTC subcutaneously (20 mg/kg) and TDF (22 mg/kg) subcutaneously ((1), (2), and (3) given 7–9 days prior to exposure)</td>
<td>24 hours after exposure</td>
<td>SHIV</td>
<td>Rectal (weekly × 14 weeks)</td>
<td>33% and 67%* in groups 1 and 2. 100%* in groups 3 and 4</td>
</tr>
<tr>
<td>(4) Subcutaneous FTC (20 mg/kg) and TDF (22 mg/kg) given weekly 2 hours before exposure$^{47}$</td>
<td>1 preexposure dose and second postexposure dose 2 hours after</td>
<td>SHIV</td>
<td>Rectal (weekly × 14 weeks)</td>
<td>50%–83%*</td>
</tr>
<tr>
<td>Intermittent tenofovir–emtricitabine before exposure (1, 3, or 7 days before exposure)$^{48}$</td>
<td>Second postexposure dose 2 hours after</td>
<td>SHIV</td>
<td>Rectal (weekly × 14 weeks)</td>
<td>100%*</td>
</tr>
<tr>
<td>(1) Oral FTC/TDF 3 days prior to exposure$^{49}$</td>
<td>SHIV – FTC resistant isolate with M184V mutation</td>
<td>SHIV</td>
<td>Vaginal single challenge</td>
<td>45%–56%, No statistically significant difference in infection rate between groups</td>
</tr>
<tr>
<td>Oral CCR5 antagonist either given on the day of challenge or 4 days before challenge$^{50}$</td>
<td>Daily until 10 days after challenge</td>
<td>SHIV</td>
<td>Vaginal single challenge</td>
<td>45%–56%, No statistically significant difference in infection rate between groups</td>
</tr>
</tbody>
</table>

Note: *statistically significant.

Abbreviations: TDF, tenofovir; SIV, simian immunodeficiency virus; FTC, emtricitabine; SHIV, simian human immunodeficiency virus; CCR5, C–C chemokine receptor type 5.
compared to condoms are significant drawbacks. In the Center for the AIDS Programme of Research in South Africa 004 trial, 41% of women exhibited low adherence, defined as using the gel <50% during intercourse. In day-to-day use outside of a clinical trial, one would expect this number to be even lower, thereby compromising the benefits of this intervention.

**Oral prophylaxis**

**Theoretical considerations with preexposure prophylaxis**

The use of oral antiretroviral drugs as PreP has a foundation in existing knowledge. These agents have been used in preventing perinatal transmission of HIV and acquisition of the virus after percutaneous occupational exposures.

It is critical to understand the pathogenesis of HIV in order to understand why PreP works. After exposure to HIV, systemic infection usually does not occur immediately, but is delayed 1–3 days. During this period, a small population of “founder cells” appear to be responsible for spreading the infection to a cluster of nearby cells, which then leads to viral dissemination and systemic viremia as early as 5 days after exposure. The administration of antiretrovirals for PreP when populations of founder cells are being established may disrupt the infection cycle and help the immune system eradicate the virus.

The choice of an ideal agent for oral PreP is dependent on multiple factors including: (1) whether it acts pre-integration or post-integration (2) the pharmacokinetics of the agent, including its ability to achieve high concentrations in cervicovaginal fluid and the rectum, (3) barriers to resistance, (4) pill burden, and (5) safety and tolerability.

An agent with activity against HIV prior to its establishment in the host cell (ie, pre-integration) would be ideal for use as PreP because it would prevent the establishment of HIV infection in cells. Protease inhibitors are post-integration agents, and their use in this field will likely be limited, although currently there is no available data to support this assertion.

Additionally, the optimal PreP should have ideal pharmacokinetic properties, including rapid distribution and long half-life into tissues. FTC and TDF, for example, both have long intracellular half-lives (40–100 hours), with rapid distribution into tissues, and achieve high concentrations in the cervicovaginal fluid. Studies measuring levels of antiretrovirals in cervicovaginal fluid have shown that NRTIs achieve the highest levels when compared to protease inhibitors and NNRTIs. More recent data has revealed high concentrations of both maraviroc, a CCR5 antagonist, and raltegravir, an integrase inhibitor, in both the male and female genital tracts. Maraviroc has also achieved high rectal concentration in healthy men. See Figure 1 for a comparison of these agents and their concentration in the genital tract relative to blood.

Of the antiretroviral agents mentioned, NRTIs and CCR5 antagonists are the most appealing for PreP for the reasons already cited, as well as for their high barrier to resistance and relatively good safety profile. Maraviroc is currently a twice-daily medication, which is a drawback that might hinder larger trials looking at its efficacy for PreP. However, it is unclear whether or not maraviroc could be given once daily for prevention instead of treatment and this needs further investigation.

![Figure 1](image-url)  
**Figure 1** Concentration of select oral antiretrovirals in the genital tract of men and women relative to blood. Adapted from references 35–40.

**Notes:** In general, the nucleoside reverse transcriptase inhibitors seem to concentrate better in the genital tract of men and women than protease inhibitors and non-nucleoside reverse transcriptase inhibitors (the latter two are excluded from this graph). Some more recent data, although there was significant variability between subjects, shows that raltegravir and maraviroc each can achieve significant levels as well.

*Abacavir achieves lower levels in the female genital tract relative to blood (~40%).
study. Given that CCR5-tropic viruses represent the majority of transmitted viruses, maraviroc may have a unique niche in PreP since it does not form the backbone of current antiretroviral regimens as do the NRTIs.

Once-a-day integrase inhibitors, which block viral integration into the host cell, have been developed. Despite their pre-integration efficacy, integrase inhibitors are not ideal, given their low barrier to resistance, which makes their long-term efficacy questionable. The pharmacokinetics of these agents have also not been as well studied.

For these reasons, the majority of animal studies and clinical trials studying PreP have looked at agents like TDF or FTC/TDF, although oral maraviroc may have a potential role for prophylaxis, which will be discussed.

**Preclinical animal studies of PreP**

Both macaque and mouse models have been used to study the potential efficacy of PreP, mainly using TDF and FTC. In macaque monkeys, SIV and SHIV have been used as nonhuman primate models to mimic human immunodeficiency virus. SHIV more closely mimics the various aspects of human infection.

Data from the macaque monkey trials with oral antiretrovirals is summarized in Table 2. Initially, studies in SIV focused on a single agent, TDF, which showed efficacy at preventing SIV but, at oral doses comparable to human doses, only revealed partial efficacy at preventing SHIV infection. This prompted many of the most recent animal studies and human clinical trials to look at the combination of FTC/TDF.

In general, the macaque studies have shown that the combination of FTC/TDF can be very effective in preventing the acquisition of SIV and SHIV after oral, rectal, or vaginal exposure. Increased efficacy seems to correlate with increased dosages of antiretrovirals. One study utilizing intermittent weekly dosing of subcutaneous TDF and FTC before and after viral exposure showed equivalent protection from SHIV infection as provided by the same medications before and after viral exposure showed equivalent protection from SHIV infection as provided by the same medications.

Humanized mouse models that harbor HIV-susceptible human cells have also been used to examine PreP. Studies using this model have demonstrated the efficacy of TDF and FTC/TDF in preventing vaginal, rectal, and intravenous transmission of HIV. One study used this model and demonstrated that oral raltegravir and maraviroc protected against a vaginal HIV-1 challenge.

**Clinical studies**

The data from animal studies, as well as the success of antiretrovirals in preventing mother-to-child transmission and postexposure prophylaxis after occupational exposure, has prompted many clinical trials looking at the use of PreP in high-risk HIV populations. To date, these trials have all evaluated the efficacy of either TDF or FTC/TDF.

The first published clinical report of PreP compared daily TDF versus placebo in 936 West African women from Ghana, Cameroon, and Nigeria in preventing HIV transmission via vaginal intercourse. Two women in the TDF group and six women in the placebo group became infected, but this result was not statistically significant. The study terminated early due to the closure of two of the clinical sites, which limited the planned person-years of follow-up and the power of the study.

The Preexposure Prophylaxis Initiative (iPrEx) study, a subsequent landmark multinational study, evaluated the safety and efficacy of FTC/TDF in 2499 HIV seronegative men or transgender women who have sex with men, a group with a high number of incident infections worldwide. One hundred patients became infected during follow-up, 36 in the FTC/TDF group and 64 in the placebo group, indicating a statistically significant 44% prevention efficacy. In subgroup analyses, efficacy was higher among subjects that reported that they had previously had unprotected receptive anal intercourse and those that reported greater than 90% compliance with pill use. Of those patients in the FTC/TDF group that became infected with HIV, only three had detectable levels of drug in their blood, none with cell-associated drug levels higher than the median levels for seronegative controls.

Preliminary data from additional clinical trials has yielded conflicting data in other high-risk groups. The TDF2 study followed 1200 heterosexual men and women in Botswana randomized to receive FTC/TDF versus placebo, and found a significantly decreased risk of HIV acquisition (nine patients in the FTC/TDF arm as opposed to 24 patients in placebo). As women are one of the main target groups for PreP, it is important to note that the protective effects of FTC/TDF were seen in both women and men, although the study was underpowered to evaluate sex-based differences. These results are in contrast to the FEM-PrEP trial, a phase III clinical trial of oral FTC/TDF in nearly 2000 women at high risk for HIV infection.
<table>
<thead>
<tr>
<th>Trial</th>
<th>Study population</th>
<th>Preexposure prophylaxis</th>
<th>Current status</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bangkok Tenofovir Study (CDC)</td>
<td>2400 IVDU in Bangkok, Thailand</td>
<td>Oral daily TDF</td>
<td>Fully enrolled. Trial results expected in 2012</td>
</tr>
<tr>
<td>iPrEx Rollover Study (NIH, BMGF)</td>
<td>2500 MSM in multiple countries</td>
<td>Oral daily FTC/TDF in all HIV-negative patients from original iPrEx</td>
<td>44% reduced risk of infection in FTC/TDF group. Rollover study results expected early 2013</td>
</tr>
<tr>
<td>TDF2 (CDC)</td>
<td>1200 heterosexual men and women in Botswana</td>
<td>Oral daily FTC/TDF</td>
<td>TDF with 62% protective efficacy. Full data awaiting publication</td>
</tr>
<tr>
<td>Partners PreP (BMGF)</td>
<td>4700 serodiscordant heterosexual couples in Kenya and Uganda</td>
<td>Oral daily TDF or oral FTC/TDF</td>
<td>TDF with 62% protection efficacy and FTC/TDF with 73% protection efficacy. Full results expected 2013</td>
</tr>
<tr>
<td>VOICE (MTN 003)</td>
<td>4200 heterosexual women in various countries in Africa</td>
<td>Topical TDF vaginal gel or oral TDF or oral FTC/TDF</td>
<td>Results expected 2013</td>
</tr>
<tr>
<td>PreP in Young MSM (ATN 082)</td>
<td>99 young MSM</td>
<td>Oral FTC/TDF</td>
<td>Data collected. Analysis ongoing.</td>
</tr>
<tr>
<td>ADAPT (HPTN 067)</td>
<td>360 participants (MSM and heterosexual women) in South Africa and Thailand</td>
<td>Oral FTC/TDF (daily dosing, time-driven dosing, event-driven dosing)</td>
<td>Enrolling (phase II trial)</td>
</tr>
<tr>
<td>HPTN 069</td>
<td>400 MSM in the United States</td>
<td>MVC or MVC/FTC or MVC/TDF or FTC/TDF</td>
<td>In development (phase II trial)</td>
</tr>
<tr>
<td>IPERGAY (ANRS)</td>
<td>300 MSM in France and Canada</td>
<td>Oral FTC/TDF with intermittent dosing</td>
<td>Enrolling</td>
</tr>
</tbody>
</table>

**Notes:** *Vaginal gel arm discontinued early due to review of interim data showing predicted lack of efficacy compared to placebo; **TDF arm discontinued early due to review of interim data showing predicted lack of efficacy compared to placebo.

**Abbreviations:** CDC, Centers for Disease Control; IVDU, intravenous drug user; TDF, tenofovir; NIH, National Institute of Health; BMGF, Bill and Melinda Gates Foundation; MSM, men who have sex with men; FVC, emtricitabine; VOICE, Vaginal and Oral Interventions to Control the Epidemic; MTN, Microbicide Trials Network; ADAPT, Alternative Dosing to Augment PrEP pill-Taking; ATN, Adolescent Trials Network; HPTN, HIV Prevention Trials Network; ANRS, Agence Nationale de Recherches sur le Sida et les Hepatites Virales.
This study was ended prematurely early in April of 2011 after a preliminary review of the data revealed equal rates of infections in the TDF and placebo groups. The study was deemed unlikely to be able to demonstrate efficacy if continued, and the study is being reviewed to assess the reasons for this surprising result.

Another phase III trial, the Partners PreP study, compared TDF, FTC/TDF, and placebo in preventing HIV transmission in HIV-discordant heterosexual African couples. Both the TDF and FTC/TDF arms had a statistically significant decreased incidence of HIV infections versus placebo, with a 62% HIV protection efficacy noted in the TDF arm and a 73% protection efficacy noted in the FTC/TDF group.

Like the Partners study, the results of HPTN 052, although not a PreP trial, support the efficacy of early antiretroviral drugs in reducing rates of sexual transmission of HIV-1 among serodiscordant couples. In this multinational study of 1763 couples, the HIV-infected partners were randomized to receive early antiretroviral therapy or delayed treatment once CD4 criteria were met. Only one of the 28 overall partner-linked infections occurred in the group that was randomized to receive early antiretroviral therapy, which was statistically significant.

The discrepancy between the FEM-PreP trial and the other clinical trials in African countries showing the clinical efficacy of FTC/TDF is worth discussing. The most plausible explanation is that the subjects in FEM-PreP had lower adherence than they self-reported. The measurement of serum levels and the tissue concentration of these drugs are a better means of assessing adherence than self-reporting. There may be other factors that we do not yet understand in the FEM-PreP population that, even with reasonable adherence, might contribute to decreased levels of antiretrovirals systemically and, more importantly, in the cervicovaginal fluid. A recent study, for example, revealed significantly lower levels of tenofovir diphosphate in the vaginal tissue of patients receiving oral tenofovir as opposed to vaginally applied tenofovir. Achieving levels of active drug in the cervicovaginal tissue is a valid concern, although it does not fully explain the discrepancy between the various studies. The data from the FEM-PreP trial is being analyzed by its investigators to better try to identify the reason for the discordant results in this study compared to others.

With a small number of clinical trials completed showing the efficacy and relative safety of oral PreP with FTC/TDF in high-risk groups for infection with HIV, further studies are in progress. These include studies of MSM, high-risk women, and injection drug users, and evaluate strategies for intermittent PreP. Information on phase II and phase III clinical trials of oral PreP are found in Table 2. One of these trials, the Vaginal and Oral Interventions to Control the Epidemic (VOICE) study, aimed to compare the efficacy of oral tenofovir, oral truvada, and vaginal tenofovir as PreP to each other and to placebo. During interim reviews by an independent safety board, the trial discontinued both the oral tenofovir arm and the vaginal gel arm since review of the early data indicated lack of efficacy of either compared to placebo. Potential explanations for the lack of efficacy of oral tenofovir include the varying degrees of active tenofovir drug in the cervical tissue of women when administered in an oral formulation and intermittent or poor adherence.

Mathematical models have examined PreP and seen the potential for a significant public health benefit to its implementation, both in the United States and developing countries. With a product demonstrating 90% efficacy and with 75% coverage of the general population, models forecast a 74% decline in the number of cumulative HIV infections in 10 years. These models are encouraging, but do not factor in some of the important considerations with regard to safety, the development of resistance to PreP medications, and potential changes in sexual-risk behavior over time, considerations that will be discussed.

### Considerations for the long-term use of PreP

#### Safety

Most of the clinical trials looking at the use of FTC/TDF for PreP seem to indicate that use for over a year seems relatively safe. The CAPRISA trial demonstrated that no statistically significant adverse reaction was seen more frequently in the TDF gel group compared to placebo. With regards to oral PreP, the iPreEx study revealed that the only significant adverse reactions encountered by subjects were nausea and weight loss, with a trend observed of elevated creatinine levels in the FTC/TDF group that was not clinically significant. It should be noted, however, that the majority of this study population was less than 40 years old. As with populations that tend to make up clinical trials, one could also presume that they would have been a healthier population than the real-world population that would be using these medications. Patients using PreP, therefore, may encounter more side effects or drug–drug interactions than could be expected within the confines of a study. Another important consideration is that we have extensive knowledge about the side-effect profile of FTC/TDF, but there is the potential that medication use in a patient population with an intact immune system could lead
to unforeseen side effects, as we have seen previously with hepatotoxicity developing in patients with high CD4 counts starting nevirapine.65

Safety information available from the Partners PrEP study, the study of West African women, the United States Extended Safety Trial, and the TDF2 study of heterosexual men and women in Botswana all seem to demonstrate short-term safety of PreP. The Partners PreP prophylaxis trial, for example, only showed increased diarrhea with active agents.61 According to the Centers for Disease Control, preliminary data from the US Extended Safety Trial (CDC 4323) of 400 MSM receiving oral tenofovir as PreP also did not raise significant safety concerns.66

None of the longer-term cumulative toxicities have been addressed by the clinical trials to date. The trend to an elevated creatinine in the iPrEx with less than perfect adherence study raises concerns for cumulative toxicities including renal failure, tubulointerstitial disease, electrolyte derangements, bone disease, and potential flares of hepatitis B that should not be underestimated. Recently presented and published data has demonstrated reductions in bone mineral density in seronegative men participating in clinical trials with TDF and FTC/TDF.67,68 In the tenofovir PreP study among MSM in San Francisco, a statistically significant decline in bone mineral density in the femoral neck and hip was seen compared to placebo. The implications for bone health over longer periods of time in a PreP setting need to be studied further, but one would expect further deteriorations in both health and clinically significant fractures over time.

One area of safety that has been discussed, but not studied extensively, is the impact of the use of TDF and FTC/TDF in cases of hepatitis B co-infection where the patient is unaware of their hepatitis status. In the TDF trial of West African women, this side effect was not seen. But given the activity of TDF and FTC/TDF against hepatitis B, the incidence of hepatitis B flares, and the development of hepatitis B resistance to these medications, further investigation is required.

Resistance

One of the biggest concerns with HIV PreP is the development of resistance to these medications in those who become infected. Especially in the developing world, resistance to first- and second-line regimens has significant implications for treatment options for HIV-infected patients. FTC and TDF are two of the most potent and well-tolerated medications used worldwide for HIV treatment, and we cannot lose the ability to prescribe these medications for the treatment of HIV.

FTC has signature resistance mutations at codon 184 with a single nucleotide change that result in amino acid changes from methionine to isoleucine (M184I) or valine (M184V).69,70 Either of these mutations confers a high level of resistance to either FTC or lamivudine (3TC).71 Monotherapy studies have shown that M184V mutations can develop within approximately 15 days of treatment with lamivudine.70 Despite the high level of resistance to these medications, studies have shown that their continuation in regimens does confer a benefit with regard to viral suppression, likely due to decreased replicative capacity of the virus.71,72 Additionally, hypersusceptibility to other components of an antiretroviral regimen can occur with these mutations (eg, TDF, zidovudine [AZT]).73 One animal macaque study exemplified this point by showing that rectal transmission of an FTC-resistant isolate with an M184V mutation did not occur in macaques receiving prophylaxis with FTC/TDF, although it did occur in all of the controls.49

There are two specific mutations that influence the efficacy of tenofovir: K65R and K70E/G. The K65R mutation involves a change at the second nucleotide codon position from lysine to arginine and causes intermediate-level resistance to tenofovir.74,75 There is some data to support that this mutation is more likely to occur in patients with subtype C of HIV, the prevalent subtype seen in Sub-Saharan Africa and India, that are not fully suppressed on their antiretroviral regimen.76,77 Single nucleotide alterations in codon 70 lead to an amino acid shift from lysine to glutamate and a second transition required to yield glycine. This produces decreased susceptibility to tenofovir and other antiretrovirals, including FTC. Mutations associated with TDF do not occur as rapidly as the 184V/I with FTC, with monotherapy studies predicting development of these mutations in the 4- to 8-week timeframe.78,79

Dual therapy with FTC/TDF in patients who are intermittently adherent to these medications will likely lead to the development of an M184V mutation within several weeks, with almost all patients exhibiting resistance within a year.73,80 In the iPrEx study, two of the men had seronegative HIV infection at the time they started to receive active drug, and both were noted to have M184V mutations by week 4.35 In the TDF2 study, one participant had unrecognized HIV infection at the time they started to receive active drug, and both were noted to have M184V mutations by week 4.34 In an area like southern Africa, where the C subtype of HIV predominates, one might expect higher rates of TDF resistance to develop. It will, therefore, be interesting to see the resistance mutations that develop with longer-term follow-up, especially in the trials based in Sub-Saharan Africa.
An M184V mutation in a patient in the developing world can have significant implications for treatment, since most first-line regimens involve two NRTIs (usually a combination of TDF or AZT and lamivudine or FTC) with an NNRTI. Although the M184V virus would exhibit hypersusceptibility to TDF or AZT and would be less fit for replication, a regimen based on one active NRTI and an NNRTI, given the low barrier to resistance of the NNRTI, runs the risk of treatment failure. One mathematical model looking to predict the effect of PreP on the HIV epidemic in MSM in San Francisco forecasts decreased transmission of HIV, but an increased proportion of new infections caused by resistant strains.81

Outside a clinical trial, one could expect more resistant mutations to develop due to poorer adherence and compliance with follow-up. In order to avoid the consequences for resistance in patients on a partially suppressive antiretroviral regimen with undiagnosed infection, regular HIV testing should be linked to the use of PreP. The development of an infrastructure to provide HIV testing at least every 6 months in subjects receiving PreP is critical to prevent resistance with implementation of PreP.

Cost-effectiveness and considerations for the implementation of PreP

The high cost of implementation of PreP is an important consideration in its potential adoption as a prevention strategy in high-risk populations. The wholesale cost of FTC/TDF is about US$900 per month, not factoring in other costs, including personnel and infrastructure costs, counseling, routine testing, and surveillance for adverse reactions.82 Using data from the iPrEx study, it is estimated that about 44 people would have to receive PreP to prevent one infection, which reflects a cost of over $500,000 over a 1-year period.83 This cost is about 20 times the amount that it takes to treat someone with HIV for a year. In this model, PreP is not cost-effective.

It is also unlikely that these medications for PreP will be covered by private insurance, which leaves public financing as the only option for implementing a PreP program. With the current economic downturn, public funding may not be readily available. Some state drug assistance programs set up to treat individuals already infected with HIV are running out of resources, with large waiting lists for antiretroviral treatment. In resource-limited settings, it would seemingly be more difficult to set up the infrastructure for a PreP program and to garner the funds for such an endeavor when there are not sufficient funds to treat individuals already infected with HIV. Currently, the World Health Organization does not even mandate treatment for HIV-infected partners in serodiscordant couples, despite the results of the HIV Prevention Trials Network 052 study which demonstrated that treatment of the HIV-infected partner in serodiscordant couples has a significant impact on decreasing transmission of HIV.59

Aside from the lack of cost-effectiveness of a PreP strategy, there are other important public health and population considerations for implementation. This includes the optimal target group for treatment (high-risk MSM participating in receptive anal intercourse, female and male sex workers). Which practitioners would be in a position to offer PreP? What training would they require? Also, one would need to decide whether to use a daily pill strategy or an intermittent-pill strategy, the latter model showing efficacy in animal models but not yet in humans.

PreP may have implications on an individual level as well. Concerns have been raised that the availability of PreP may lead to more risky sexual practices through behavioral disinhibition, also known as risk compensation.84 The concept is that people might feel protected by PreP and stop using barrier methods of protection, which have been proven to be more efficacious. Risk compensation has not yet been seen in placebo-controlled clinical trials, although some data exists that it is a genuine concern in high-risk populations.55 The attitudes of high-risk individuals to PreP, actual levels of adherence, and the role of risk compensation need to be studied further. In the clinical trials of PreP already completed, other preventive strategies for decreasing HIV transmission are reinforced to subjects as part of the study protocol. It is not yet clear whether or not these same measures would be built into a program for PreP in a real-world setting.

Future areas of interest

The role of PreP as a strategy that can be widely implemented for HIV prevention remains to be determined. From a research standpoint, there are other drugs and molecules that could be and are being evaluated in clinical studies for PreP, including CCR5 antagonists. Biologic molecules that have an effect on CCR5 expression could potentially be another area of interest that will help avoid the concerns about antiretroviral drug resistance.

Despite the data that has demonstrated the efficacy of PreP in higher-risk populations, there are many unknown factors with regards to longer-term use of oral antiretrovirals for PreP, including safety and resistance concerns. Equally important, there are significant public health concerns relating to the implementation of such a strategy. PreP needs to
be evaluated alone and in combination with other approaches such as universal testing and early treatment of HIV infections to determine the best means to decrease the incidence of HIV infections worldwide.

**Disclosure**

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