PTID:	Visit #·

Adverse Events Log [Ongoing Logs]

0000	
	DATE SENSIT

DATE SENSITIVE DATA ENTRY. Please enter the data from the paper CRF into REDCap ASAP.

01	REDCap entry Date:	$oldsymbol{0}$ You do not need to enter a date on this paper
		CRF, but you will be prompted to click the "Today"
		button when entering this AE in REDCap
02	Date site was informed of AE:	
		/ / (dd/mm/yyyy)
03	Adverse Event Description:	
04	AE onset date:	
		/(dd/mm/yyyy)
05	Body system:	☐ Constitutional
		☐ Cardiovascular
		□ Digestive
		☐ Endocrine
		☐ Hemic/Lymphatic ☐ Metabolic/Nutritional
		☐ Musculoskeletal
		□ Nervous
		☐ Respiratory
		☐ Skin/Appendages
		☐ Special Senses (5 senses + equilibrium)
		☐ Urogenital
		□ Infection
		□ HEENT
		☐ Other (answer 05a)
00	95a. Complete only if other body system:	
		•
	Other body system:	
•		
06	Severity:	☐ Grade 1 - Mild
		☐ Grade 2 - Moderate
		☐ Grade 3 - Severe
		☐ Grade 4 - Life threatening
		☐ Grade 5 - Death
07	Was this AE a worsening of a pre-existing condition?	☐ Yes
		□ No
80	Study Product Administration:	☐ No change
		☐ Held (2nd ring not administered)
		☐ Permanently discontinued
		☐ Not applicable
09	Status:	☐ Continuing
		☐ Continuing at end of study participation
		☐ Death (answer 9a)
		☐ Severity/frequency increased
		☐ Resolved/Stabilized (answer 09a)

MAT	RIX-003 Adverse Events Log	PTID:	Visit #:
Adve	erse Events Log (continued)		
00	Pa. Complete only if status marked "Death" or "Resolved/Stabilized":		
	Status/Outcome Date://(dd/mm/yyyy)		
10	Treatment: Mark 'none' or all that apply:	 □ None □ Ring removed □ Medications (answer 10a) □ New/Prolonged hospitalization □ Procedure/Surgery (answer 10c) □ Other (answer 10d) 	(answer 10b)
	10a. Complete only if treatment marked "medications" at question 10: Medications, specify:		
	10b. Complete only if treatment marked "new/prolonged hospitalization" at q New/Prolonged hospitalization, specify: Brief details.		
	10c. Complete only if treatment marked "procedure/surgery" at question 10: Procedure or surgery, specify:		
	10d. Complete only if treatment marked "other" at question 10: Other treatment, specify:		
11	This AE was first reported at:	Usit 2 (enrollment) Usit 3 (phone call) Usit 4 (clinic visit) Usit 5 (clinic visit) Usit 6 (clinic visit) Usit 7 (phone call) Usit 8 (clinic visit) Usit 9 (clinic visit/IDI) Interim Visit Unscheduled phone contact Other (answer 11a)	

Is this AE serious according to ICH guidelines?

 $\mathbb{O}_{ extstyle{1}}$ 11a. Complete only if AE first reported at was marked "other":

Other, specify: _____

☐ Yes (answer 12a)

□ No

MATRIX-003 | Adverse Events Log

Adverse Events Log (continued)

☐ Important Medical Event	e risk of death) on of existing hospitalization <i>(answer 12b)</i> capacity or substantial disruption of the ability to conduct normal life functions
Hospitalization admission date: Hospitalization discharge date:	marked "hospitalization/prolongation of existing hospitalization": // (dd/mm/yyyy) // (dd/mm/yyyy)
	marked "other":
12a. Continued. Complete only if this AE is serious. Has the participant had any diagnostic test related to this SAE?	according to ICH guidelines: ting or labs done
Indicate diagnostic tests and labs done, inc Update as needed. Additional space for notes if need	cluding results if known:

Visit #: ____

MATE	RIX-003 Adverse Events Log		PTID:	Visit #:
Adve	rse Events Log (continued)			
13	Date participant had first intravaginal ring study product inserted?	//	(dd/mm/yyyy)	
14	Has participant had 2nd intravaginal ring study product inserted?	☐ Yes (answer 14a) ☐ No		
①	14a. Complete only if participant had 2 nd intravaginal ring study product in: Date participant had second intravaginal ring study product inserted?		(dd/mm/yyyy)	
() F	telatedness			
15	Relatedness to Study Product/Procedure: • Relatedness to be determined by a study clinician.		□ Not related □ Related	
16	Justification of relatedness (for both related and not related):			

17

Name of Clinician determining relatedness:

Adverse Events Log (continued)

Please include a narrative documenting any additional treatment, hospitalization, or outcomes for this AE/SAE. Add additional notes as needed. End each note with your name or initials and the date.

18	Comment (1):	
		\
		1

19	Comment (2):

MATRIX-003	Adverse	Events Log
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PTID: _____ Visit #: ____

Adverse Events Log (continued)

20	Comment (3):



CRF Completed By: _____ (initials)

CRF Completion Date: __ _ / __ _ _ / __ _ _ (dd/mm/yyyy)