

DS--Disposition

'DSTERM = INFORMED CONSENT OBTAINED



| | | |
|---|---|--|
| AT3SC002: Demographic Data and Informed Consent Recording (DM) [frmDM] | | |
| Demographics [sctDM] | | |
| 1.* ✓ | Date of birth [Date of birth] | [dtmBRTH] (DD/MM/YYYY) Req / Req / Req (1930-2023) |
| 2.* ✓ | Subject signed informed consent on [Date of informed consent] | [dtmIFC] (DD/MM/YYYY) Req / Req / Req (2015-2023) |
| 3.* | Age (calculated) [read-only] [Age] | [numAGE] N3 (years) |
| 4.* ✓ | Sex at birth [Sex at birth] | [IstSXBIRTH] [A:M] Male |
| 5.* ✓ | Race [Race] | [IstRC] [A:REPORTED] [IstRACE] Reported, please specify: [A:WHITE] White [A:BLACK] Black or African-American [A:ASIAN] Asian [A:INDIAN] American Indian or Alaska Native [A:HAWAIIAN] Native Hawaiian or other Pacific Islander [A:OTHER] [txtRACEOTH] Other, please specify: A60 [A:NOT REPORTED] Not Available/Not 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] Part B [A:PART C] Part C [A:PART D] Part D |
| 8.* ✓ | Did the patient have an interruption between parent study and roll-over to this study? <i>More than a month between completion of parent study treatment and roll-over to this study will be considered an interruption.</i> [Subject initiates dosing] | [IstDAYS] [A:Y] Yes [A:N] No |
| 9.* ✓ | What is the subject's HIV status? [HIV status] | [IstHIV] [A:POSITIVE] Positive [A:NEGATIVE] Negative |
| Will the Whole Blood Clot Formation test (ROTEM) be performed at this site? [sctDYN] | | |
| 10.* ✓ | Will the Whole Blood Clot Formation test (ROTEM) be performed at this site? [ROTEM available] | [IstWBCLT] [A:Y] Yes [A:N] No |
| Subject Identifiers [sctSUBID] | | |
| 11.* ✓ | Subject ID (3-digits) [read-only] [Short Subject ID] | [txtSHORTID1] A3 |
| 12. ✓ | Site and Subject ID from parent study [Parent study ID] | [txtSUBJDPS] 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 |
| Key: [*] = Item is required [✓] = Source verification required Note: Source verification critical settings made in InForm will override any settings made in Central Designer. | | |

| | | | | | |
|---|--|--|--|--|--|
| Codelist Values Tables: Demographic Data and Informed Consent Recording | | | | | |
| | | | | | |

DS--Disposition

RELREC--Related Records

AT3SC002: Drug Termination (DRUG TERM) [frmDRUGTERM]

Study Drug Completion / Termination [sctDRUGTERM]

1.*
✓

Date of study drug termination
[Date of study drug termination]

[dtmDRUGTERM] (DD/MM/YYYY)
Req / Req / Req (2015-2023)

DSTERM='COMPLETED'. If LSTDRUGCOMP_C = 'COMPLETED' then DSTERM='COMPLETED'

2.*
✓

Please specify if the patient completed the entire course of ALN-AT3SC treatment as specified in the protocol or if the patient discontinued ALN-AT3SC early
[Please specify if the patient completed the entire course of ALN-AT3SC treatment]

[lstDRUGCOMP]
[A: COMPLETED] Completed DSTERM='COMPLETED'

[A: DISCONTINUED] [lstDISCRS]
Discontinued - please specify the primary reason for early termination
[A: ADVERSE EVENT] [cmpAE] DSTERM= AVDERSE EVENT
Adverse Event
Please specify AE#(s) - as recorded on AE form
[numAESEQ1] [numAESEQ2] [numAESEQ3] [numAESEQ4] [numAESEQ5] [numAESEQ6]

N3

N3

N3

N3

N3

N3

[A: DEATH] Death DSTERM= DEATH
[A: LOST TO FOLLOW-UP] Lost to follow-up DSTERM=FOLLUW-UP
[A: NON-COMPLIANT] Non-Compliance with Study Drug
[A: PHYSICIAN DECISION] [txtDISCPHYSDEC]
Physician Decision, please specify DSTERM= TXTDISCPHYSDEC

A200

[A: PROTOCOL VIOLATION] [txtDISCPV]
Protocol Violation, please specify

A200

[A: STUDY TERMINATED] Study Terminated by Sponsor
[A: SUBJECT WITHDREW CONSENT] Subject Withdrew Consent
[A: OTHER] [txtDISCOTH]
Other, please specify

A200

[A: AT] More than 1 AT measurement <15%
[A: EMERGENCY] Temporary discontinuation due to a regional or national emergency

3.*
✓

Will the patient continue to use commercially available fitusiran?
[Will the patient continue to use commercially available fitusiran?]

[lstCOMMERC]
[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

Key: [*] = Item is required [✓] = Source verification required
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| Codelist Values Tables: Drug Termination | | | | | | |
|--|--------------------|--------|---|--------------------------|-------------------------|-----------------------|
| Codelist RefName | Codelist Data Type | Subset | Label | Code | Codelist Item RefName | Data Variable RefName |
| clCompDisc | String | | Completed | COMPLETED | citmCompDisc_Comp | lstDRUGCOMP |
| | | | Discontinued | DISCONTINUED | citmCompDisc_Disc | |
| clDISCRS | String | | Adverse Event | ADVERSE EVENT | citmDISCRS_AE | lstDISCRS |
| | | | 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 | Y | citmYesNo_Y | lstCOMMERC |

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21-Sep-21

DS--Disposition

AT3SC002: Trial Termination (TERM) [frmTERM]

Trial 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]

[IstCOMPDISC]
[A: COMPLETED]

[dtmCOMP] (DD/MM/YYYY)

Completed, date of study completion | Req ▾ / | Req ▾ / | Req ▾ (2015-2023)

[dtmDISC] (DD/MM/YYYY)

Discontinued, date of study discontinuation | Req ▾ / | Req ▾ / | Req ▾ (2015-2023)

[IstDISCRS]

Discontinued - please specify the primary reason for early termination

[A: ADVERSE EVENT]

[cmpAE]

Adverse Event

DSTERM=AVSER EVENT

Please specify AE#(s) - as recorded on AE form

[numAESEQ1] [numAESEQ2] [numAESEQ3] [numAESEQ4] [numAESEQ5] [numAESEQ6]

N3 N3 N3 N3 N3 N3

Death

DSTERM=DEATH

Lost to follow-up

Non-Compliance with Study Drug

[txtDISCPHYSEDEC]

Physician Decision, please specify

A200

[txtDISCPV]

Protocol Violation, please specify

A200

Study Terminated by Sponsor

Subject Withdrew Consent

[txtDISCOTH]

Other, please specify

A200

More than 1 AT measurement <15%

Temporary discontinuation due to a regional or national emergency

[A: DEATH]

[A: LOST TO FOLLOW-UP]

[A: NON-COMPLIANT]

[A: PHYSICIAN DECISION]

[A: PROTOCOL VIOLATION]

[A: STUDY TERMINATED]

[A: SUBJECT WITHDREW CONSENT]

[A: OTHER]

[A: AT]

[A: EMERGENCY]

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

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| Codelist Values Tables: Trial Termination | | | | | | |
|---|--------------------|--------|---|--------------------------|-------------------------|-----------------------|
| Codelist RefName | Codelist Data Type | Subset | Label | Code | Codelist Item 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 | |
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| | | | 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 | Y | citmYesNo_Y | IstROLLOVER |

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21-Sep-21