

Source Documentation

Visit 5 – Day 35 End of Study Rescheduled On- Site Visit within 30 days of Telehealth Visit

Date of Visit: ____/____/____ Visit Type: On-Site VisitVisit Not Performed: Provide reason if visit completed: _____

(If visit not performed, enter as a protocol violation on the Protocol Violation log page in EDC and provide reason not performed)

DISCHARGE:Has the patient been discharged since Visit 4? No / previously discharged Yes

If yes, discharge date: ____/____/____ Discharge time: _____ am / pm

Facility to which Patient was transferred or discharged:

- Home
- Nursing facility
- Rehab
- Hospice facility
- Other hospital unit: specify: _____

PHYSICAL EXAMINATION:

Body System	Overall interpretation, findings, abnormalities etc.	*Clinically Significant?
Skin (Record Infusion Site Assessment)		
Lungs		
Cardiovascular		
Abdomen		
Upper Extremities		
Lower Extremities		
Other:		

*If any abnormal findings are clinically significant and are changes from baseline, record as AEs

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SIGNATURE SECTION – Physical Examination Completed by:

_____/_____/_____
Printed Name and qualification(s) Signature Date

VITAL SIGNS

Time completed: _____ am / pm

Blood pressure: _____ / _____ mmHg

Location: Arm Leg
 Right Left

Heart rate: _____ bpm

Respiratory rate: _____ breaths / min

Temperature: _____ °F or °C

Temperature Method: Oral Rectal Axillary Tympanic (Ear)
 Cutaneous Infrared Other: _____

Please record any clinically relevant findings on the Adverse Event page

ECG

Was the ECG taken? Yes No: Reason: _____

Time of ECG: _____

Subject Position:

Supine Semi-Recumbent
 Sitting Other: _____

Tracing to be reviewed/assessed as clinically significant or non-clinically significant and initialized and dated by investigator; file original and record any findings judged clinically relevant as AEs Transcribe all findings from the ECG in EDC. QTcF and QTcB will be auto calculated in EDC

INFORMED CONSENT FOR GENETIC TESTING

Has the patient consented to sampling for genetic testing No Yes NA (previously taken)

If yes, was sample collected? No Yes – record detail on re-consent page in EDC

CENTRAL LABORATORY SAMPLES

Date Central Lab Samples drawn, if different from Day 35 visit.

Date of Visit: _____/_____/_____ Time completed: _____ am / pm

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Have central lab samples been completed per protocol? Yes No

If no, record reason and add as a protocol violation: _____

Plasma Biomarker Time completed: _____ am / pm

PK Ticagrelor Total Time completed: _____ am / pm

PK MEDI2452 Time completed: _____ am / pm

PK Ticagrelor Free Time completed: _____ am / pm

Chemistry Time completed: _____ am / pm

Coagulation Time completed: _____ am / pm

Immunogenicity Time completed: _____ am / pm

PREGNANCY TEST

Serum Pregnancy Test Collected No Yes N/A comment: _____

INFUSION SITE ASSESSMENT

Time completed: _____ am / pm

Did the subject have any reactions? No yes- record as Adverse Events

ANTICOAGULANT THERAPY

Did patient restart antiplatelet therapy since last visit?

No - reason not restarted:

- Not indicated
- Continued bleed risk
- Pending surgery or other procedure
- Other _____

Yes – record details on Concomitant medication page

Name of antiplatelet therapy started: _____

CONCOMITANT MEDICATIONS

Any new, or changes, in concomitant medications, Adverse Events or Infusion Site Reactions since last visit? No Yes*

**If yes, complete the applicable worksheet*

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TRIAL COMPLETION

Date of Trial Completion ____/____/____ (date of last contact with patient or date of death)

Did the subject complete the treatment as planned? Yes No if no: please check one of the reasons:

- Adverse event (specify): _____
- Death
- Medication Error
- Withdrawal by Subject
- Other: _____

Did Subject Complete Study as Plan (to day 35)? Yes No if no: please check one of the reasons:

- | | |
|--|---|
| <input type="checkbox"/> Study terminated by sponsor | <input type="checkbox"/> Non-compliance with study drug |
| <input type="checkbox"/> Death | <input type="checkbox"/> Physician decision |
| <input type="checkbox"/> Lost to follow-up | <input type="checkbox"/> Pregnancy |
| <input type="checkbox"/> Informed consent withdrawn | <input type="checkbox"/> Protocol Violation |
| <input type="checkbox"/> Lack of efficacy | <input type="checkbox"/> Adverse event |
| <input type="checkbox"/> Other: _____ | |

VITAL STATUS

Reminders: Patients who withdraw from the study early should be requested to be contacted for vital status at the end of their planned study period (Day 35).

The status of this patient is: Alive Deceased* Lost to follow-up

*If patient has died, complete an SAE report

Date of death: _____ Cause of death: _____

SIGNATURE SECTION –Source Documents Completed by:

_____	_____	____/____/____
Printed Name and qualification(s)	Signature	Date

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PI SIGNATURE SECTION – Principal Investigator signature is required below to signify review:

_____	_____	____/____/____
Printed Name	Signature	Date