PTID:

Instructions: Use the table below to document a woman participant's eligibility status for MTN-011 study participation. Initial and date below each set of "yes/no" checkboxes upon assessment of each eligibility criterion. Once ineligibility status is determined, the form may be stopped and the remaining questions may be left blank.

Inclusion Criteria	Screening Visit	Enrollment Visit
1. <i>a.</i> Able and willing to provide written informed consent to be screened for and take part in the study	Yes No	Yes No
<i>b.</i> Able and willing to provide adequate locator information, as defined by the site SOPs.	review and proceed accordingly	
2. a. Per participant report, no STIs in the 6 months prior to Screening		not required
b. Per participant report, no non-therapeutic intravenous drug use in the 18 months prior to Screening		not required
<i>c.</i> Per participant report, in a mutually monogamous relationship with a partner of the opposite sex for 6 months prior to Screening and the intent to stay in this relationship for the next 4 months.		not required
3. At Screening and Enrollment, both partners independently report not using barrier contraception and/or barrier protection as part of the normal coitus routine and report the intent to continue said sexual practice for the duration of study participation.		
4. HIV-uninfected, based on testing performed by study staff at Screening (per algorithm in protocol appendix III)		not required
 Agrees not to participate in other research studies involving drugs, medical devices, genital or rectal products, or large blood draw studies during study participation. 		

Note: In order for the participant to be <u>eligible</u>, all of the responses to items 1-5 above <u>must be "yes"</u>.

Inclusion Criteria – cont'd	Screening Visit	Enrollment Visit
	Yes No	Yes No
6. a. Age 21 through 46 years (inclusive) at Screening, verified per site SOPs.		not required
b. Pap results in the 12 calendar months prior to Screening consistent with Grade 0 according to the FGGT* or satisfactory evaluation with no treatment required of non-Grade 0 Pap result per American Society for Colposcopy and Cervical Pathology (ASCCP) guidelines in the 12 calendar months prior to the Screening Visit		not required
Note : Women with a documented normal results with the 12 months prior to screening need not have a Pap smear during the S abnormal Pap smears can be enrolled upon completion of the initial phase of evaluation if no current treatment is indicated (bae management of abnormal cervical cytology). Need for a repeat Pap within 6 months does not preclude enrollment prior to that results and the second s	ed on local standard	ofcare for
c. Must be currently using effective non-barrier contraception, other than a contraceptive vaginal ring, for at least three months prior to Screening (i.e. oral contraceptive, patch, injectable hormones, subdermal implants, intrauterine device, female or male sterilization) and intending to use this method for the course of the study.		
<i>d.</i> Per participant report, regular menstrual cycles with at least 21 days between menses (does not apply to participants who report using a progestin-only method of contraception at screening, e.g., Depo-Provera)		not required
Note: This criterion is not applicable to participants using continuous combination oral contraceptive pills, as the absence of regular menstrual cycles is an expected, normal consequence in the context		
e. Anatomy sufficient for performing pelvic examinations and for collection vaginal and cervical specimens.		not required
f. Must also agree to abstain from intercourse (oral, anal, or penile-vaginal) and other vaginal practices (e.g. masturbation, douching, tampon use, application of lubricants/spermicides or other related practices) 72 hours prior to each follow-up visit. Group 2 participants must agree to also abstain from the aforementioned practices throughout the at-home gel use period.		

Note: In order for the participant to be eligible, all of the responses to items 6 above must be "yes".

*Female Genital Grading Table for Use in Microbicide Studies Addendum 1 to the DAIDS Table for Grading Adult and Pediatric Adverse Events, Version 1.0, December 2004 (Clarification dated August 2009).

Exclusion Criteria	Screening Visit	Enrollment Visit
	Yes No	Yes No
Participant report of any of the following: 1.		not required
a. Known allergy to study product (ever)		
b. Post-exposure prophylaxis (PEP) for HIV exposure within 6 months prior to Screening		not required
 Participation in any other research study involving drugs, medical devices, or genital products 30 days or less prior to Enrollment 	review and proceed accordingly	
<i>d.</i> Plans to relocate away from the study site in the next 4 months.		not required
e. History of domestic violence with current partner (ever)		not required
f. Systemic or topical antimicrobials within the last 7 days prior to Enrollment	review and proceed accordingly	
g. Currently using or planning to use pharmacologic immune modulator(s)		
2. At Screening or Enrollment, symptomatic urinary tract infection (UTI)		
Note : Otherwise eligible participants diagnosed with UTI during screening are offered treatment and may be enrolled after complete resolved as long as treatment is completed and all symptoms have resolved with 30 days of obtaining informed consent for Screen		symptoms have

Note: In order for the participant to be <u>eligible</u>, all of the responses to items 1-2 above <u>must be "no"</u>.

Exclusion Criteria cont'd	Screening Visit	Enrollment Visit
	Yes No	Yes No
3. At Screening, has a positive hepatitis B surface antigen (HbsAg) test result		not required
 At Screening and Enrollment, has an STI or reproductive tract infection (RTI) requiring treatment per current CDC guidelines 		
5. Genital signs and/or symptoms of Grade 2 or higher		
Note: For female participants, cervical bleeding associated with speculum insertion and/or specimen collected judged to be with clinical judgment of the IoR/designee is considered expected non-menstrual bleeding and is not exclusionary. Note: Otherwise eligible participants with exclusionary genital findings may be enrolled after the findings have improved to a non-exast long as treatment is completed and all symptoms have resolved within 30 days of obtaining informed consent for Screening/Enrol	xclusionary severity	Ū
6. Has any other condition that, in the opinion of the IoR/designee, would preclude informed consent, make study participation unsafe, complicate interpretation of study outcome data, or otherwise interfere with achieving study objectives		
 7. a. Participant report (or clinical finding) of the following: <i>i.</i> Last pregnancy outcome 90 days or less prior to Enrollment 	review and proceed accordingly	
ii. Currently pregnant		
Note: Self-reported pregnancy is adequate for exclusion from the study. A documented negative pregnancy test performed by stud	ly staff is required fo	or inclusion.
iii. Currently breastfeeding		
<i>iv.</i> Intends to become pregnant in the next 4 months	review and proceed accordingly	

Note: In order for the participant to be eligible, all of the responses to items 3-7a(iv) above must be "no".

Exclusion Criteria cont'd	Screening Visit	Enrollment Visit
	Yes No	Yes No
 V. Gynecologic or genital procedure (e.g. tubal ligation, dilation and curettage) within the prior 30 days to Enrollment 	review and proceed accordingly	
Note: This does not include biopsy for the evaluation of an abnormal pap results or endometrial biopsy that occurred more than 7 days prior to Enrollment.		
 7. b. Any of the following laboratory abnormalities at Screening: <i>i.</i> Hemoglobin less than 10.0 g/dl 		review and proceed accordingly
<i>ii.</i> Platelet count less than 100,000/mm ³		review and proceed accordingly
Note: Otherwise eligible participants with an exclusionary test may be re-tested during the screening process.		
7. c. Use of a vaginal douche or other Intravaginal products (excluding tampon use) in the 30 days prior to Enrollment	review and proceed accordingly	
7. d. Currently menopausal or perimenopausal		

Note: In order for the participant to be <u>eligible</u>, all of the responses to items 7a(v)-7d above <u>must be "no"</u>.

At enrollment visit, participant is found to meet all eligibility criteria:

Signature of staff member

Date

Signature of Investigator of Record (or designee)

Date