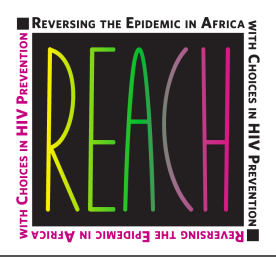
# Things to Do Things to Consider

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Meeting the HIV Prevention Needs of Adolescent Girls and Young Women South Africa Stakeholders Meeting on REACH 28 February 2017, Johannesburg



### Getting Ready for REACH

### **Overall Study Timeline**

February 2017	<ul> <li>Protocol Version 1.0 approved by NIAID/NIH and sent to sites</li> </ul>
March – August 2017	<ul> <li>Ethics and MCC submissions and approvals</li> </ul>
September – December 2017	<ul> <li>Final preparations at sites</li> </ul>
Before end of 2017	<ul> <li>Sites expected to begin enrolling participants</li> </ul>
Mid-2020	<ul> <li>Study follow-up completed: 18 months per participant</li> </ul>

### Ethics and in-country approvals

- In South Africa, DTHF and Wits RHI will each submit the protocol and consent materials (informed assent, informed consent and parental/guardian permission form) to Ethics Committee
  - Wits Human Research Ethics Committee
  - University of Cape Town Human Research Ethics Committee
  - We aim for submissions March/April
- The IRB/EC may have questions, request additional information, or request revisions
  - Process can take about 3 months
  - Because the study involves minors, review may take longer
- Concurrently, we will seek approval from the Medicines Control Council of South Africa, on behalf of both DTHF and Wits RHI

#### The process is important because...

The protocol review process is vital to ensuring that a participant's safety, rights and welfare are protected, and that the research being conducted has scientific merit





#### We don't wait – the work has already begun!

- Each site must prepare several documents, procedures and forms
- The clinic, pharmacy, laboratory must meet specific criteria and be approved for the study
- For REACH, we will be working to ensure youth-friendly clinic space
- Additional staff may be hired to ensure the team has people with the right skills, experience and personal qualities for the study
- Site staff will undergo days of training, conduct mock participant visits and counseling sessions

#### Community Engagement Plans

- Outreach activities with CAB and community stakeholders ramp up
- Creation and EC approvals of community and participant education materials
- Planning for participant and community engagement
  - To support recruitment, retention, adherence
  - To ensure community support
  - To address concerns and rumors up front

### There is a lot to do!

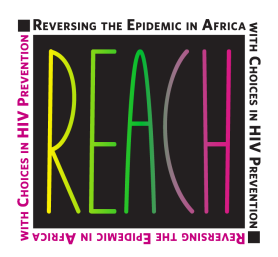
It may take up to 6 months from the time of submission before the site is ready to begin enrolling participants





Doctors, nurses, counselors, pharmacy staff, lab technicians, clerks, drivers, and custodians -everyone must be ready before the study can begin to ensure successful implementation and conduct of the study.

## Questions?



#### The Realities and Requirements for Participating in REACH

### Who may take part in REACH?

- For a young woman to join the study, she must meet the following requirements:
  - Be between 16-21 years old
  - Be able to comply with study requirements
  - Not be HIV-infected
  - Have had sex at least once
  - Not be pregnant
  - Been using an effective contraception for at least 2 months before enrollment
  - Not have any active STIs or other infections
  - Be generally healthy, as per physical and pelvic exams, as well as laboratory tests

#### A research study isn't for everyone

- For REACH, girls must be willing to come to the study clinic every month, for 1 ½ years
  - There are many medical tests and exams some study visits may take several hours
- They must have experienced sex and for those under age 18, must have a parent's permission
  - This means girls will need to admit they are having sex
- Participants should be willing to use the study products as directed
  - And if unable to use the products, to be open and honest about her experiences
  - Researchers need to understand these challenges so they can try to address them

#### Services provided as part of the study

- Monthly visits will always include:
  - HIV and pregnancy testing
  - HIV/STI risk reduction, contraceptive counseling and provision of condoms
  - Individualized product adherence counseling
- Contraception counseling and services
- Social harms assessments, physical exams and pelvic exams will also be conducted at some visits
- Referrals will be provided for any issue the site cannot manage or cannot include as part of the study, such as HIV counseling and treatment, pregnancy and antenatal care and/or specialized counseling or other services.

### Deciding to join the study

- In-depth informed assent and consent discussions with potential participants and their parents will be critical to ensuring that participants clearly understand the study procedures, time commitment, and importance of a successful study.
- Participation is voluntary. Likewise, participants may choose to leave the study at any time



- □ Enrolling in a research study is a commitment
- Enrolling in REACH requires commitment, but also a sense of purpose – to wanting to make a difference in the lives of adolescent girls and young women in their communities and around the world





## Questions?