Annotated Study Book - AT3SC002

DS--Disposition

'DSTERM = INFORMED CONSENT OBTAINED

AT:	3SC002: Demographic Data and Informed Consent Recordin	g (DM) [frmDM] INFORMED CONSENT OBTAINED
Den	nographics [sctDM]	
1.*	Date of birth [Date of birth]	[dtmBRTH] (DD/MM/YYYY) Req
2.* •	Subject signed informed consent on [Date of informed consent]	[dtmIFC] (DD/MM/YYYY) Req / Req (2015-2023)
3.*	Age (calculated) [read-only] [Age]	[numAGE] N3 (years)
4.* ✓	Sex at birth [Sex at birth]	[IstSXBRTH] [A:M] Male
5.* •	Race [Race]	[IstRC] [A: REPORTED]
6.*	Ethnicity [Ethnicity]	[IstETHNICITY] [A:HISPANIC OR LATINO] Hispanic or Latino [A:NOT HISPANIC OR LATINO] Not Hispanic or Latino [A:NA] Not Available/Not Reported
7.* •	Part in parent study? [Part parent study]	[IstPART] [A:PART B]
8.* •	Did the patient have an interruption between parent study and roll-over to this study? More than a month between completion of parent study treatment and roll-over to this study will be considered an interruption. [Subject initiates dosing]	[IstDAYS] [A:Y]
9.* •	What is the subject's HIV status? [HIV status]	[IstHIV] [A:POSITIVE] O Positive [A:NEGATIVE] Negative
Will	the Whole Blood Clot Formation test (ROTEM) be performed at this site? [sctDN	
10.*	Will the Whole Blood Clot Formation test (ROTEM) be performed at this site? [ROTEM available]	[IstWBCLT] [A:Y]
Sub	ject I dentifiers [sctSUBID]	
11.*	Subject ID (3-digits) [read-only] [Short Subject ID]	[txtSHORTID1] A3
12. •	Site and Subject ID from parent study [Parent study ID]	[txtsubjidps] A7
13.*	Site and Subject ID [hidden] [Site and Subject ID]	[txtSUBJID] A7
14.	For office use only [hidden] [For office use only]	[numEMAIL1] N3
15.	For office use only [hidden] [For office use only]	[numEMAIL2] N3
16.	For office use only [hidden] [For office use only]	[numEMAIL3] N3
Ke No	y: [*] = Item is required [✔] = Source verification required tee: Source verification critical settings made in InForm will override any settings made in Central De	esigner.
Cod	delist Values Tables: Demographic Data and Informed Conse	ent Recording

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DS--Disposition
RELREC--Related Records

	T3SC002: Drug Termination (DRUG TERM) [frmDRUGTERM]		
	udy Drug Completion / Termination [sctDRUGTERM]		
1.	Date of study drug termination [Date of study drug termination]	[dtmDRUGTERM] (DD/MM/YYYY) Req	DSTERM='COMPLETED'. If LSTDRUGCOMP_C = 'COMPLETED' then DSTERM='COMPLET
2. 🗸		[A: ADVERSE EVENT] [A: DEATH] DISTI [A: LOST TO FOLLOW-UF [A: NON-COMPLIANT] [A: PHYSICIAN DECISION [A: PROTOCOL VIOLATION [A: STUDY TERMINATED] [A: SUBJECT WITHDREW [A: OTHER] DISTIENT [A: AT] DISTERM	DISTERM=ADVERSE EVENT Adverse Event Please specify AE#(s) - as recorded on AE form [numAESEQ1] [numAESEQ2] [numAESEQ3] [numAESEQ4] [numAESEQ4] N3
3.	Will the patient continue to use commercially available fitusiran? [Will the patient continue to use commercially available fitusiran?]	[IstCOMMERC] [A:Y] Yes [A:N] No	
4.	For office use only [hidden] [For office use only]	[numEMAIL1] N3	
5.	For office use only [hidden] [For office use only]	[numEMAIL2] N3	
6.	For office use only [hidden] [For office use only]	[numEMAIL3] N3	
	Yey: [*] = Item is required [✓] = Source verification required source verification critical settings made in InForm will override any settings made in Central D	resigner.	

Codelist RefName	Codelist Data Type	Subset	Label	Code	Codelist Item RefName	Data Variable RefName
clCompDisc	String	Jubset	Completed	COMPLETED	citmCompDisc_Comp	IstDRUGCOMP
			Discontinued	DISCONTINUED	citmCompDisc_Disc	
clDISCRS	String		Adverse Event	ADVERSE EVENT	citmDISCRS_AE	IstDISCRS
			Death	DEATH	citmDISCRS_Death	
			Lost to follow-up	LOST TO FOLLOW-UP	citmDISCRS_LostFU	
			Non-Compliance with Study Drug	NON-COMPLIANT	citmDISCRS_NonCompliant	
			Physician Decision	PHYSICIAN DECISION	citmDISCRS_PhysDec	
			Protocol Violation	PROTOCOL VIOLATION	citmDISCRS_PV	
			Study Terminated by Sponsor	STUDY TERMINATED	citmDISCRS_StudyTerm	
			Subject Withdrew Consent	SUBJECT WITHDREW CONSENT	citmDISCRS_WDConsent	
			Other	OTHER	citmDISCRS_Other	
			More than 1 AT measurement <15%	AT	citmDISCRS_AT	
			Temporary discontinuation due to a regional or national emergency	EMERGENCY	citmDISCRS_EMERGENCY	
clYesNo	String		Yes	Υ	citmYesNo_Y	IstCOMMERC

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DS--Disposition

ΑТ	3SC002: Trial Termination (TERM) [frmTERM]	
Tria	al Completion / Discontinuation [sctTERM]	
1.*	Please specify if the subject completed the entire course of the trial as specified in the protocol or if the subject discontinued from the trial early [Completed/Discontinued]	Completed, date of study completion Req V Req V Req V Countrier
2.* •	Will the subject roll-over to the next OLE study? [Will the subject roll-over to the next OLE study?]	[IstROLLOVER] [A:Y] ○ Yes [A:N] ○ No
3.	For office use only [hidden] [For office use only]	[numEMAIL1] N3
4.	For office use only [hidden] [For office use only]	[numEMAIL2] N3
5.	For office use only [hidden] [For office use only]	[numEMAIL3] N3
	ey: [*] = Item is required [🗸] = Source verification required stee: Source verification critical settings made in InForm will override any settings made in Central Des	igner.

Codelist RefName	Codelist Data Type	Subset	Label	Code	Codelist I tem RefName	Data Variable RefName
clCompDisc	String		Completed	COMPLETED	citmCompDisc_Comp	IstCOMPDISC
			Discontinued	DISCONTINUED	citmCompDisc_Disc	
clDISCRS	String		Adverse Event	ADVERSE EVENT	citmDISCRS_AE	IstDISCRS
			Death	DEATH	citmDISCRS_Death	
			Lost to follow-up	LOST TO FOLLOW-UP	citmDISCRS_LostFU	
			Non-Compliance with Study Drug	NON-COMPLIANT	citmDISCRS_NonCompliant	
			Physician Decision	PHYSICIAN DECISION	citmDISCRS_PhysDec	
			Protocol Violation	PROTOCOL VIOLATION	citmDISCRS_PV	
			Study Terminated by Sponsor	STUDY TERMINATED	citmDISCRS_StudyTerm	
			Subject Withdrew Consent	SUBJECT WITHDREW CONSENT	citmDISCRS_WDConsent	
			Other	OTHER	citmDISCRS_Other	
			More than 1 AT measurement <15%	AT	citmDISCRS_AT	
			Temporary discontinuation due to a regional or national emergency	EMERGENCY	citmDISCRS_EMERGENCY	
clYesNo	String		Yes	Υ	citmYesNo_Y	IstROLLOVER