Adolescent pre-exposure prophylaxis for HIV prevention: current perspectives

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Abstract: Adolescents are a critical population that is disproportionately impacted by the HIV epidemic. More than 2 million adolescents between the age group of 10 and 19 years are living with HIV, and millions are at risk of infection. HIV risks are considerably higher among girls, especially in high-prevalence settings such as eastern and southern Africa. In addition to girls, there are other vulnerable adolescent subgroups, such as teenagers, who use intravenous (IV) drugs, gay and bisexual boys, transgender youth, male sex workers, and people who fall into more than one of these categories. Pre-exposure prophylaxis (PrEP) is a new intervention for people at high risk for acquiring HIV, with an estimated HIV incidence of >3%. Recent data from trials show evidence of the efficacy of PrEP as a powerful HIV prevention tool in high-risk populations, including men who have sex with men, HIV-1-serodiscordant heterosexual couples, and IV drug users. The reported efficacy in those trials of the daily use of oral tenofovir, alone or in combination with emtricitabine, to prevent HIV infection ranged from 44% to 75% and was heavily dependent on adherence. Despite the proven efficacy of PrEP in adult trials, concerns remain about its feasibility in real-life scenarios due to stigma, cost, and limited clinician experience with PrEP delivery. Recent studies are attempting to expand the inquiry into the efficacy of such HIV prophylaxis approaches in adolescent populations, but there are still many gaps in knowledge, and no country has yet approved it for use with adolescents. The aim of this review was to identify and summarize the evidence from studies on PrEP for adolescents. We have compiled and reviewed published studies focusing on safety, feasibility, adherence to therapeutics, self-perception, and legal issues related to PrEP in people aged between 10 and 24 years. Keywords: HIV prevention, pre-exposure prophylaxis (PrEP), adolescents, youth

Introduction
Adolescents are a critical population that is disproportionately impacted by the HIV epidemic. AIDS is the leading cause of death among adolescents in sub-Saharan Africa and is the second cause of mortality among youth worldwide.1 Over 2 million adolescents between the age group of 10 and 19 years are living with HIV, and millions more are at risk of infection every year.2

Due to fundamental neurological, psychological, and behavioral changes that take place during adolescence,3 this group is considered highly vulnerable to contracting HIV.4,5 Infection risks are considerably higher among girls, especially in high-prevalence settings such as eastern and southern Africa. Due to preexisting, gender-based inequalities such as exclusion and sexual violence, this subgroup is disproportionately affected by HIV. In 2013, two-thirds of new HIV infections reported in people between the age group of 15 and 19 years were adolescent girls, particularly in sub-Saharan Africa.6

References

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Furthermore, besides young women, there are other youth populations that exhibit distinct sociobehavioral patterns that increase their risk to contracting the virus, such as intravenous drug users, gay and bisexual-identifying teenagers, adolescent sex workers, and those who belong to multiple risk categories. Their public health outcomes can be influenced by evidence-based primary preventive strategies, so the focus should be on them.

Goal #3 of the United Nations’ “Sustainable Development Goals” (SDG3) – to ensure healthy lives and promote well-being for all ages – requires as a target that by 2030 there should be an end to the epidemic of AIDS. To achieve this ambitious goal, each age group, population, and geographical location will require specific and flexible strategies.7

Pre-exposure prophylaxis (PrEP) as an HIV prevention strategy

PrEP is a new intervention for people at high risk to contracting HIV, with an estimated HIV incidence of >3%, based on the use of oral antiretroviral medications to help prevent the infection. A daily use of two combined antiretroviral drugs (tenofovir disoproxil fumarate [TDF]/emtricitabine [FTC]) was approved as PrEP in 2012, sold under the name Truvada.8

Recent data from clinical trials have demonstrated the efficacy of PrEP as a powerful HIV prevention tool in some high-risk populations including men who have sex with men,9 HIV-1-serodiscordant heterosexual couples,10 and intravenous drug users.11 PrEP effectively reduced HIV infection in four of six trials conducted with the adult population.9–11

The reported efficacy in those randomized trials using TDF was as a single agent or in combination with FTC to prevent HIV infection ranged from 44% to 75% and was heavily dependent on patient adherence (Table 1).

The Phase III randomized clinical trial PrEP initiative (iPrEx) highlighted the importance of adherence with overall protective efficacy of 44%, but as high as 92% in those with detectable drug levels.9 Two trials (the FEMPrEP12 and VOICE13) showed no significant protective effect in women, yet less than 30% adhered to the prescribed intervention, as observed through drug concentration analysis.

Despite the proven efficacy of PrEP in adult trials, there are concerns about real-life feasibility due to several barriers including stigma, cost, and limited experience in PrEP delivery by HIV providers. However, some of these concerns abated after demonstration projects showed high clinical effectiveness (>80%).14,15

Recent studies are attempting to expand the investigation into the efficacy of such HIV prophylaxis approaches in adolescent populations. However, there are still many knowledge gaps, and no country has yet approved it for use with adolescents. Guidelines have recommended the use of PrEP as part of prevention for persons at substantial risk for HIV infection without limiting the recommendations to specific populations.16

The aim of this review was to identify and summarize the evidence from studies on PrEP for adolescents. We mapped published studies focusing on the safety, feasibility, acceptability, adherence to therapeutics, self-perception, and legal issues related to PrEP in people aged between 10 and 24 years.

Criteria for considering studies for this review

Type of studies

We considered the following study designs for inclusion:

1. Primary studies, such as randomized, quasi-randomized, and nonrandomized clinical trials, longitudinal observational (historical cohorts, prospective cohorts, case–control, and before-and-after studies), analytic cross-sectional studies, and non-comparative studies (case and series reports or single experimental cohorts).

2. Secondary studies, including systematic reviews, guidelines, or economic evaluations. We did not consider narrative reviews, case reports, comments, or editorials.

<table>
<thead>
<tr>
<th>Table 1 Clinical trials of PrEP in the population at risk of HIV infection</th>
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<tbody>
<tr>
<td>Study</td>
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<tr>
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<tr>
<td>Partners PrEP (TDF + FTC or TDF)</td>
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<tr>
<td>Botswana TDF2 (TDF + FTC)</td>
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<td>Bangkok (TDF)</td>
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<td>iPrEx (TDF + FTC)</td>
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<td>FemPrEP (TDF + FTC)</td>
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<td>VOICE (TDF + FTC or TDF gel)</td>
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Abbreviations: FTC, emtricitabine; PrEP, pre-exposure prophylaxis; iPrEx, PrEP initiative; TDF, tenofovir disoproxil fumarate.
Type of participants
Although the focus was primarily on the second decade of life, in some cases data on youth or young people were included. This inclusion is generally due to data being aggregated in ways that did not distinguish adolescent years specifically.

Defining overlapping terms “adolescents,” “youth,” and “young people”
The World Health Organization (WHO) defines adolescent as those people between 10 and 19 years of age. The United Nations define youth as 15–24 years, and the term “young people” is used by WHO and others to combine adolescents and youth.17

Outcomes of interest
We considered all effectiveness and safety outcomes for therapeutic studies addressing PrEP use. For all studies, we collected data linking PrEP use to adherence, adolescent self-perception, satisfaction, health care provider attitudes related to youth, and legal issues.

Strategy for searching studies
We conducted a systematic search of studies on 23 and 24 January 2017 in the following databases: MEDLINE (via PubMed, from 1946), EMBASE (via Elsevier, from 1974), Cochrane Library (via Wiley, issue 1, 2017), Literatura Latino-Americana em Ciências da Saúde e do Coração (LILACS; via Biblioteca Virtual em Saúde [BVS], from 1966), and ADOLEC (via BVS). We also searched for ongoing or unpublished trials in the US National Institutes of Health Ongoing Trials Register (www.clinicaltrials.gov) and the WHO International Clinical Trials Registry Platform (ICTRP; http://apps.who.int/trialssearch/). The search strategies are available as Appendix 1.

Study selection
Two reviewers (DMM and RR) independently assessed titles and abstracts from all records retrieved by the literature search for eligibility according to the inclusion criteria. Studies selected at this first stage were then evaluated in its full text to confirm eligibility. Disagreements between reviewers were resolved using consensus-based models for decision making. The selection process was performed using the Rayyan platform.18

Data extraction and presentation of findings
The authors extracted the data on publication, methods, and results of included studies. The findings are summarized in Tables 1 and 2 using a narrative approach.

The search strategies retrieved 521 records. The PRISMA diagram is shown in Figure 1.

From the 44 studies initially retrieved by the search strategy, 31 were excluded, due to inclusion of participants older than 25 years, without depicting young people separately in the analysis. Of the 13 selected studies, there were six registered clinical trials, and two of which evaluated safety, feasibility, and the acceptability of daily oral TDF/FTC (Truvada®) specifically among adolescents.

The Adolescent Medicine Trials Network [ATN] protocol 113 (NCT01769456)19 recruited participants from six study sites in the USA, and the CHAMPS PillPlus (NCT02213328),20 in two peri-urban settings in South Africa. Other selected trials included participants up to 24 years of age combining behavioral HIV prevention interventions and oral PrEP. The trials and other studies in this review are summarized in Table 2. They also addressed several domains, including feasibility, medical adherence, the acceptability of oral daily TDF or TDF/FTC, patterns of PrEP use, patterns of sexual risk behavior, risk compensation, as well as cognitive and emotional processes in response to using PrEP. The ATN 082 study compared proven behavioral HIV prevention interventions (Many Men, Many Voices [3MV]) alone and in combination with PrEP or placebo.21

There are three selected trials that are still ongoing, with complete results unavailable as of yet. One of which is evaluating the differences in adherence to PrEP with or without the use of short message service (SMS) reminders (NCT02915367).22 The HPTN 082 trial is comparing standard adherence support versus enhanced counseling based on feedback from observed drug levels.23 Besides, qualitative research is availing the factors influencing PrEP initiation and adherence.

The NCT0281024924 has a qualitative design aims to identify and understand African American young men who have sex with men (YMSM)’s cognitive and emotional processes in response to using PrEP to reduce their risk for HIV, as well as to identify what factors influence African American YMSM’s likely use of PrEP.

The systematic review retrieved studies’ referencing values and preferences among populations that might benefit from PrEP, as well as health care providers who may prescribe PrEP. Of the 76 included studies, 71 addressed adult populations. Just five studies focused on adolescent girls and young women.25 The substudy of Crew 450, an ongoing longitudinal cohort study, describes PrEP interests among racially and ethnically diverse YMSM aged 16–20 years, utilizing a PrEP interest scale with a score of the likelihood of using PrEP.26

Fisher et al27 provided empirical data on sexual and gender minority youth (SGMY) self-consent that can assist
### Table 2 Summary of the included studies

<table>
<thead>
<tr>
<th>Study/country</th>
<th>Type of study</th>
<th>Population/age</th>
<th>Aims/outcomes</th>
<th>Description</th>
<th>Results</th>
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<tbody>
<tr>
<td>Project PrEPare</td>
<td>Open-label demonstration project and Phase II safety study</td>
<td>Completed: 78 young HIV-negative MSM 35% Black/ African American 45% Latino/ Hispanic, 15–17 years (ATN 113); 18–22 years (ATN 110)</td>
<td>To evaluate: Safety of daily oral TDF/FTC (Truvada®) Patterns of use Rates of adherence Patterns of sexual risk behavior</td>
<td>Study subjects engaged in a behavioral risk-reduction intervention before PrEP. Monthly visits for the first 12 weeks of study. Quarterly visits thereafter until 48 weeks of study.</td>
<td>Acceptability of PrEP was high: -56% with TDF diphosphate drug levels consistent with ≥4 pills per week (at week 24) and 34% at week 48. Noticeable drop-off occurring at week 24</td>
</tr>
<tr>
<td>ATN 113 (NCT01769456) and ATN 110 (NCT01772823)</td>
<td>Demonstration project and Phase II safety study</td>
<td>Completed: 68 youth (mean age = 19.97 years; 53% African American, 40% Latino)</td>
<td>To compare an efficacious behavioral HIV prevention intervention (3MV) alone, 3MV combined with PrEP (TDF/FTC), and 3MV combined with placebo.</td>
<td>3MV intervention: conducted as a 2-day seminar throughout centrally located communities in Chicago with approximately eight participants per session. After 3MV, study participants were randomized in blocks of six within each site to one of three study arms: 1) daily combo FTC and TDF as PrEP, 2) placebo pill control (with HIV behavioral intervention), or 3) “no pill” control (subjects receive HIV behavioral intervention but no pills)</td>
<td>The feasibility of enrolling at-risk youth, particularly YMSM of color has been demonstrated. The acceptability of the group intervention along with counseling and testing was high. Self-reported medication adherence and corresponding plasma drug concentrations were low. Behavioral disinhibition was not seen. Decrease in sexual risk behavior was seen over time</td>
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<tr>
<td>Project PrEPare</td>
<td>Prospective, open-label study</td>
<td>Completed: HIV-negative adolescents 98 female 50 male 15–19 years Median age of 18 years (17–19 years)</td>
<td>To evaluate: Safety, feasibility, acceptability of daily oral TDF/FTC (Truvada®)</td>
<td>Daily oral PrEP as part of an HIV prevention package that included condoms and STI screening and treatment. Adherence support using SMS, adherence clubs, real-time feedback on drug levels</td>
<td>Estimated date for final data collection date for primary outcome measure: August 2017</td>
</tr>
<tr>
<td>ATN 082 (NCT01033942)</td>
<td>Randomized multisite prospective study</td>
<td>Enrolled: adolescent and young adult women (16–25 years)</td>
<td>To assess the acceptance rate, adherence, acceptability, and continuation of oral PrEP among young Southern African women</td>
<td>Standard adherence support versus enhanced counseling based on feedback from observed drug levels in the first 2 months after PrEP initiation. Qualitative research about factors influencing PrEP initiation and adherence (for those who initially declined PrEP)</td>
<td>Ongoing</td>
</tr>
<tr>
<td>CHAMPS PillsPlus</td>
<td>Prospective, open-label study</td>
<td>Enrolled: 78 young HIV-negative MSM 35% Black/ African American 45% Latino/ Hispanic, 15–17 years (ATN 113); 18–22 years (ATN 110)</td>
<td>To evaluate: Safety of daily oral TDF/FTC (Truvada®) Patterns of use Rates of adherence Patterns of sexual risk behavior</td>
<td>Study subjects engaged in a behavioral risk-reduction intervention before PrEP. Monthly visits for the first 12 weeks of study. Quarterly visits thereafter until 48 weeks of study.</td>
<td>Acceptability of PrEP was high: -56% with TDF diphosphate drug levels consistent with ≥4 pills per week (at week 24) and 34% at week 48. Noticeable drop-off occurring at week 24</td>
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<tr>
<td>(clinical trial: NCT02123128)</td>
<td>Situation: active, not recruiting</td>
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<tr>
<td>Country: South Africa (Johannesburg and Cape Town)</td>
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<tr>
<td>HPTN 082 (clinical trial: NCT01033942)</td>
<td>Randomized multisite prospective study</td>
<td>Situatin: active, not recruiting</td>
<td>To assess the acceptance rate, adherence, acceptability, and continuation of oral PrEP among young Southern African women</td>
<td>Standard adherence support versus enhanced counseling based on feedback from observed drug levels in the first 2 months after PrEP initiation. Qualitative research about factors influencing PrEP initiation and adherence (for those who initially declined PrEP)</td>
<td>Ongoing</td>
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<tr>
<td>Country: South Africa and Zimbabwe</td>
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Table 2 (Continued)

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<tr>
<th>Study/country</th>
<th>Type of study</th>
<th>Population/age</th>
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<th>Results</th>
</tr>
</thead>
<tbody>
<tr>
<td>Clinical trial</td>
<td>Interventional, randomized,</td>
<td>Young women (18–24 years)</td>
<td>1) To evaluate the difference in adherence to PrEP by the study arm: SMS reminders versus no reminders 2) Acceptability 3) Risk perception</td>
<td>Intervention: behavioral. Adherence Wisepill data and TDF concentration on dried blood spots</td>
<td>Ongoing</td>
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<tr>
<td>(NCT02915367) Country: Kenya</td>
<td>open-label</td>
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<tr>
<td>Clinical trial</td>
<td>Observational, with a</td>
<td>20–30 African American YMSM aged 16–24 years</td>
<td>1) To identify and understand African American YMSM’s cognitive and emotional processes in response to using PrEP to reduce their risk for HIV 2) To identify what factors (sociocultural, individual, experiences in health care, socioeconomic) influence African American YMSM’s likely use of PrEP</td>
<td>Individual interviews</td>
<td>Ongoing</td>
</tr>
<tr>
<td>(NCT02810249) Country: USA</td>
<td>qualitative design</td>
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<tr>
<td>Koechlin et al25</td>
<td>Systematic review</td>
<td>78 studies evaluated hypothetical use of PrEP (five studies were conducted among “adolescent girls and young women,” 26 studies included individuals who actually took PrEP or placebo)</td>
<td>To describe the evidence about values and preferences among populations that might benefit from PrEP and among health care providers who may prescribe PrEP</td>
<td>Three electronic databases of articles and HIV-related conference abstracts (January 1990–April 2015) Type of studies: 68 quantitative 24 qualitative 12 mixed methods</td>
<td>Participants showed strong interest in PrEP (aged 14–24 years). They appreciated the “privacy” of a pill; some girls expressed hesitation with regard to PrEP. Participants would be willing to take PrEP if it was free. One study showed 20% of young women expecting to use condoms less frequently if they took PrEP.</td>
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<tr>
<td>Substudy of Crew 450 (an ongoing longitudinal cohort study, N = 451 YMSM); Chicago, USA</td>
<td>Observational, cross-sectional</td>
<td>YMSM aged 16–20 years N = 184</td>
<td>To describe PrEP interests among racially and ethnically diverse YMSM aged 16–20 years using a new measure that may have utility in the future studies</td>
<td>Computer-assisted self-interview with audio instructions. PrEP interest scale with a score of the likelihood of using PrEP</td>
<td>Participants with better HIV knowledge had higher interests in PrEP. Higher level of education was related to an increased likelihood of taking PrEP. No relationship between most of the reported risk behaviors and mean PrEP interest scores. Indirectly proportional relationship between unprotected anal intercourse with PrEP interest</td>
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(Continued)
Despite the encouraging results of PrEP trials in adults, adolescents have the ability to self-consent as compared to age- and population-appropriate procedures. Guardian permission as a significant barrier to research participation.

Finally, the study by Mack et al describes clinician attitudes toward, and practices around, PrEP use in youth.

Despite the encouraging results of PrEP trials in adults, their relevance to the realities of adolescents and young adults

### Table 2 (Continued)

<table>
<thead>
<tr>
<th>Study/country</th>
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<th>Population/age</th>
<th>Aims/outcomes</th>
<th>Description</th>
<th>Results</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fisher et al&lt;sup&gt;37&lt;/sup&gt;</td>
<td>Qualitative study, using</td>
<td>Enrolled: 60</td>
<td>To provide empirical data on SGMY self-consent that can assist IRBs on strategies to increase their research participation</td>
<td>Online survey and asynchronous focus group questions after watching a video about PrEP. Discussion addressed: guardian permission, random assignment, privacy concerns, and PrEP medication adherence</td>
<td>Adolescents have the ability to self-consent as compared to age- and population-appropriate procedures. Guardian permission as a significant barrier to research participation</td>
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<tr>
<td>NY, USA</td>
<td>a Web-based qualitative/</td>
<td>sexually active SGMY 14–17 years</td>
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<td>mixed methods analysis program</td>
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<tr>
<td>Mack et al&lt;sup&gt;30&lt;/sup&gt;</td>
<td>Qualitative study</td>
<td>South Africa:</td>
<td>To explore the potential role of choice in women’s use of HIV prevention methods. To present the analysis of groups’ attitudes toward ARV-based HIV prevention</td>
<td>All focus groups were audio-recorded, transcribed, and translated into English. Transcripts were coded using a codebook and QSR NVivo 9.0 (QSR International, Melbourne, Australia); generated code reports; and conducted inductive thematic analysis to identify major trends and themes</td>
<td>All groups expressed strong interest in PrEP products. Adolescent girls believed that it would be possible to obtain the products more privately than condoms. Some girls stated that they would be interested in using PrEP only after seeing other girls use it. Young women showed concern that it would be challenging to negotiate PrEP use with a partner. Minors’ access to PrEP without parental consent is unclear. No state expressly prohibits minors’ access to PrEP. All jurisdictions allow adolescents to consent to diagnosis and treatment of STIs; only eight jurisdictions allow consent to prophylactic services.</td>
</tr>
<tr>
<td>Country: Kenya and South</td>
<td>Kenya (adults) and South Africa (adolescents and young women)</td>
<td>(N = 36 participants). Two FGs with adolescent aged 14–17 years and two with young women aged 18–24 years</td>
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<td>Africa</td>
<td>Qualitative study</td>
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<td>Culp and Caucci&lt;sup&gt;28&lt;/sup&gt;</td>
<td>Analytic, WestlawNext, a subscription-only online legal research service</td>
<td>A 50-state analysis of minor consent laws and its implications for PrEP</td>
<td>To analyze laws related to a minor’s ability to consent to medical care, including HIV diagnostic testing and treatment, and its implications for PrEP</td>
<td>Staff collected all statutes and regulations about an adolescent’s ability to consent to HIV diagnosis, treatment, and prevention. On laws current as of December 31, 2011</td>
<td>No state expressly prohibits minors’ access to PrEP. All jurisdictions allow adolescents to consent to diagnosis and treatment of STIs; only eight jurisdictions allow consent to prophylactic services.</td>
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<tr>
<td>Country: USA</td>
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<tr>
<td>Clinical trial (ATN 113</td>
<td>Qualitative study</td>
<td>17 research personnel at 13 ATN sites considering “implementation” of ATN 113</td>
<td>To examine factors related to the process by which IRBs and research personnel made decisions regarding whether to approve and implement the protocol. Emphasis on the issue of adolescent consent with high-risk minor participants</td>
<td>Semi-structured interviews and the receipt of IRB-related correspondence and documents generated during the review process</td>
<td>Of 13 ATN sites in 12 states: seven received IRB approval for ATN 113, three were denied approval, and three received a formal disposition.</td>
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<td>113 substudy)</td>
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**Abbreviations:** ATN, Adolescent Medicine Trials Network; ARV, antiretroviral; FGs, focal groups; FTC, emtricitabine; IRBs, institutional review boards; PrEP, pre-exposure prophylaxis; SGMY, sexual and gender minority youth; SNS, short message service; STI, sexually transmitted infection; TDF, tenofovir disoproxil fumarate; 3MV, Many Men, Many Voices; YMSM, young men who have sex with men.
is not directly transferable. Thus, data from studies including youth and adolescents are critical for implementation of PrEP in such a highly vulnerable population.

At the time of this review, there were few PrEP studies among adolescents with available results. One such study, Project PrEPare (ATN protocol 113), was the first adolescent PrEP study among YMSM aged 15–17 years. Some findings of ATN 113 were initially presented at the Conference on Retroviruses and Opportunistic Infections (CROI).31

Two interconnected issues are important to highlight in this trial. The first is related to adherence: long-term, high adherence has proved to be essential for PrEP to be successful. Most of the adolescents participating in project PrEPare achieved protective drug levels during monthly visits, but adherence decreased when visits became quarterly. Among MSM aged 18–22 years, TDF in the plasma was detectable in only 20% of participants at week 24.21 In the follow-up study, <34% had drug levels consistent with 4+ pills/week at week 48.31 Such a finding is in contrast to adult trends seen in recent trials where adherence by drug detection was quantified at 51–100%.32

Other important findings relate to stigma as a barrier to effective health interventions. Adherent participants were significantly less likely than non-adherents to report concerns about what people would think when finding they were taking PrEP pills. They anticipated the perception of stigma, as living with HIV meant living with the fear and effects of stigmatization, with consequent social rejection, discrimination, or violence.33 Likewise, previous studies have shown that associating PrEP with sexual risk taking may reduce the motivation of young people to seek PrEP or to maintain PrEP use for fear of stigmatization.34,35 The process of enrolling adolescents in this trial was more time-consuming than in the parallel study group of 18–22 years (ATN protocol 110).36 The adolescents were allowed to provide self-consent to participate. This approach reduced by 50% the number of eligible sites cleared by their IRBs to be part of the study, as related in the qualitative substudy of ATN 113.
The complexity of the legal, ethical, and practical barriers to conducting PrEP studies with adolescents are evident, as is the need for coherent and comprehensive standards for participation.28,29

The second trial, CHAMPS PillPlus (NCT02213328), included both male and female HIV-negative adolescents aged 16–19 years.30 Those under the age of 18 years had to obtain parental/guardian consent to participate in the study, but this requirement did not disturb the enrollment. Considering that adherence to daily medication regimens poses a significant challenge for adolescents, this study provided support using SMSs, adherence clubs, trained youth-friendly counselors, and real-time feedback on drug levels. A baseline analysis of data from the study indicates a reasonable uptake and use of PrEP, based on initial plasma TDF levels. Those findings are encouraging, considering results from iPrEx indicating that PrEP reduced the risk of HIV infection by 92% in participants with detectable drug levels in plasma.9

Both ATN 11319 and CHAMPS20 trials showed a high level of acceptability to PrEP among YMSM, girls, and young women. Although some girls expressed hesitation with regard to PrEP, they appreciated the privacy of a pill and would be willing to take PrEP if it was provided for free in the USA. YMSM with better HIV knowledge and a higher level of education showed more interest and were more likely to take PrEP. Irrespective of prior knowledge, they expressed a desire for more information about this new prophylactic strategy. Prudent information for young PrEP users would include minimum necessary frequency of use and if gender would influence the overall dose to achieve similar levels of protection.38,39 Available data suggest that more consistent dosing is required to achieve sufficient levels of TDF in vaginal tissue when compared to rectal tissues.40 However, as demonstrated in the efficacy clinical trials of PrEP, women who were adherent to a daily PrEP regimen were strongly protected against HIV.10,11,41

Project PrEPare also demonstrated the feasibility of enrolling and retaining a young cohort of YMSM into a PrEP trial despite an intensive visit schedule.21 The data focus on YMSM and the general group of adolescents at risk for HIV infection. Nonetheless, the scientific health literature shows a lack of qualitative information on the attitudes and perceptions toward PrEP acceptability in other subgroups of adolescents such as intravenous drug users, transgender youth, adolescent sex workers, or teenagers in other cultural contexts.

Other ongoing, adolescent-inclusive trials hopefully will deliver more data to the discussion. One such study, HPTN082,23 is focused on acceptance and adherence among adolescents and young women aged 16–25 years.

Haberer et al42 showed that scheduled SMS reminders improved highly active antiretroviral therapy (HAART) in the context of real-time monitoring in an adult population. The trial NCT02915367 is assessing the use of SMS reminders (a behavioral intervention) to improve medication compliance in young people.22 A better comprehension of useful indicators of adherence among adolescents at high risk for HIV infection is needed to ensure the effectiveness of PrEP programs.

Young people are not at uniform risk throughout adolescence, and PrEP does not need to be lifelong. As pointed by Hosek et al,21 “PrEP for young people may be best viewed as a time-limited strategy that can bridge the developmental period between sexual debut and adulthood.” This idea may have a strong impact on acceptability and points out the difference between the use of PrEP and HAART for treatment.

Combining behavioral and biomedical approaches may be critical to successful implementation of PrEP programs among youth populations. One such trial, ATN 082, shows the feasibility of enrolling at-risk youth with behavioral intervention plus biomedical prevention (PrEP) with high acceptability; however, the self-reported pill adherence and corresponding plasma drug concentrations from participants were low. More optimistic findings, however, were the reduction in behavioral disinhibition with decrease in risky sexual behavior over time.21

One of the five studies included in the systematic review by Koechlin et al25 was not retrieved by our search strategy, so it does not appear individualized in Table 2. The adolescent girls showed interest in PrEP as protection from HIV, although a few participants appeared confused about if PrEP would protect them from pregnancy as well.30

One of the major concerns that could undermine PrEP benefits regarding risky sexual behavior is risk compensation, that is, PrEP could reduce condom use and increase the risk for other sexually transmitted infections (STIs).43–46 Rubtsova et al47 found that 20% of young women expected to use condoms less frequently if receiving PrEP. The data of adolescent girls from South Africa corroborate in part with this concern. They found PrEP particularly appealing because it would eliminate concerns about being seen while obtaining condoms from clinics.30 The use of condoms is closely related to exposure to partner violence. Even the fear of violence is linked to women’s reluctance or inability to negotiate condoms or to use contraceptives.8 Indeed, if people were using condoms consistently, the rates of new HIV infections would not remain at 2 million new infections.
Adherence | PrEP for youth
---|---
Risk compensation | Reduce risk of HIV infection
Stigma | Provide autonomy
Ethical and legal issues | Can be used in a period of risk

Figure 2 Factors that influence acceptability and implementation of PrEP in adolescents.

Note: Arguments in favor and the issues that should be better addressed.

Abbreviation: PrEP, pre-exposure prophylaxis.
support sound decision making by health care professionals, aiming to not only protect individuals but also contribute to the control of the HIV pandemic.

Some argue that due to cost and adherence barriers, PrEP will not benefit those most in need and may actually enhance existing HIV-associated disparities.62

The inclusion of PrEP in public programs will require guidance to maximize impact and effectiveness based on region, population group, and individual risk.63 As Aggleton64 has remarked, “PrEP is an HIV prevention strategy that may be useful for some people in some contexts some of the time.” It is clear that the benefits of PrEP will vary across different programs and global regions.

Conclusion

Oral PrEP offers an exciting opportunity for adolescents and young adults to access a new prevention option, respecting their particularities and vulnerabilities. Detailed information about the results of available studies should be adequately disseminated to both the youth population and their health care providers to facilitate effective and safe delivery of PrEP. Adherence to PrEP might be facilitated by technology, including SMS and wireless electronic monitoring, as well as counseling and adherence feedback (by measuring drug concentration when feasible). The focus should not be to use PrEP indefinitely, but rather to align PrEP use with periods of high risk for HIV infection. It is essential to reject stereotypes and sex-negative messaging in guiding decisions of having PrEP access and uptake, since they may contribute to enhance stigma. Overall, it is evident that substantial more research is needed focusing on adolescents and PrEP.

Each country and program should research and understand the peculiar characteristics of their adolescent target populations—what they think, how they feel, and their perspectives surrounding PrEP and risky sexual behavior. With this information, policy makers and clinicians will be better equipped to serve and implement specific strategies that respond adequately to their health care needs.

Disclosure

The authors report no conflicts of interest in this work.

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