

INFORMED CONSENT

- Please insert patient signed ICF

ADVERSE EVENTS

- Adverse Event Source Worksheet V3.0, 06DEC2024
- Adverse Event Source V1.0, 06Dec2024
- Neurological Event V3.0, 06Dec2024
- Systemic Embolism V3.0, 06Dec2024
- Chemistry - Cardiac Enzymes V2.0, 06Dec2024
- Pericardial Effusion Event V3.0, 06Dec2024
- Death V3.0, 06Dec2024

CONCOMITANT MEDICATIONS

- Concomitant Medication V3.0, 06Dec2024

PROTOCOL DEVIATIONS

- Protocol Deviation V3.0, 06Dec2024

SCREENING

- Screening Data Worksheet V5.0, 06Dec2024
- Shared Decision-Making Source V1.0, 06Dec2024
- Inclusion/Exclusion Criteria V3.0, 06Dec2024
- Medical History V6.0, 06Dec2024
- Vital Signs V3.0, 06Dec2024
- Physical Examination – Review of Systems V4.0, 06Dec2024
- CHA₂DS₂VASc V4.0, 06Dec2024
- HAS-BLED V4.0, 06Dec2024
- ECG V2.0, 06Dec2024
- Echo/CT Screening V5.0, 06Dec2024
- NIHSS V2.0, 06Dec2024
- QVSFS V2.0, 06Dec2024
- mRS V2.0, 06Dec2024

PROCEDURE DAY PACKET (SEE BINDER POCKET)

- Sonographer Worksheet V5.0, 08Jun2023
- Procedure V5.0, 06Dec2024
- Additional Procedure Worksheet V1.0, 06Dec2024
- Implant LAA Measurements V3.0, 06Dec2024
- Echo/CT INDEX PROCEDURE V4.0, 06Dec2024
- CLASS Device Deficiency V2.0, 06Dec2024
- CLAAS Implant V2.0, 06Dec2024
- CLAAS Delivery System V2.0, 06Dec2024
- Control Implant V3.0, 06Dec2024
- Patient Population V1.0, 06Dec2024
- Implant Echo Exclusion Criteria V3.0, 06Dec2024
- Procedure Lab Assessments V1.0, 06Dec2024
- **Implant Card**

PRE-DISCHARGE

- Vital Signs V3.0, 06Dec2024
- Echo/CT PRE-DISCHARGE V4.0, 06Dec2024
- NIHSS V2.0, 06Dec2024
- mRS V2.0, 06Dec2024

7 DAY

- Visit Information V4.0, 06Dec2024
- QVSFS V2.0, 06Dec2024

45 DAY

- Visit Information V4.0, 06Dec2024
- QVSFS V2.0, 06Dec2024
- Echo/CT FOLLOW-UP V4.0, 06Dec2024
- Vital Signs V3.0, 06Dec2024

6 MONTH

- Visit Information V4.0, 06Dec2024
- QVSFS V2.0, 06Dec2024

12 MONTH

- Visit Information V4.0, 06Dec2024
- QVSFS V2.0, 06Dec2024
- Echo/CT FOLLOW-UP V4.0, 06Dec2024
- Vital Signs V3.0, 06Dec2024

18 MONTH

- Visit Information V4.0, 06Dec2024
- QVSFS V2.0, 06Dec2024
- NIHSS V2.0, 06Dec2024
- mRS V2.0, 06Dec2024

2 YEAR THROUGH 5 YEAR

- Visit Information V4.0, 06Dec2024
- QVSFS V2.0, 06Dec2024

STUDY EXIT

- Study Exit V2.0, 06Dec2024

INFORMED CONSENT

Please insert patient signed ICF here

ADVERSE EVENTS

| | |
|--|---|
| AE EDC Event Number | |
| Status of Adverse Event | <input type="checkbox"/> New adverse event <input type="checkbox"/> Worsening of pre-existing condition |
| AE Event Term | |
| AE Description | |
| Suspected Cause | |
| Date of Site Awareness of AE | ___/___/___ (DD/MMM/YYYY) |
| Date Sponsor Notified of AE | ___/___/___ (DD/MMM/YYYY) |
| AE Onset Date | ___/___/___ (DD/MMM/YYYY) |
| Severity <i>Refer to Protocol Appendix A 21.1.1</i> | <input type="checkbox"/> Mild <input type="checkbox"/> Moderate <input type="checkbox"/> Severe |
| Action Taken <i>(Check all that apply)</i> | <input type="checkbox"/> None <input type="checkbox"/> Hospitalization < 24 hours <input type="checkbox"/> Hospitalization > 24 hours <input type="checkbox"/> Study Medication prescribed <input type="checkbox"/> Study Medication dose changed <input type="checkbox"/> Study Medication stopped <input type="checkbox"/> Percutaneous intervention Specify: _____ <input type="checkbox"/> Surgical intervention Specify: _____ <input type="checkbox"/> Transfusion Number of units of red blood cells (xx): _____ Number of units of platelets or FFP (xx): _____ <input type="checkbox"/> Other, specify: _____ |

| | | | | | | | | | | | | | |
|--|---|---|---|---|---|---|---|--|---|--|---|---|---|
| <p>Were any of the following performed? <i>(Please ensure any images are uploaded into Imaging Module)</i></p> | <p><input type="checkbox"/> None <input type="checkbox"/> Cardiac Angiography <input type="checkbox"/> Cardiac MRI <input type="checkbox"/> Cardiac Echo/CT <input type="checkbox"/> Brain Imaging <input type="checkbox"/> ECG <input type="checkbox"/> Ultrasound <input type="checkbox"/> Pathologic Examination</p> | | | | | | | | | | | | |
| <p>Is this a Serious Adverse Event (SAE)?</p> | <p><input type="checkbox"/> Yes <input type="checkbox"/> No</p> | | | | | | | | | | | | |
| | <table border="1" style="width: 100%;"> <tr> <td style="width: 70%; text-align: center;">Led to subject death <i>(If yes, complete Death Form)</i></td> <td style="width: 30%; vertical-align: top;"> <input type="checkbox"/> Yes <input type="checkbox"/> No </td> </tr> <tr> <td style="text-align: center;">A life-threatening illness or injury</td> <td style="vertical-align: top;"> <input type="checkbox"/> Yes <input type="checkbox"/> No </td> </tr> <tr> <td style="text-align: center;">A permanent impairment of a body structure or body function, including chronic diseases</td> <td style="vertical-align: top;"> <input type="checkbox"/> Yes <input type="checkbox"/> No </td> </tr> <tr> <td style="text-align: center;">A medical or surgical intervention to prevent life-threatening illness, injury or permanent impairment of body structure or function</td> <td style="vertical-align: top;"> <input type="checkbox"/> Yes <input type="checkbox"/> No </td> </tr> <tr> <td style="text-align: center;">In-subject hospitalization or prolongation of existing hospitalization</td> <td style="vertical-align: top;"> <input type="checkbox"/> Yes <input type="checkbox"/> No </td> </tr> <tr> <td style="text-align: center;">Fetal distress, fetal death or congenital anomaly or birth defect including physical or mental impairment</td> <td style="vertical-align: top;"> <input type="checkbox"/> Yes <input type="checkbox"/> No </td> </tr> </table> | Led to subject death <i>(If yes, complete Death Form)</i> | <input type="checkbox"/> Yes <input type="checkbox"/> No | A life-threatening illness or injury | <input type="checkbox"/> Yes <input type="checkbox"/> No | A permanent impairment of a body structure or body function, including chronic diseases | <input type="checkbox"/> Yes <input type="checkbox"/> No | A medical or surgical intervention to prevent life-threatening illness, injury or permanent impairment of body structure or function | <input type="checkbox"/> Yes <input type="checkbox"/> No | In-subject hospitalization or prolongation of existing hospitalization | <input type="checkbox"/> Yes <input type="checkbox"/> No | Fetal distress, fetal death or congenital anomaly or birth defect including physical or mental impairment | <input type="checkbox"/> Yes <input type="checkbox"/> No |
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| In-subject hospitalization or prolongation of existing hospitalization | <input type="checkbox"/> Yes <input type="checkbox"/> No | | | | | | | | | | | | |
| Fetal distress, fetal death or congenital anomaly or birth defect including physical or mental impairment | <input type="checkbox"/> Yes <input type="checkbox"/> No | | | | | | | | | | | | |
| <p>Is Event cardiovascular or neurological in etiology?</p> | <p><input type="checkbox"/> Yes, Cardiovascular <input type="checkbox"/> Yes, Neurological <input type="checkbox"/> No</p> | | | | | | | | | | | | |
| <p>Adverse Event of Special Interest?</p> | <table border="1" style="width: 100%;"> <tr> <td style="width: 20%; vertical-align: top;"> <input type="checkbox"/> Yes <input type="checkbox"/> No </td> <td style="width: 20%; vertical-align: top;"> <p>If yes, check all that apply</p> </td> <td style="width: 60%; vertical-align: top;"> <input type="checkbox"/> Bleeding Event <input type="checkbox"/> Myocardial Infarction Were cardiac enzymes drawn? <input type="checkbox"/> Yes <i>(Complete Cardiac Enzyme Form)</i> <input type="checkbox"/> No <input type="checkbox"/> Unknown <input type="checkbox"/> Pericardial Effusion <input type="checkbox"/> Neurological Event <input type="checkbox"/> Vascular Complication <input type="checkbox"/> Systemic Embolization <input type="checkbox"/> Device Embolization </td> </tr> </table> | <input type="checkbox"/> Yes <input type="checkbox"/> No | <p>If yes, check all that apply</p> | <input type="checkbox"/> Bleeding Event <input type="checkbox"/> Myocardial Infarction Were cardiac enzymes drawn? <input type="checkbox"/> Yes <i>(Complete Cardiac Enzyme Form)</i> <input type="checkbox"/> No <input type="checkbox"/> Unknown <input type="checkbox"/> Pericardial Effusion <input type="checkbox"/> Neurological Event <input type="checkbox"/> Vascular Complication <input type="checkbox"/> Systemic Embolization <input type="checkbox"/> Device Embolization | | | | | | | | | |
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|-----------------------------------|--|--|
| Related to study device? | <input type="checkbox"/> Yes | |
| | <input type="checkbox"/> No | |
| | Relationship to implant? | <input type="checkbox"/> Not related <input type="checkbox"/> Possible <input type="checkbox"/> Probable <input type="checkbox"/> Causal relationship |
| | Relationship to access sheath? | <input type="checkbox"/> Not related <input type="checkbox"/> Possible <input type="checkbox"/> Probable <input type="checkbox"/> Causal relationship |
| | Relationship to delivery system? | <input type="checkbox"/> Not related <input type="checkbox"/> Possible <input type="checkbox"/> Probable <input type="checkbox"/> Causal relationship |
| Relationship to hydraulic loader? | <input type="checkbox"/> Not related <input type="checkbox"/> Possible <input type="checkbox"/> Probable <input type="checkbox"/> Causal relationship <input type="checkbox"/> N/A – Subject did not receive CLAAS | |
| Related to Study Procedure? | <input type="checkbox"/> Yes | |
| | <input type="checkbox"/> Possible <input type="checkbox"/> Probable <input type="checkbox"/> Causal relationship <input type="checkbox"/> No, Not related | |
| Related to Study Medication? | <input type="checkbox"/> Yes | |
| | <input type="checkbox"/> Possible <input type="checkbox"/> Probable <input type="checkbox"/> Causal relationship <input type="checkbox"/> No, Not related | |

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| AE Outcome | <input type="checkbox"/> Recovered/Resolved <input type="checkbox"/> Recovered/Resolved with Sequelae <input type="checkbox"/> Ongoing <input type="checkbox"/> Fatal (<i>Complete Death Form and Study Exit Forms</i>) <input type="checkbox"/> Ongoing at end of study |
| If recovered/resolved, describe how resolution was confirmed: | |
| AE End Date | ___/___/___ (DD/MMM/YYYY) |

RC/ RA Signature

___/___/___
Date (DD/MMM/YYYY)

Investigator Signature

___/___/___
Date (DD/MMM/YYYY)

Site Number: _____ Subject ID: _____

| AE # | AE Term | AE Status | Cause | Date Aware | Date Entered | Severity | Serious | Onset Date | Resolved Date |
|--------------------------------|---------------------------------|---|---------------------------------------|--|---|--|--|--|--|
| | | <input type="checkbox"/> New <input type="checkbox"/> Pre-Existing | | | | <input type="checkbox"/> Mild <input type="checkbox"/> Moderate <input type="checkbox"/> Severe | <input type="checkbox"/> No <input type="checkbox"/> Yes * Circle all that apply 1 2 3 4 5 6 7 | | |
| | Relationship to implant? | <input type="checkbox"/> Not related <input type="checkbox"/> Possible <input type="checkbox"/> Probable <input type="checkbox"/> Causal relationship | Relationship to access sheath? | <input type="checkbox"/> Not related <input type="checkbox"/> Possible <input type="checkbox"/> Probable <input type="checkbox"/> Causal relationship | Relationship to delivery system? | <input type="checkbox"/> Not related <input type="checkbox"/> Possible <input type="checkbox"/> Probable <input type="checkbox"/> Causal relationship | Relationship to Study Procedure? | <input type="checkbox"/> Not related <input type="checkbox"/> Possible <input type="checkbox"/> Probable <input type="checkbox"/> Causal relationship | Relationship to Study Medication? |
| Investigator Signature: | | | | | | | | | |
| | | <input type="checkbox"/> New <input type="checkbox"/> Pre-Exist | | | | <input type="checkbox"/> Mild <input type="checkbox"/> Moderate <input type="checkbox"/> Severe | <input type="checkbox"/> No <input type="checkbox"/> Yes * Circle all that apply 1 2 3 4 5 6 7 | | |
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|-----------------------------------|--|--|
| Related to study device? | <input type="checkbox"/> Yes | |
| | <input type="checkbox"/> No | |
| | Relationship to implant? | <input type="checkbox"/> Not related <input type="checkbox"/> Possible <input type="checkbox"/> Probable <input type="checkbox"/> Causal relationship |
| | Relationship to access sheath? | <input type="checkbox"/> Not related <input type="checkbox"/> Possible <input type="checkbox"/> Probable <input type="checkbox"/> Causal relationship |
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| Related to Study Procedure? | <input type="checkbox"/> Yes | |
| | <input type="checkbox"/> Possible <input type="checkbox"/> Probable <input type="checkbox"/> Causal relationship <input type="checkbox"/> No, Not related | |
| Related to Study Medication? | <input type="checkbox"/> Yes | |
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| AE Outcome | <input type="checkbox"/> Recovered/Resolved <input type="checkbox"/> Recovered/Resolved with Sequelae <input type="checkbox"/> Ongoing <input type="checkbox"/> Fatal (<i>Complete Death Form and Study Exit Forms</i>) <input type="checkbox"/> Ongoing at end of study |
| If recovered/resolved, describe how resolution was confirmed: | |
| AE End Date | ___/___/___ (DD/MMM/YYYY) |

RC/ RA Signature

___/___/___
Date (DD/MMM/YYYY)

Investigator Signature

___/___/___
Date (DD/MMM/YYYY)

Site Number: _____ Subject ID: _____

| AE # | AE Term | AE Status | Cause | Date Aware | Date Entered | Severity | Serious | Onset Date | Resolved Date |
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| | Relationship to implant? | <input type="checkbox"/> Not related <input type="checkbox"/> Possible <input type="checkbox"/> Probable <input type="checkbox"/> Causal relationship | Relationship to access sheath? | <input type="checkbox"/> Not related <input type="checkbox"/> Possible <input type="checkbox"/> Probable <input type="checkbox"/> Causal relationship | Relationship to delivery system? | <input type="checkbox"/> Not related <input type="checkbox"/> Possible <input type="checkbox"/> Probable <input type="checkbox"/> Causal relationship | Relationship to Study Procedure? | <input type="checkbox"/> Not related <input type="checkbox"/> Possible <input type="checkbox"/> Probable <input type="checkbox"/> Causal relationship | Relationship to Study Medication? |
| Investigator Signature: | | | | | | | | | |
| | | <input type="checkbox"/> New <input type="checkbox"/> Pre-Exist | | | | <input type="checkbox"/> Mild <input type="checkbox"/> Moderate <input type="checkbox"/> Severe | <input type="checkbox"/> No <input type="checkbox"/> Yes <i>* Circle all that apply</i> 1 2 3 4 5 6 7 | | |
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| Investigator Signature: | | | | | | | | | |

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| AE EDC Event Number | |
| Status of Adverse Event | <input type="checkbox"/> New adverse event <input type="checkbox"/> Worsening of pre-existing condition |
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| Severity <i>Refer to Protocol Appendix A 21.1.1</i> | <input type="checkbox"/> Mild <input type="checkbox"/> Moderate <input type="checkbox"/> Severe |
| Action Taken <i>(Check all that apply)</i> | <input type="checkbox"/> None <input type="checkbox"/> Hospitalization < 24 hours <input type="checkbox"/> Hospitalization > 24 hours <input type="checkbox"/> Study Medication prescribed <input type="checkbox"/> Study Medication dose changed <input type="checkbox"/> Study Medication stopped <input type="checkbox"/> Percutaneous intervention Specify: _____ <input type="checkbox"/> Surgical intervention Specify: _____ <input type="checkbox"/> Transfusion Number of units of red blood cells (xx): _____ Number of units of platelets or FFP (xx): _____ <input type="checkbox"/> Other, specify: _____ |

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| <p>Were any of the following performed? <i>(Please ensure any images are uploaded into Imaging Module)</i></p> | <input type="checkbox"/> None <input type="checkbox"/> Cardiac Angiography <input type="checkbox"/> Cardiac MRI <input type="checkbox"/> Cardiac Echo/CT <input type="checkbox"/> Brain Imaging <input type="checkbox"/> ECG <input type="checkbox"/> Ultrasound <input type="checkbox"/> Pathologic Examination | | | | | | | | | | | | |
| <p>Is this a Serious Adverse Event (SAE)?</p> | <input type="checkbox"/> Yes <input type="checkbox"/> No | | | | | | | | | | | | |
| | <table border="1" style="width: 100%;"> <tr> <td style="width: 70%; text-align: center;">Led to subject death <i>(If yes, complete Death Form)</i></td> <td style="width: 30%; vertical-align: top;"> <input type="checkbox"/> Yes <input type="checkbox"/> No </td> </tr> <tr> <td style="text-align: center;">A life-threatening illness or injury</td> <td style="vertical-align: top;"> <input type="checkbox"/> Yes <input type="checkbox"/> No </td> </tr> <tr> <td style="text-align: center;">A permanent impairment of a body structure or body function, including chronic diseases</td> <td style="vertical-align: top;"> <input type="checkbox"/> Yes <input type="checkbox"/> No </td> </tr> <tr> <td style="text-align: center;">A medical or surgical intervention to prevent life-threatening illness, injury or permanent impairment of body structure or function</td> <td style="vertical-align: top;"> <input type="checkbox"/> Yes <input type="checkbox"/> No </td> </tr> <tr> <td style="text-align: center;">In-subject hospitalization or prolongation of existing hospitalization</td> <td style="vertical-align: top;"> <input type="checkbox"/> Yes <input type="checkbox"/> No </td> </tr> <tr> <td style="text-align: center;">Fetal distress, fetal death or congenital anomaly or birth defect including physical or mental impairment</td> <td style="vertical-align: top;"> <input type="checkbox"/> Yes <input type="checkbox"/> No </td> </tr> </table> | Led to subject death <i>(If yes, complete Death Form)</i> | <input type="checkbox"/> Yes <input type="checkbox"/> No | A life-threatening illness or injury | <input type="checkbox"/> Yes <input type="checkbox"/> No | A permanent impairment of a body structure or body function, including chronic diseases | <input type="checkbox"/> Yes <input type="checkbox"/> No | A medical or surgical intervention to prevent life-threatening illness, injury or permanent impairment of body structure or function | <input type="checkbox"/> Yes <input type="checkbox"/> No | In-subject hospitalization or prolongation of existing hospitalization | <input type="checkbox"/> Yes <input type="checkbox"/> No | Fetal distress, fetal death or congenital anomaly or birth defect including physical or mental impairment | <input type="checkbox"/> Yes <input type="checkbox"/> No |
| Led to subject death <i>(If yes, complete Death Form)</i> | <input type="checkbox"/> Yes <input type="checkbox"/> No | | | | | | | | | | | | |
| A life-threatening illness or injury | <input type="checkbox"/> Yes <input type="checkbox"/> No | | | | | | | | | | | | |
| A permanent impairment of a body structure or body function, including chronic diseases | <input type="checkbox"/> Yes <input type="checkbox"/> No | | | | | | | | | | | | |
| A medical or surgical intervention to prevent life-threatening illness, injury or permanent impairment of body structure or function | <input type="checkbox"/> Yes <input type="checkbox"/> No | | | | | | | | | | | | |
| In-subject hospitalization or prolongation of existing hospitalization | <input type="checkbox"/> Yes <input type="checkbox"/> No | | | | | | | | | | | | |
| Fetal distress, fetal death or congenital anomaly or birth defect including physical or mental impairment | <input type="checkbox"/> Yes <input type="checkbox"/> No | | | | | | | | | | | | |
| <p>Is Event cardiovascular or neurological in etiology?</p> | <input type="checkbox"/> Yes, Cardiovascular <input type="checkbox"/> Yes, Neurological <input type="checkbox"/> No | | | | | | | | | | | | |
| <p>Adverse Event of Special Interest?</p> | <table border="1" style="width: 100%;"> <tr> <td style="width: 20%; vertical-align: top;"> <input type="checkbox"/> Yes <input type="checkbox"/> No </td> <td style="width: 20%; vertical-align: top;"> <p>If yes, check all that apply</p> </td> <td style="width: 60%; vertical-align: top;"> <input type="checkbox"/> Bleeding Event <input type="checkbox"/> Myocardial Infarction Were cardiac enzymes drawn? <input type="checkbox"/> Yes <i>(Complete Cardiac Enzyme Form)</i> <input type="checkbox"/> No <input type="checkbox"/> Unknown <input type="checkbox"/> Pericardial Effusion <input type="checkbox"/> Neurological Event <input type="checkbox"/> Vascular Complication <input type="checkbox"/> Systemic Embolization <input type="checkbox"/> Device Embolization </td> </tr> </table> | <input type="checkbox"/> Yes <input type="checkbox"/> No | <p>If yes, check all that apply</p> | <input type="checkbox"/> Bleeding Event <input type="checkbox"/> Myocardial Infarction Were cardiac enzymes drawn? <input type="checkbox"/> Yes <i>(Complete Cardiac Enzyme Form)</i> <input type="checkbox"/> No <input type="checkbox"/> Unknown <input type="checkbox"/> Pericardial Effusion <input type="checkbox"/> Neurological Event <input type="checkbox"/> Vascular Complication <input type="checkbox"/> Systemic Embolization <input type="checkbox"/> Device Embolization | | | | | | | | | |
| <input type="checkbox"/> Yes <input type="checkbox"/> No | <p>If yes, check all that apply</p> | <input type="checkbox"/> Bleeding Event <input type="checkbox"/> Myocardial Infarction Were cardiac enzymes drawn? <input type="checkbox"/> Yes <i>(Complete Cardiac Enzyme Form)</i> <input type="checkbox"/> No <input type="checkbox"/> Unknown <input type="checkbox"/> Pericardial Effusion <input type="checkbox"/> Neurological Event <input type="checkbox"/> Vascular Complication <input type="checkbox"/> Systemic Embolization <input type="checkbox"/> Device Embolization | | | | | | | | | | | |

| | | |
|-----------------------------------|--|--|
| Related to study device? | <input type="checkbox"/> Yes | |
| | <input type="checkbox"/> No | |
| | Relationship to implant? | <input type="checkbox"/> Not related <input type="checkbox"/> Possible <input type="checkbox"/> Probable <input type="checkbox"/> Causal relationship |
| | Relationship to access sheath? | <input type="checkbox"/> Not related <input type="checkbox"/> Possible <input type="checkbox"/> Probable <input type="checkbox"/> Causal relationship |
| | Relationship to delivery system? | <input type="checkbox"/> Not related <input type="checkbox"/> Possible <input type="checkbox"/> Probable <input type="checkbox"/> Causal relationship |
| Relationship to hydraulic loader? | <input type="checkbox"/> Not related <input type="checkbox"/> Possible <input type="checkbox"/> Probable <input type="checkbox"/> Causal relationship <input type="checkbox"/> N/A – Subject did not receive CLAAS | |
| Related to Study Procedure? | <input type="checkbox"/> Yes | |
| | <input type="checkbox"/> Possible <input type="checkbox"/> Probable <input type="checkbox"/> Causal relationship <input type="checkbox"/> No, Not related | |
| Related to Study Medication? | <input type="checkbox"/> Yes | |
| | <input type="checkbox"/> Possible <input type="checkbox"/> Probable <input type="checkbox"/> Causal relationship <input type="checkbox"/> No, Not related | |

| | |
|---|--|
| AE Outcome | <input type="checkbox"/> Recovered/Resolved <input type="checkbox"/> Recovered/Resolved with Sequelae <input type="checkbox"/> Ongoing <input type="checkbox"/> Fatal (<i>Complete Death Form and Study Exit Forms</i>) <input type="checkbox"/> Ongoing at end of study |
| If recovered/resolved, describe how resolution was confirmed: | |
| AE End Date | ___/___/___ (DD/MMM/YYYY) |

RC/ RA Signature

___/___/___
Date (DD/MMM/YYYY)

Investigator Signature

___/___/___
Date (DD/MMM/YYYY)

Site Number: _____ Subject ID: _____

| | | | |
|--|--|--|--|
| Related AE #: _____ | | Related AE Term: _____ | |
| Neurological deficit <i>(Check all that apply)</i> | | <input type="checkbox"/> Altered mental status <input type="checkbox"/> Coordination <input type="checkbox"/> Decreased level of consciousness <input type="checkbox"/> Memory <input type="checkbox"/> Motor <input type="checkbox"/> Sensory <input type="checkbox"/> Speech <input type="checkbox"/> Swallowing <input type="checkbox"/> Visual deficit <input type="checkbox"/> Other, specify: _____ | |
| Location of neurological deficit <i>(Check all that apply)</i> | | <input type="checkbox"/> Cranial nerves/face <input type="checkbox"/> Arm <input type="checkbox"/> Leg <input type="checkbox"/> Trunk | |
| Side of neurological deficit | | <input type="checkbox"/> Left <input type="checkbox"/> Right <input type="checkbox"/> Bilateral | |
| Was a neurological consult performed? | | <input type="checkbox"/> Yes <input type="checkbox"/> No | |
| Were neurological assessments performed? | | <input type="checkbox"/> Yes <input type="checkbox"/> No | |
| | | If yes, specify <i>(Select all that apply)</i> | |
| | | <input type="checkbox"/> Neuro exam and evaluation <input type="checkbox"/> mRS <i>(Enter mRS CRF)</i> <input type="checkbox"/> NIHSS <i>(Enter NIHSS CRF)</i> <input type="checkbox"/> QVSFS <i>(Enter QVSFS CRF)</i> | |

Site Personnel Signature

____/____/_____
Date (DD/MMM/YYYY)

| | |
|---|--|
| Related AE #: _____ | Related AE Term: |
| Source of systemic embolization <i>(Select only one)</i> | <input type="checkbox"/> Cardioembolic <input type="checkbox"/> Other, specify: _____ <input type="checkbox"/> Unknown |
| End organ damage? | <input type="checkbox"/> Yes <input type="checkbox"/> No <input type="checkbox"/> Unknown |
| If yes, specify <i>(Select all that apply)</i> | <input type="checkbox"/> Pulmonary circulation/lungs <input type="checkbox"/> Coronary circulation <input type="checkbox"/> Visceral-mesenteric <input type="checkbox"/> Peripheral vasculature <input type="checkbox"/> Upper extremity <input type="checkbox"/> Lower extremity |

Note: If utilizing as source (no other source exists)- form should be signed by Site Investigator

Site Personnel Signature

___/___/_____
Date (DD/MMM/YYYY)

Source Data Transfer Tool

Site Number: _____ Subject ID: _____

Note: It is not required to complete this source worksheet if lab reports are readily available.

| | | | |
|--|---|-------------|--|
| Was the cardiac enzyme sample collected? | <input type="checkbox"/> Yes <input type="checkbox"/> No | | |
| Date of Lab | ___ / ___ / ___ (DD/MMM/YYYY) | | |
| Time of Lab (24 HR) | __ : __ | | |
| Was CK collected? | <input type="checkbox"/> Yes <input type="checkbox"/> No | Total CK: | |
| | | Unit: | <input type="checkbox"/> U/L <input type="checkbox"/> Other, specify: _____ |
| Was CK-MB collected? | <input type="checkbox"/> Yes <input type="checkbox"/> No | CK-MB: | |
| | | Unit: | <input type="checkbox"/> ng/mL <input type="checkbox"/> Other, specify: _____ |
| Was Troponin I collected? | <input type="checkbox"/> Yes <input type="checkbox"/> No | Troponin I: | |
| | | Unit: | <input type="checkbox"/> ng/mL <input type="checkbox"/> Other, specify: _____ |
| Was Troponin T collected? | <input type="checkbox"/> Yes <input type="checkbox"/> No | Troponin T: | |
| | | Unit: | <input type="checkbox"/> ng/mL <input type="checkbox"/> Other, specify: _____ |

Site Personnel Signature

___ / ___ / ___
Date (DD/MMM/YYYY)

| | | |
|---|---|--|
| Related AE #: | | |
| Type of Pericardial Effusion | <input type="checkbox"/> Circumferential <input type="checkbox"/> Non-Circumferential | |
| Size of Pericardial Effusion | <input type="checkbox"/> Trivial <input type="checkbox"/> Small (<1 cm) <input type="checkbox"/> Moderate (1-2 cm) <input type="checkbox"/> Large (>2 cm and <5cm) <input type="checkbox"/> Large (>5cm) | |
| Persistent Hypotension requiring pressor support? | <input type="checkbox"/> Yes <input type="checkbox"/> No <input type="checkbox"/> Unknown | |
| Pericardial Drainage Attempted? | <input type="checkbox"/> Yes <input type="checkbox"/> No <input type="checkbox"/> Unknown | If yes, <input type="checkbox"/> Surgical Intervention <input type="checkbox"/> Pericardiocentesis |
| | | <input type="checkbox"/> Successful Volume Removed: _____ ml Type Removed: <input type="checkbox"/> Blood <input type="checkbox"/> Blood Tinged <input type="checkbox"/> Straw Colored <input type="checkbox"/> Unsuccessful (No Volume Removed) |
| Time of occurrence? | <input type="checkbox"/> Intraprocedural <input type="checkbox"/> Acute (< or = 48 hours post-procedure) <input type="checkbox"/> Late (>48 hours post-procedure through 45 days) <input type="checkbox"/> Very Late (>45 days post procedure) | |

Site Personnel Signature

___/___/___
Date (DD/MMM/YYYY)

All deaths are considered cardiac unless an unequivocal noncardiac cause can be established. Specifically, **any unexpected death even in patients with coexisting potentially fatal noncardiac disease** (e.g., cancer, infection) **should be classified as cardiac.**

This language is from the Protocol Appendix definition around Mortality.

| | |
|-------------------------------|--|
| Date of Death | ___ / ___ / ___ (DD/MMM/YYYY) |
| Primary cause of death | <input type="checkbox"/> Cardiovascular death Any death due to proximate cardiac cause (e.g., MI, low output failure, fatal arrhythmia), unwitnessed death and death of unknown cause, AND all procedure-related deaths, including those related to concomitant treatment, should be classified as such. Death caused by noncoronary vascular causes, such as cerebrovascular disease, pulmonary embolism, ruptured aortic, aneurysm, dissecting aneurysm, or other vascular diseases should be classified as such |
| | <input type="checkbox"/> Non-Cardiovascular death Any death not covered by Cardiovascular causes- such as death caused by infection, malignancy, sepsis, pulmonary causes, accident, suicide or trauma |
| | <input type="checkbox"/> Unknown/Not Available Should only be selected if death certificate, or autopsy is not available AND the investigator is not comfortable classifying as defined above given the information available at the time of death. |
| Was an autopsy performed? | <input type="checkbox"/> Yes, date of autopsy: ___ / ___ / ___ (DD/MMM/YYYY) <input type="checkbox"/> No <input type="checkbox"/> Unknown |
| Is subject autopsy available? | <input type="checkbox"/> No <input type="checkbox"/> Yes <i>Please provide source documents for this event to Safety including autopsy if available</i> |

If utilizing as source, (no autopsy/death certificate source) is available, form should be signed by Site Investigator

_____ / _____ / _____
Site Personnel Signature **Date (DD/MMM/YYYY)**

CONCOMITANT MEDICATIONS



CONFORM Concomitant Medication

Source Data Transfer Tool

Site Number: _____ Subject ID: _____

*Note: Data collection includes the use of antiplatelet, anticoagulation and prophylactic antibiotic medication only.
Collect prescribed antiplatelet, anticoagulant, and P2Y12 therapies from subject's relevant medical history through study exit.*

| Med Number in EDC | Med Name | Dose/Units/Route | Frequency | Start Date | Stop Date |
|-------------------|----------|------------------|-----------|------------|-----------|
| 1 | | | | | |
| 2 | | | | | |
| 3 | | | | | |
| 4 | | | | | |
| 5 | | | | | |
| 6 | | | | | |
| 7 | | | | | |
| 8 | | | | | |
| 9 | | | | | |
| 10 | | | | | |



CONFORM Concomitant Medication

Source Data Transfer Tool

Site Number: _____ Subject ID: _____

| Med Number in EDC | Med Name | Dose/Units/Route | Frequency | Start Date | Stop Date |
|-------------------|----------|------------------|-----------|------------|-----------|
| 11 | | | | | |
| 12 | | | | | |
| 13 | | | | | |
| 14 | | | | | |
| 15 | | | | | |
| 16 | | | | | |
| 17 | | | | | |
| 18 | | | | | |
| 19 | | | | | |
| 20 | | | | | |



CONFORM Concomitant Medication

Source Data Transfer Tool

Site Number: _____ Subject ID: _____

| Time point Con Meds reviewed/ updated by RC | Site Personnel Signature | Date of Review |
|---|--------------------------|----------------|
| SCREENING | | |
| PROCEDURE | | |
| PRE-DISCHARGE | | |
| 7 DAY | | |
| 45 DAY | | |
| 6 MONTH | | |
| 1 YR | | |
| 18 MONTH | | |
| 2 YEAR | | |
| 3 YEAR | | |
| 4 YEAR | | |
| 5 YEAR | | |
| STUDY EXIT | | |

Data collection includes the use of antiplatelet, anticoagulation and prophylactic antibiotic medication only.



CONFORM Concomitant Medication

Source Data Transfer Tool

Site Number: _____ Subject ID: _____

*Note: Data collection includes the use of antiplatelet, anticoagulation and prophylactic antibiotic medication only.
Collect prescribed antiplatelet, anticoagulant, and P2Y12 therapies from subject's relevant medical history through study exit.*

| Med Number in EDC | Med Name | Dose/Units/Route | Frequency | Start Date | Stop Date |
|-------------------|----------|------------------|-----------|------------|-----------|
| 1 | | | | | |
| 2 | | | | | |
| 3 | | | | | |
| 4 | | | | | |
| 5 | | | | | |
| 6 | | | | | |
| 7 | | | | | |
| 8 | | | | | |
| 9 | | | | | |
| 10 | | | | | |



CONFORM Concomitant Medication

Source Data Transfer Tool

Site Number: _____ Subject ID: _____

| Med Number in EDC | Med Name | Dose/Units/Route | Frequency | Start Date | Stop Date |
|-------------------|----------|------------------|-----------|------------|-----------|
| 11 | | | | | |
| 12 | | | | | |
| 13 | | | | | |
| 14 | | | | | |
| 15 | | | | | |
| 16 | | | | | |
| 17 | | | | | |
| 18 | | | | | |
| 19 | | | | | |
| 20 | | | | | |



CONFORM Concomitant Medication

Source Data Transfer Tool

Site Number: _____ Subject ID: _____

| Time point Con Meds reviewed/ updated by RC | Site Personnel Signature | Date of Review |
|---|--------------------------|----------------|
| SCREENING | | |
| PROCEDURE | | |
| PRE-DISCHARGE | | |
| 7 DAY | | |
| 45 DAY | | |
| 6 MONTH | | |
| 1 YR | | |
| 18 MONTH | | |
| 2 YEAR | | |
| 3 YEAR | | |
| 4 YEAR | | |
| 5 YEAR | | |
| STUDY EXIT | | |

Data collection includes the use of antiplatelet, anticoagulation and prophylactic antibiotic medication only.



CONFORM Concomitant Medication

Source Data Transfer Tool

Site Number: _____ Subject ID: _____

*Note: Data collection includes the use of antiplatelet, anticoagulation and prophylactic antibiotic medication only.
Collect prescribed antiplatelet, anticoagulant, and P2Y12 therapies from subject's relevant medical history through study exit.*

| Med Number in EDC | Med Name | Dose/Units/Route | Frequency | Start Date | Stop Date |
|-------------------|----------|------------------|-----------|------------|-----------|
| 1 | | | | | |
| 2 | | | | | |
| 3 | | | | | |
| 4 | | | | | |
| 5 | | | | | |
| 6 | | | | | |
| 7 | | | | | |
| 8 | | | | | |
| 9 | | | | | |
| 10 | | | | | |



CONFORM Concomitant Medication

Source Data Transfer Tool

Site Number: _____ Subject ID: _____

| Med Number in EDC | Med Name | Dose/Units/Route | Frequency | Start Date | Stop Date |
|-------------------|----------|------------------|-----------|------------|-----------|
| 11 | | | | | |
| 12 | | | | | |
| 13 | | | | | |
| 14 | | | | | |
| 15 | | | | | |
| 16 | | | | | |
| 17 | | | | | |
| 18 | | | | | |
| 19 | | | | | |
| 20 | | | | | |



CONFORM Concomitant Medication

Source Data Transfer Tool

Site Number: _____ Subject ID: _____

| Time point Con Meds reviewed/ updated by RC | Site Personnel Signature | Date of Review |
|---|--------------------------|----------------|
| SCREENING | | |
| PROCEDURE | | |
| PRE-DISCHARGE | | |
| 7 DAY | | |
| 45 DAY | | |
| 6 MONTH | | |
| 1 YR | | |
| 18 MONTH | | |
| 2 YEAR | | |
| 3 YEAR | | |
| 4 YEAR | | |
| 5 YEAR | | |
| STUDY EXIT | | |

Data collection includes the use of antiplatelet, anticoagulation and prophylactic antibiotic medication only.

PROTOCOL DEVIATIONS

Note: Please complete only one deviation per form.

PD # in EDC _____

| | | |
|--|---|---|
| Date of Deviation | ___ / ___ / ___ (DD/MMM/YYYY) | |
| Date of Site Awareness | ___ / ___ / ___ (DD/MMM/YYYY) | |
| Time Period of Deviation | <input type="checkbox"/> Screening <input type="checkbox"/> Index Procedure <input type="checkbox"/> Discharge <input type="checkbox"/> Day 7 <input type="checkbox"/> Day 45 <input type="checkbox"/> 6 Months <input type="checkbox"/> 12 Months <input type="checkbox"/> 18 Months | <input type="checkbox"/> 2 Year <input type="checkbox"/> 3 Year <input type="checkbox"/> 4 Year <input type="checkbox"/> 5 Year <input type="checkbox"/> Not related to a study visit <input type="checkbox"/> Unscheduled visit |
| Deviation Category <i>(Select one)</i> | <input type="checkbox"/> Eligibility <input type="checkbox"/> Adverse event not reported per protocol <input type="checkbox"/> Informed Consent <input type="checkbox"/> Randomization <input type="checkbox"/> Study medications <input type="checkbox"/> Procedure/assessment complete out of window <input type="checkbox"/> Procedure/assessment done but not per protocol <input type="checkbox"/> Procedure/assessment incomplete or not done <input type="checkbox"/> Visit not done <input type="checkbox"/> Visit out of window <input type="checkbox"/> Other, specify: _____ | |
| If procedure/assessment <i>(Check all that apply)</i> | <input type="checkbox"/> Study Index Procedure <input type="checkbox"/> Physical Exam <input type="checkbox"/> Angiography <input type="checkbox"/> Echocardiography/CT <input type="checkbox"/> ECG <input type="checkbox"/> Laboratory Assessment <input type="checkbox"/> NIHSS <input type="checkbox"/> mRS <input type="checkbox"/> QVSFS <input type="checkbox"/> Other, specify: _____ | |

| | | |
|--|---|----------------------------------|
| Deviation Reason | <input type="checkbox"/> Oversight in protocol requirements <input type="checkbox"/> Subject refusal or non-compliance <input type="checkbox"/> Unable to reach subject <input type="checkbox"/> Site scheduling difficulty/error <input type="checkbox"/> Investigator decision to protect the rights, safety and welfare of subject <input type="checkbox"/> Equipment failure <input type="checkbox"/> User error <input type="checkbox"/> COVID-19 – Subject diagnosed <input type="checkbox"/> COVID-19 – Other, specify: _____ <input type="checkbox"/> Disaster/Weather related <input type="checkbox"/> Other, specify: _____ | |
| Additional Description of Deviation | | |
| Action Taken | <input type="checkbox"/> None <input type="checkbox"/> Documented site retraining <input type="checkbox"/> Subject education/review of study requirements with subject <input type="checkbox"/> Other, specify: _____ | |
| Does this Protocol Deviation (PD) require prompt reporting to the IRB? | <input type="checkbox"/> Yes <input type="checkbox"/> No | |
| | If yes, submitted on: | ____ / ____ / ____ (DD/MMM/YYYY) |

Site Personnel Signature

____ / ____ / ____
Date (DD/MMM/YYYY)

Note: Please complete only one deviation per form.

PD # in EDC _____

| | | |
|--|---|---|
| Date of Deviation | ___ / ___ / ___ (DD/MMM/YYYY) | |
| Date of Site Awareness | ___ / ___ / ___ (DD/MMM/YYYY) | |
| Time Period of Deviation | <input type="checkbox"/> Screening <input type="checkbox"/> Index Procedure <input type="checkbox"/> Discharge <input type="checkbox"/> Day 7 <input type="checkbox"/> Day 45 <input type="checkbox"/> 6 Months <input type="checkbox"/> 12 Months <input type="checkbox"/> 18 Months | <input type="checkbox"/> 2 Year <input type="checkbox"/> 3 Year <input type="checkbox"/> 4 Year <input type="checkbox"/> 5 Year <input type="checkbox"/> Not related to a study visit <input type="checkbox"/> Unscheduled visit |
| Deviation Category <i>(Select one)</i> | <input type="checkbox"/> Eligibility <input type="checkbox"/> Adverse event not reported per protocol <input type="checkbox"/> Informed Consent <input type="checkbox"/> Randomization <input type="checkbox"/> Study medications <input type="checkbox"/> Procedure/assessment complete out of window <input type="checkbox"/> Procedure/assessment done but not per protocol <input type="checkbox"/> Procedure/assessment incomplete or not done <input type="checkbox"/> Visit not done <input type="checkbox"/> Visit out of window <input type="checkbox"/> Other, specify: _____ | |
| If procedure/assessment <i>(Check all that apply)</i> | <input type="checkbox"/> Study Index Procedure <input type="checkbox"/> Physical Exam <input type="checkbox"/> Angiography <input type="checkbox"/> Echocardiography/CT <input type="checkbox"/> ECG <input type="checkbox"/> Laboratory Assessment <input type="checkbox"/> NIHSS <input type="checkbox"/> mRS <input type="checkbox"/> QVSFS <input type="checkbox"/> Other, specify: _____ | |

| | |
|--|---|
| Deviation Reason | <input type="checkbox"/> Oversight in protocol requirements <input type="checkbox"/> Subject refusal or non-compliance <input type="checkbox"/> Unable to reach subject <input type="checkbox"/> Site scheduling difficulty/error <input type="checkbox"/> Investigator decision to protect the rights, safety and welfare of subject <input type="checkbox"/> Equipment failure <input type="checkbox"/> User error <input type="checkbox"/> COVID-19 – Subject diagnosed <input type="checkbox"/> COVID-19 – Other, specify: _____ <input type="checkbox"/> Disaster/Weather related <input type="checkbox"/> Other, specify: _____ |
| Additional Description of Deviation | |
| Action Taken | <input type="checkbox"/> None <input type="checkbox"/> Documented site retraining <input type="checkbox"/> Subject education/review of study requirements with subject <input type="checkbox"/> Other, specify: _____ |
| Does this Protocol Deviation (PD) require prompt reporting to the IRB? | <input type="checkbox"/> Yes <input type="checkbox"/> No |
| | If yes, submitted on: ___ / ___ / ___ (DD/MMM/YYYY) |

Site Personnel Signature

Date (DD/MMM/YYYY)

Note: Please complete only one deviation per form.

PD # in EDC _____

| | | |
|--|---|---|
| Date of Deviation | ___ / ___ / ___ (DD/MMM/YYYY) | |
| Date of Site Awareness | ___ / ___ / ___ (DD/MMM/YYYY) | |
| Time Period of Deviation | <input type="checkbox"/> Screening <input type="checkbox"/> Index Procedure <input type="checkbox"/> Discharge <input type="checkbox"/> Day 7 <input type="checkbox"/> Day 45 <input type="checkbox"/> 6 Months <input type="checkbox"/> 12 Months <input type="checkbox"/> 18 Months | <input type="checkbox"/> 2 Year <input type="checkbox"/> 3 Year <input type="checkbox"/> 4 Year <input type="checkbox"/> 5 Year <input type="checkbox"/> Not related to a study visit <input type="checkbox"/> Unscheduled visit |
| Deviation Category <i>(Select one)</i> | <input type="checkbox"/> Eligibility <input type="checkbox"/> Adverse event not reported per protocol <input type="checkbox"/> Informed Consent <input type="checkbox"/> Randomization <input type="checkbox"/> Study medications <input type="checkbox"/> Procedure/assessment complete out of window <input type="checkbox"/> Procedure/assessment done but not per protocol <input type="checkbox"/> Procedure/assessment incomplete or not done <input type="checkbox"/> Visit not done <input type="checkbox"/> Visit out of window <input type="checkbox"/> Other, specify: _____ | |
| If procedure/assessment <i>(Check all that apply)</i> | <input type="checkbox"/> Study Index Procedure <input type="checkbox"/> Physical Exam <input type="checkbox"/> Angiography <input type="checkbox"/> Echocardiography/CT <input type="checkbox"/> ECG <input type="checkbox"/> Laboratory Assessment <input type="checkbox"/> NIHSS <input type="checkbox"/> mRS <input type="checkbox"/> QVSFS <input type="checkbox"/> Other, specify: _____ | |

| | |
|--|---|
| Deviation Reason | <input type="checkbox"/> Oversight in protocol requirements <input type="checkbox"/> Subject refusal or non-compliance <input type="checkbox"/> Unable to reach subject <input type="checkbox"/> Site scheduling difficulty/error <input type="checkbox"/> Investigator decision to protect the rights, safety and welfare of subject <input type="checkbox"/> Equipment failure <input type="checkbox"/> User error <input type="checkbox"/> COVID-19 – Subject diagnosed <input type="checkbox"/> COVID-19 – Other, specify: _____ <input type="checkbox"/> Disaster/Weather related <input type="checkbox"/> Other, specify: _____ |
| Additional Description of Deviation | |
| Action Taken | <input type="checkbox"/> None <input type="checkbox"/> Documented site retraining <input type="checkbox"/> Subject education/review of study requirements with subject <input type="checkbox"/> Other, specify: _____ |
| Does this Protocol Deviation (PD) require prompt reporting to the IRB? | <input type="checkbox"/> Yes <input type="checkbox"/> No |
| | If yes, submitted on: ___ / ___ / ___ (DD/MMM/YYYY) |

Site Personnel Signature

Date (DD/MMM/YYYY)

SCREENING

Informed Consent

| | |
|--|---|
| Subject to be enrolled as | <input type="checkbox"/> Roll-In <input type="checkbox"/> Randomized |
| Protocol Version Activated to at time of Informed Consent: | |
| Site ICF Version /IRB Approval Date DDMMYYYY | |
| Was this subject screened previously? | <input type="checkbox"/> Yes <input type="checkbox"/> No |
| | Previous Subject ID: _____ - _____ |

Randomization: N/A

Randomization shall be within 90 days of informed consent. The LAA occlusion procedure shall take place within and including 14 days from the date of randomization.

Randomization takes place in MEDIDATA Conform Study Data Base. Reference MOPs Binder, as needed

Print off Randomization eCRF and place in Subject Binder.

Screening Demographics

| | |
|--|---|
| If female, is subject of childbearing age? | <input type="checkbox"/> Yes <input type="checkbox"/> No |
|--|---|

Pregnancy Test

| | |
|----------------------------------|---|
| If yes, was pregnancy test done? | <input type="checkbox"/> Yes <input type="checkbox"/> No <i>If no and the female is of child-bearing age, complete a protocol deviation</i> <input type="checkbox"/> N/A Reason N/A: |
| Date of pregnancy test | ___ / ___ / ___ (DD/MMM/YYYY) |
| Result | <input type="checkbox"/> Positive <i>(Check I&E Criteria!)</i> <input type="checkbox"/> Negative |

Documentation of Shared Decision Making

Source must be present in Subject Record to document that INCLUSION 6 has been met.
Deemed appropriate for LAA closure by the site investigator and a clinician not a part of the procedural team using a shared decision-making process in accordance with standard of care

Confirmation that shared decision-making already documented in other medical records

Site Personnel Signature

___/___/___
Date (DD/MMM/YYYY)

| | |
|---|---|
| <p>Have all the inclusion criteria and none of the exclusion criteria, as specified by the protocol, been met for this subject?</p> | <p><input type="checkbox"/> Yes <input type="checkbox"/> No <input type="checkbox"/> N/A – Inclusion and Exclusion Criteria not assessed</p> |
| <p>What primary imaging modality was used to assess Echo Exclusion Criteria?</p> | <p><input type="checkbox"/> TTE – Transthoracic echocardiogram <input type="checkbox"/> CT <input type="checkbox"/> MRI <input type="checkbox"/> TEE – Transesophageal echocardiogram <input type="checkbox"/> None</p> |

Inclusion Criteria

Potential subjects must meet **ALL** of the following criteria to be eligible for inclusion in the study:

| Inclusion Criteria | Yes | No | N/A – Not assessed |
|--|--------------------------|--------------------------|--------------------------|
| 1. Male or nonpregnant female aged ≥ 18 years? | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> |
| 2. Documented non-valvular AF (paroxysmal, persistent, or permanent)? | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> |
| 3. High risk of stroke or systemic embolism, defined as CHA ₂ DS ₂ -VASc score of ≥ 3 ? | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> |
| 4. Has an appropriate rationale to seek a non-pharmacologic alternative to long-term oral anticoagulation? | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> |
| 5. Deemed by the site investigator to be suitable for short term oral anticoagulation therapy but deemed less favorable for long-term oral anticoagulation? | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> |
| 6. Deemed appropriate for LAA closure by the site investigator and a clinician not a part of the procedural team using a shared decision-making process in accordance with standard of care? | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> |
| 7. Able to comply with the protocol-specified medication regimen and follow-up evaluations? | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> |
| 8. The subject (or legally authorized representative, where allowed) has been informed of the nature of the study, agrees to its provisions, and has provided written informed consent approved by the appropriate Institutional Review Board (IRB)/Regional Ethics Board (REB)? | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> |

Exclusion Criteria

*Potential subjects will be excluded if **ANY** of the following conditions apply*

| Exclusion Criteria | Yes | No | N/A – Not assessed |
|--|--------------------------|--------------------------|--------------------------|
| 1. Pregnant or nursing subjects and those who plan pregnancy in the period up to 1 year following index procedure? Female subjects of childbearing potential must have a negative pregnant test (per site standard test) within 7 days prior to index procedure | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> |
| 2. Anatomic conditions that would prevent performance of an LAA occlusion procedure (e.g., prior atrial septal defect [ASD] or high-risk patent foramen ovale [PFO], surgical repair or implanted closure device, or obliterated or ligated left atrial appendage)? | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> |
| 3. Atrial fibrillation that is defined by a single occurrence or that is transient or reversible (e.g., secondary thyroid disorders, acute alcohol intoxication, trauma, recent major surgical procedures)? | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> |
| 4. A medical condition (other than atrial fibrillation) that mandates long-term oral anticoagulation (e.g., history of unprovoked deep vein thrombosis or pulmonary embolism, or mechanical heart valve)? | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> |
| 5. History of bleeding diathesis or coagulopathy, or subjects in whom antiplatelet and/or anticoagulant therapy is contraindicated? | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> |
| 6. Documented active infection? | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> |
| 7. Symptomatic carotid artery disease (defined as >50% stenosis with symptoms of ipsilateral transient or visual TIA evidence by amaurosis fugax, ipsilateral hemispheric TIAs or ipsilateral stroke)? If subject has a history of carotid stent or endarterectomy, the subject is eligible if there is <50% stenosis at the site of prior treatment | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> |
| 8. Recent (within 30 days of index procedure) or planned (within 60 days post-procedure) cardiac or non-cardiac interventional or surgical procedure? | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> |
| 9. Recent (within 30 days of index procedure) stroke or transient ischemic attack? | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> |
| 10. Recent (within 30 days of index procedure) myocardial infarction? | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> |
| 11. Vascular access precluding delivery of implant with catheter-based system? | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> |
| 12. Severe heart failure (New York Heart Association Class IV)? | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> |
| 13. Prior cardiac transplant, history of mitral valve replacement or transcatheter mitral valve intervention, or any mechanical valve implant? | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> |
| 14. Renal insufficiency, defined as estimated glomerular filtration rate (eGFR) <30 mL/min/1.73m ² (by the Modification of Diet in Renal Disease equation)? | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> |

| Exclusion Criteria | Yes | No | N/A – Not assessed |
|--|--------------------------|--------------------------|--------------------------|
| 15. Platelet count < 75,000 cells/mm ³ or > 700,000 cells/ mm ³ , or white blood cell count < 3,000 cells/ mm ³ ? | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> |
| 16. Known allergy, hypersensitivity or contraindication to aspirin, heparin, or device materials (e.g., nickel, titanium) or that would preclude any P2Y12 inhibitor therapy, or the subject has contrast sensitivity that cannot be adequately pre-medicated? | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> |
| 17. Actively enrolled or plans to enroll in a concurrent clinical study in which the active treatment arm may confound the results of this trial? | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> |
| 18. Unable to undergo general anesthesia? | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> |
| 19. Known other medical illness or known history of substance abuse that may cause non-compliance with the protocol or protocol-specified medication regimen, confound the data interpretation, or is associated with a life expectancy of less than 5 years? | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> |
| 20. A condition which precludes adequate transesophageal echocardiographic (TEE) assessment? | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> |

Screening Echocardiographic Exclusion Criteria

*This is based on historical imaging (performed within 6 months prior to consent) at Screening. Cardiac CT or TEE can be used to assess all criteria TTE and MRI studies are limited to the confirmed assessment of #3 and #4. Potential subjects will be excluded if **ANY** of the following conditions are known to apply*

| Exclusion Criteria | Yes | No | N/A – Not assessed |
|---|--------------------------|--------------------------|--------------------------|
| 1. Left atrial appendage cannot accommodate either a commercially available device of the CLAAS device per manufacturer IFU (e.g., the anatomy and sizing must be appropriate for both devices in order to be enrolled in the trial)? | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> |
| 2. Intracardiac thrombus or dense spontaneous echo contrast consistent with thrombus, as visualized by TEE prior to implant? | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> |
| 3. Left ventricular ejection fraction (LVEF) < 30%? | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> |
| 4. Existing circumferential pericardial effusion > 10 mm or symptomatic pericardial effusion, signs, or symptoms of acute or chronic pericarditis, or evidence of tamponade physiology? | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> |
| 5. Atrial septal defect that warrants closure? | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> |
| 6. High risk patent foramen ovale (PFO), defined as an atrial septal aneurysm (exclusion > 15 mm or length > 15 mm) or large shunt (early [within 3 beats] and/or substantial passage of bubbles, e.g., > 20)? | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> |
| 7. Moderate or severe mitral valve stenosis (mitral valve area < 1.5 cm ²)? | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> |
| 8. Complex atheroma with mobile plaque of the descending aorta and/or aortic arch? | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> |
| 9. Evidence of cardiac tumor? | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> |

Reminder: If a significant cardiac event (potentially related to a change in cardiac status, e.g., CHF decompensation) occurs after cardiac imaging is obtained and before randomization takes place- then imaging should be repeated.

Site Personnel Signature

___/___/_____
Date (DD/MMM/YYYY)

Investigator Signature

___/___/_____
Date (DD/MMM/YYYY)

Site Number: _____ Subject ID: _____

Documentation of Shared Decision Making

N/A, shared decision-making already documented in other medical records

Implanting Physician Name (First Last)

Implanting Physician Specialty

Interventional Cardiology
 Electrophysiology

Referring Physician (First Last)

Referring Physician Specialty

Attestation:

Based on my review of the patient's medical history, and in conjunction with a formal and shared decision-making process involving the patient and multidisciplinary team, the patient is suitable for the following:

LAA Closure
 Short Term Oral Anticoagulation

Source must be present in Subject Record, or Subject Study Binder to document that INCLUSION 6 has been met. If utilizing this source, i.e., no source in other MR is available, this Attestation Source should be signed by Subject's Implanting Study Investigator or the Principal Investigator

Site Personnel Signature

___/___/_____
Date (DD/MMM/YYYY)

| | | | |
|---|--|--|--|
| Date Medical History Performed | ___ / ___ / ___ (DD/MMM/YYYY) | | |
| Rationale for seeking a non-pharmacologic alternative to OAC (Check all that apply) | <input type="checkbox"/> Drug regimen not compatible with OAC <input type="checkbox"/> Non-compliance to medication or monitoring schedule <input type="checkbox"/> History of bleeding or high bleeding risk <input type="checkbox"/> Renal failure <input type="checkbox"/> High fall risk <input type="checkbox"/> Other, specify: _____ | | |
| Documented type of non-valvular atrial fibrillation: | <input type="checkbox"/> Paroxysmal <input type="checkbox"/> Persistent <input type="checkbox"/> Permanent | | |
| Does the subject have a medical condition that mandates long term oral anticoagulation? | <input type="checkbox"/> Yes (<i>Review for I&E!</i>) <input type="checkbox"/> No | | |
| Diabetes mellitus (DM)? | <input type="checkbox"/> Yes <input type="checkbox"/> No <input type="checkbox"/> Unknown If yes, please select one: | <input type="checkbox"/> Insulin dependent diabetes mellitus (IDDM) <input type="checkbox"/> Type I DM <input type="checkbox"/> Type II DM <input type="checkbox"/> Unknown <input type="checkbox"/> Non-insulin Dependent Diabetes Mellitus How is NIDDM controlled? <input type="checkbox"/> Diet <input type="checkbox"/> Oral Hypoglycemics <input type="checkbox"/> Unknown <input type="checkbox"/> Unknown | |
| History of hypertension (Systolic BP > 140 mmHg, or Diastolic BP >90 mmHg)? | <input type="checkbox"/> Yes <input type="checkbox"/> No <input type="checkbox"/> Unknown | If yes, currently requires medication? | <input type="checkbox"/> Yes <input type="checkbox"/> No <input type="checkbox"/> Unknown |
| History of hyperlipidemia (medical diagnosis) or total cholesterol >200? | <input type="checkbox"/> Yes <input type="checkbox"/> No <input type="checkbox"/> Unknown | If yes, currently requires medication? | <input type="checkbox"/> Yes <input type="checkbox"/> No <input type="checkbox"/> Unknown |
| History of peripheral vascular disease? | <input type="checkbox"/> Yes <input type="checkbox"/> No <input type="checkbox"/> Unknown | If yes, prior intervention? | <input type="checkbox"/> Yes (check all that apply) <input type="checkbox"/> Percutaneous <input type="checkbox"/> Surgical <input type="checkbox"/> Unknown <input type="checkbox"/> No <input type="checkbox"/> Unknown |

| | | | | |
|--|---|--|--|---|
| History of carotid artery disease? | <input type="checkbox"/> Yes <input type="checkbox"/> No <input type="checkbox"/> Unknown | If yes, location | <input type="checkbox"/> Right <input type="checkbox"/> Left <input type="checkbox"/> Bilateral | |
| | | If yes, prior intervention? | <input type="checkbox"/> Yes, specify: <input type="checkbox"/> Endarterectomy <input type="checkbox"/> Stent <input type="checkbox"/> No <input type="checkbox"/> Unknown | |
| Prior cerebral vascular accident? | <input type="checkbox"/> Yes <input type="checkbox"/> No <input type="checkbox"/> Unknown | If yes, date of most recent CVA: | ___ / ___ / ___ (DD/MMM/YYYY) | |
| | | If yes, is imaging available? | <input type="checkbox"/> Yes Date of most recent Brain Scan MRI or CT Imaging: ___ / ___ / ___ (DD/MMM/YYYY) <input type="checkbox"/> No | |
| | | If yes, specify type (Check all that apply) | <input type="checkbox"/> Ischemic <input type="checkbox"/> Hemorrhagic <input type="checkbox"/> Unknown | |
| Prior traumatic intracranial hemorrhage? | <input type="checkbox"/> Yes <input type="checkbox"/> No <input type="checkbox"/> Unknown | If yes, date of most recent intracranial hemorrhage: | ___ / ___ / ___ (DD/MMM/YYYY) | |
| | | If yes, is imaging available? | <input type="checkbox"/> Yes Date of most recent imaging: ___ / ___ / ___ (DD/MMM/YYYY) <input type="checkbox"/> No | |
| | | If yes, specify type (Check all that apply) | <input type="checkbox"/> Spontaneous <input type="checkbox"/> Traumatic | |
| Prior transient ischemic attack? | <input type="checkbox"/> Yes <input type="checkbox"/> No <input type="checkbox"/> Unknown | If yes, date of most recent TIA: | ___ / ___ / ___ (DD/MMM/YYYY) | |
| History of coronary artery disease? | <input type="checkbox"/> Yes <input type="checkbox"/> No <input type="checkbox"/> Unknown | If yes, current anginal status | <input type="checkbox"/> Asymptomatic <input type="checkbox"/> Stable Angina <input type="checkbox"/> Unstable Angina | |
| | | If yes, prior coronary artery intervention? | <input type="checkbox"/> Yes <input type="checkbox"/> No <input type="checkbox"/> Unknown | If yes, specify type: <input type="checkbox"/> Percutaneous <input type="checkbox"/> Surgical |

| | | | |
|---|---|--|---|
| History of congestive heart failure? | <input type="checkbox"/> Yes <input type="checkbox"/> No <input type="checkbox"/> Unknown | If yes, NYHA Functional Class | <input type="checkbox"/> Class I <input type="checkbox"/> Class II <input type="checkbox"/> Class III <input type="checkbox"/> Class IV (<i>Review for I&E!</i>) |
| What is the most recently documented LVEF (%)? (xx) | _____ % | ___ / ___ / _____ (DD/MMM/YYYY) | |
| History of intracardiac mass, thrombus or vegetation? | <input type="checkbox"/> Yes <input type="checkbox"/> No <input type="checkbox"/> Unknown | If yes, specify location | <input type="checkbox"/> Left Ventricle <input type="checkbox"/> Left Atrium <input type="checkbox"/> Left Atrial Appendage <input type="checkbox"/> Other, specify: _____ |
| History of severe valvular heart disease? | <input type="checkbox"/> Yes <input type="checkbox"/> No <input type="checkbox"/> Unknown | If yes, specify type (<i>Check all that apply</i>) | <input type="checkbox"/> Aortic valve stenosis <input type="checkbox"/> Aortic valve regurgitation <input type="checkbox"/> Mitral valve stenosis <input type="checkbox"/> Mitral valve regurgitation <input type="checkbox"/> Tricuspid valve stenosis <input type="checkbox"/> Tricuspid valve regurgitation <input type="checkbox"/> Unknown |
| Does the subject have history of prior cardiac transplant, history of mitral valve replacement or transcatheter mitral valve intervention, or any mechanical valve implant? | <input type="checkbox"/> Yes (<i>Review for I&E!</i>) <input type="checkbox"/> No | | |
| History of procedure to convert atrial fibrillation to atrial flutter? | <input type="checkbox"/> Yes <input type="checkbox"/> No <input type="checkbox"/> Unknown | If yes, specify type | <input type="checkbox"/> Cardioversion <input type="checkbox"/> Ablation |
| History of acute or chronic pericarditis? | <input type="checkbox"/> Yes <input type="checkbox"/> No <input type="checkbox"/> Unknown | | |
| Has the subject had a cardiac or non-cardiac intervention or surgical procedure within 30 days of the index procedure? | <input type="checkbox"/> Yes (<i>Review for I&E!</i>) <input type="checkbox"/> No | | |
| Does the subject have a planned surgical procedure within 60 days AFTER the date of the planned Index Procedure Date? | <input type="checkbox"/> Yes (<i>Review for I&E!</i>) <input type="checkbox"/> No | | |

| | | | |
|---|--|---------------------------|---|
| History of myocardial infarction? | <input type="checkbox"/> Yes <input type="checkbox"/> No <input type="checkbox"/> Unknown | If yes, most recent date: | ____/____/____ (DD/MMM/YYYY) |
| History of cardiomyopathy? | <input type="checkbox"/> Yes <input type="checkbox"/> No <input type="checkbox"/> Unknown | | |
| History of patent foramen ovale (PFO) | <input type="checkbox"/> Yes <input type="checkbox"/> No <input type="checkbox"/> Unknown | If yes, treated? | <input type="checkbox"/> Yes <input type="checkbox"/> No <input type="checkbox"/> Unknown |
| History of atrial septal defect (ASD)? | <input type="checkbox"/> Yes <input type="checkbox"/> No <input type="checkbox"/> Unknown | If yes, treated? | <input type="checkbox"/> Yes <input type="checkbox"/> No <input type="checkbox"/> Unknown |
| History of gastrointestinal bleeding? | <input type="checkbox"/> Yes <input type="checkbox"/> No <input type="checkbox"/> Unknown | | |
| History of other form of recurrent systemic bleeding? | <input type="checkbox"/> Yes <input type="checkbox"/> No <input type="checkbox"/> Unknown | | |
| History of anemia requiring transfusion? | <input type="checkbox"/> Yes <input type="checkbox"/> No <input type="checkbox"/> Unknown | | |
| History of renal disease? | <input type="checkbox"/> Yes <input type="checkbox"/> No <input type="checkbox"/> Unknown | | |
| History of malignancy? | <input type="checkbox"/> Yes <input type="checkbox"/> No <input type="checkbox"/> Unknown | | |
| History of dementia? | <input type="checkbox"/> Yes <input type="checkbox"/> No <input type="checkbox"/> Unknown | | |
| Does subject have history of COVID-19? | <input type="checkbox"/> Yes <input type="checkbox"/> No <input type="checkbox"/> Unknown <input type="checkbox"/> Patient declined to answer | | |
| Has subject received COVID-19 vaccination? | <input type="checkbox"/> Yes <input type="checkbox"/> No <input type="checkbox"/> Unknown <input type="checkbox"/> Patient declined to answer | | |

Site Personnel Signature

____/____/____
Date (DD/MMM/YYYY)

| | |
|-----------------------------------|---|
| Were vital signs performed? | <input type="checkbox"/> Yes <input type="checkbox"/> No |
| Date of vital sign measurements | ___ / ___ / ___ (DD/MMM/YYYY) |
| Height (xxx.xx) | _____ (cm / in) (circle one) |
| Weight (xxx.x) | _____ (kg / lb) (circle one) |
| BMI (xx.x) | _____ (kg/m ²) |
| Systolic Blood Pressure (xxx) | _____ (mmHg) |
| Diastolic Blood Pressure (xxx) | _____ (mmHg) |
| Heart Rate (xxx) | _____ (bpm) |

Site Personnel Signature

___ / ___ / ___
Date (DD/MMM/YYYY)

Should be performed as per Standard of Care

| | |
|--|---|
| Was Physical Examination - Review of Systems performed? | <input type="checkbox"/> Yes <input type="checkbox"/> No |
| Date of examination | ___/___/___ (DD/MMM/YYYY) |
| If subject has suspected incident of neurologic event based off responses to QVSFS, NIHSS or other signs/symptoms, was neurologic exam performed by neurologist/clinical designee? | <input type="checkbox"/> N/A – Patient doesn't have suspected incident of neurologic event <input type="checkbox"/> Yes <input type="checkbox"/> No |
| Date of neurologic examination | ___/___/___ (DD/MMM/YYYY) |

| Body System Examined | Normal | Abnormal (CS) | Abnormal (NCS) | Not Done | Description of abnormal findings |
|---|--------------------------|--------------------------|--------------------------|--------------------------|----------------------------------|
| <input type="checkbox"/> General Appearance | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> | |
| <input type="checkbox"/> Cardiovascular | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> | |
| <input type="checkbox"/> Dermatological | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> | |
| <input type="checkbox"/> Ears/Nose/Throat | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> | |
| <input type="checkbox"/> Gastrointestinal | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> | |
| <input type="checkbox"/> Genito-urinary | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> | |
| <input type="checkbox"/> Musculoskeletal | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> | |
| <input type="checkbox"/> Neurological | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> | |
| <input type="checkbox"/> Respiratory | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> | |

CS = Clinically significant
NCS = Not clinically significant

Site Personnel Signature

___/___/___
Date (DD/MMM/YYYY)

Note: It is not required to complete this source worksheet if the information below is clearly documented in other records.

| | |
|--|---|
| Was CHA ₂ DS ₂ VASc completed? | <input type="checkbox"/> Yes <input type="checkbox"/> No (<i>Complete Protocol Deviation Form</i>) |
| Date completed | ___ / ___ / ___ (DD/MMM/YYYY) |
| If CHA ₂ DS ₂ VASc is selected (<i>Select all that apply</i>) | <input type="checkbox"/> Age (years): <input type="checkbox"/> <65 <input type="checkbox"/> 65-74 <input type="checkbox"/> ≥75 <input type="checkbox"/> Female sex <input type="checkbox"/> Congestive Heart Failure history <input type="checkbox"/> Hypertension history <input type="checkbox"/> Stroke or TIA symptoms previously <input type="checkbox"/> Vascular disease history <input type="checkbox"/> Diabetes mellitus history |
| Score | <i>Auto-calculated in EDC</i> |

Site Personnel Signature

___ / ___ / ___
Date (DD/MMM/YYYY)

Note: It is not required to complete this source worksheet if the information below is clearly documented in other records.

| | |
|--|--|
| Was the HAS-BLED Score completed? (Select only one) | <input type="checkbox"/> Yes <input type="checkbox"/> No <i>If no, complete a protocol deviation</i> |
| Date completed | ___ / ___ / ___ (DD/MMM/YYYY) |
| HAS-BLED Score (Check all that apply) | <input type="checkbox"/> None of the below <input type="checkbox"/> Hypertension (<i>Uncontrolled, >160 mmHg systolic</i>) <input type="checkbox"/> Renal disease (<i>Dialysis, transplant, CR >2.26 mg/dL or >200 µmol/L</i>) <input type="checkbox"/> Liver disease (<i>Cirrhosis or bilirubin >2x normal with AST/ALT/AP >3x normal</i>) <input type="checkbox"/> Stroke history <input type="checkbox"/> Prior major bleed or predisposition to bleeding <input type="checkbox"/> Labile INR (<i>Unstable/high INRs, time in therapeutic range <60%</i>) <input type="checkbox"/> Age > 65 years <input type="checkbox"/> On medications that predispose to bleeding (<i>aspirin, clopidogrel, NSAIDs</i>) <input type="checkbox"/> Alcohol use (<i>≥8 drinks/week</i>) |
| Score | <i>Auto-Calculated in EDC</i> |

Site Personnel Signature___ / ___ / ___
Date (DD/MMM/YYYY)

| | | | |
|--|---|--|---|
| Was ECG performed? | <input type="checkbox"/> Yes <input type="checkbox"/> No (<i>Complete a protocol deviation form</i>) | | |
| Date of ECG | ___ / ___ / ___ (DD/MMM/YYYY) | | |
| Sinus rhythm | <input type="checkbox"/> Yes <input type="checkbox"/> No | | |
| Atrial Arrhythmia | <input type="checkbox"/> Yes <input type="checkbox"/> No | Atrial fibrillation | <input type="checkbox"/> Yes <input type="checkbox"/> No |
| | | Atrial flutter | <input type="checkbox"/> Yes <input type="checkbox"/> No |
| | | Paroxysmal atrial fibrillation/flutter | <input type="checkbox"/> Yes <input type="checkbox"/> No |
| | | Atrial tachycardia | <input type="checkbox"/> Yes <input type="checkbox"/> No |
| Junctional rhythm | <input type="checkbox"/> Yes <input type="checkbox"/> No | | |
| AV node conduction disturbance/heart block | <input type="checkbox"/> Yes <input type="checkbox"/> No | If yes, what degree? | <input type="checkbox"/> 1 st Degree <input type="checkbox"/> 2 nd Degree <input type="checkbox"/> 3 rd Degree |
| Paced Rhythm | <input type="checkbox"/> Yes <input type="checkbox"/> No | | |
| Q-Wave present | <input type="checkbox"/> Yes <input type="checkbox"/> No | | |
| Left bundle branch block present | <input type="checkbox"/> Yes <input type="checkbox"/> No | | |
| Right bundle branch block present | <input type="checkbox"/> Yes <input type="checkbox"/> No | | |

Site Personnel Signature

___ / ___ / ___
Date (DD/MMM/YYYY)

This Worksheet is to be used at the Screening Visit.

| | |
|---|--|
| Was Echocardiogram/CT performed? | <input type="checkbox"/> Yes <input type="checkbox"/> No <i>(Complete protocol deviation form)</i> |
| Are the required images for this visit available? | <input type="checkbox"/> Yes <input type="checkbox"/> No <i>(Complete protocol deviation form)</i> |
| Were images uploaded into the Imaging Module? | <input type="checkbox"/> Yes <input type="checkbox"/> No |
| Date echocardiogram/CT completed | ___ / ___ / ___ (DD/MMM/YYYY) |
| Imaging Type | <input type="checkbox"/> TTE – Transthoracic echocardiogram <input type="checkbox"/> TEE – Transesophageal echocardiogram <input type="checkbox"/> Cardiac CT <input type="checkbox"/> Cardiac MRI <input type="checkbox"/> Brain CT <input type="checkbox"/> MRI |
| Left atrial appendage visible | <input type="checkbox"/> Yes <input type="checkbox"/> No <input type="checkbox"/> Not Available |

If available, confirm if the following was noted on echo/CT:

| | | | |
|---|--|--------------------------|---|
| Dense spontaneous echo contrast consistent with thrombus? | <input type="checkbox"/> Yes <i>(Review for I&E!)</i> <input type="checkbox"/> No <input type="checkbox"/> Not Available | | |
| Intra-cardiac thrombus | <input type="checkbox"/> Yes <i>(Review for I&E!)</i> <input type="checkbox"/> No <input type="checkbox"/> Not available | If yes, confirm location | <input type="checkbox"/> Left atrium <input type="checkbox"/> Left atrial appendage <input type="checkbox"/> Left ventricle <input type="checkbox"/> Right atrium <input type="checkbox"/> Right ventricle <input type="checkbox"/> Other, specify: _____ |
| Intra-cardiac vegetation | <input type="checkbox"/> Yes <input type="checkbox"/> No <input type="checkbox"/> Not available | If yes, confirm location | <input type="checkbox"/> Left atrium <input type="checkbox"/> Left atrial appendage <input type="checkbox"/> Left ventricle <input type="checkbox"/> Right atrium <input type="checkbox"/> Right ventricle <input type="checkbox"/> Other, specify: _____ |

| | | | |
|--|---|--|---|
| Patent foramen ovale warranting closure? | <input type="checkbox"/> Yes <input type="checkbox"/> No <input type="checkbox"/> Not available | If yes, is this a high risk PFO? | <input type="checkbox"/> Yes (<i>Review for I&E!</i>) <input type="checkbox"/> No |
| Atrial septal defect? | <input type="checkbox"/> Yes <input type="checkbox"/> No <input type="checkbox"/> Not available | If yes, specify | <input type="checkbox"/> Right to left shunt present <input type="checkbox"/> Left to right shunt present <input type="checkbox"/> Bidirectional shunt present <input type="checkbox"/> Unable to determine |
| | | If yes, does defect warrant closure? | <input type="checkbox"/> Yes (<i>Review for I&E!</i>) <input type="checkbox"/> No |
| Pericardial effusion present? | <input type="checkbox"/> Yes <input type="checkbox"/> No <input type="checkbox"/> Not available | If yes, select type | <input type="checkbox"/> Circumferential <input type="checkbox"/> Loculated |
| | | If yes, select size | <input type="checkbox"/> Trivial <input type="checkbox"/> Small (<1 cm) <input type="checkbox"/> Moderate (1-2 cm) <input type="checkbox"/> Large (>2 cm and <5cm) <input type="checkbox"/> Large (> 5 cm) (<i>Review for I&E!</i>) |
| | | If yes, Do any of the following apply? Check all that apply (<i>Review for I&E!</i>) | <input type="checkbox"/> Symptomatic <input type="checkbox"/> Sign or symptom of acute or chronic pericarditis <input type="checkbox"/> Evidence of tamponade physiology |

Site Personnel Signature

___/___/___
Date (DD/MMM/YYYY)

Original source should be obtained from a direct laboratory report from Subject Medical Record. Laboratory results to be reviewed by delegated investigator (either directly on lab report or as Source here) as relates to subject safety and INC/EXC Criteria.

*Laboratory Collection at screening collected per standard of care **up to 60 days prior to consent***

Date of Hematology __ __ / __ __ __ / __ __ __ __ (DD/MMM/YYYY)

Not Done (ENTER PD)

| Laboratory Assessment | Results | Clinically Significant |
|------------------------------|----------------|---|
| Hemoglobin | | <input type="checkbox"/> Yes <input type="checkbox"/> No |
| Henatocrit | | <input type="checkbox"/> Yes <input type="checkbox"/> No |
| WBC | | <input type="checkbox"/> Yes <input type="checkbox"/> No |
| Platelet Count | | <input type="checkbox"/> Yes <input type="checkbox"/> No |

CHEMISTRY – SERUM CREATININE OR GFR eGFR

Date of serum Chemistry __ __ / __ __ __ / __ __ __ __ (DD/MMM/YYYY)

Not Done (ENTER PD if neither Cr or GFR/eGFR were not obtained)

| Laboratory Assessment | Results | Clinically Significant |
|------------------------------|----------------|---|
| Creatinine | | <input type="checkbox"/> Yes <input type="checkbox"/> No |
| GFR or eGFR | | <input type="checkbox"/> Yes <input type="checkbox"/> No |

COAGULATION as Relevant

| | |
|---------------------------|--|
| Was INR sample collected? | <input type="checkbox"/> Yes <input type="checkbox"/> No <input type="checkbox"/> Not applicable |
| Date of INR | ___/___/___ (DD/MMM/YYYY) |

| <i>Laboratory Assessment</i> | <i>Results</i> | <i>Clinically Significant</i> |
|------------------------------|----------------|---|
| INR | | <input type="checkbox"/> Yes <input type="checkbox"/> No |

Site Personnel Signature

___/___/___
Date (DD/MMM/YYYY)

Reminder: Pre-procedure oral anticoagulation (Warfarin or DOAC) should be managed as per site protocol. Warfarin should be discontinued in accordance with site standard of care practices including the monitoring of INR levels on the day of the procedure.

Was the NIHSS assessment completed?

- Yes
 No (*Complete Protocol Deviation form*)

Date of NIHSS assessment

___ / ___ / ___ (DD/MM/YYYY)

| 1(a) – Level of consciousness | |
|---|------------------------------|
| Alert, keenly responsive | <input type="checkbox"/> (0) |
| Not alert; but arousable by minor stimulation to obey, answer or respond | <input type="checkbox"/> (1) |
| Not alert' requires repeated stimulation to attend, or is obtunded and requires strong or painful stimulation to make movements (not stereotyped) | <input type="checkbox"/> (2) |
| Responds only with reflex motor or autonomic unresponsive, flaccid or areflexic | <input type="checkbox"/> (3) |
| 1(b) – Level of consciousness questions | |
| Answers both questions correctly | <input type="checkbox"/> (0) |
| Answers one question correctly | <input type="checkbox"/> (1) |
| Answers neither question correctly | <input type="checkbox"/> (2) |
| 1(c) – Level of consciousness command | |
| Performs both tasks correctly | <input type="checkbox"/> (0) |
| Performs one task correctly | <input type="checkbox"/> (1) |
| Performs neither task correctly | <input type="checkbox"/> (2) |
| 2 – Best gaze | |
| Normal | <input type="checkbox"/> (0) |
| Partial gaze palsy; gaze is abnormal in one or both eyes, but forced deviation or total gaze paresis is not present | <input type="checkbox"/> (1) |
| Forced deviation, or total gaze paresis not overcome by the oculoccephalic maneuver | <input type="checkbox"/> (2) |
| 3 – Visual | |
| No visual loss | <input type="checkbox"/> (0) |
| Partial hemianopia | <input type="checkbox"/> (1) |
| Complete hemianopia | <input type="checkbox"/> (2) |
| Bilateral hemianopia (blind including cortical blindness) | <input type="checkbox"/> (3) |

| 4 – Facial palsy | |
|---|-------------------------------|
| Normal symmetrical movements | <input type="checkbox"/> (0) |
| Minor paralysis (flattened nasolabial fold, asymmetry on smiling) | <input type="checkbox"/> (1) |
| Partial paralysis (total or near-total paralysis of lower face) | <input type="checkbox"/> (2) |
| Complete paralysis of one or both sides (absence of facial movement in the upper and lower face) | <input type="checkbox"/> (3) |
| 5(a) – Motor arm - left | |
| No drift, limb holds 90 (or 45) degrees for 10 full seconds | <input type="checkbox"/> (0) |
| Drift; limb holds 90 (or 45) degrees, but drifts down before full 10 seconds; does not hit bed or other support | <input type="checkbox"/> (1) |
| Some effort against gravity; limb cannot get to or maintain (if cued) 90 (or 45) degrees; drifts down to bed, but has some effort against gravity | <input type="checkbox"/> (2) |
| No effort against gravity, limb falls | <input type="checkbox"/> (3) |
| No movement | <input type="checkbox"/> (4) |
| Amputation or joint fusion, Explain | <input type="checkbox"/> (UN) |
| 5(b) – Motor arm - right | |
| No drift, limb holds 90 (or 45) degrees for 10 full seconds | <input type="checkbox"/> (0) |
| Drift; limb holds 90 (or 45) degrees, but drifts down before full 10 seconds; does not hit bed or other support | <input type="checkbox"/> (1) |
| Some effort against gravity; limb cannot get to or maintain (if cued) 90 (or 45) degrees; drifts down to bed, but has some effort against gravity | <input type="checkbox"/> (2) |
| No effort against gravity, limb falls | <input type="checkbox"/> (3) |
| No movement | <input type="checkbox"/> (4) |
| Amputation or joint fusion, Explain: | <input type="checkbox"/> (UN) |

| 6(a) Motor leg - left | |
|--|-------------------------------|
| No drift, leg holds 30-degree position for full 5 seconds | <input type="checkbox"/> (0) |
| Drift: leg falls by the end of the 5-second period but does not hit bed | <input type="checkbox"/> (1) |
| Some effort against gravity; leg falls to bed by 5 seconds, but has some effort against gravity | <input type="checkbox"/> (2) |
| No effort against gravity, leg falls to bed immediately | <input type="checkbox"/> (3) |
| No movement | <input type="checkbox"/> (4) |
| Amputation or joint fusion, Explain: | <input type="checkbox"/> (UN) |
| 6(a) Motor leg - right | |
| No drift, leg holds 30-degree position for full 5 seconds | <input type="checkbox"/> (0) |
| Drift: leg falls by the end of the 5-second period but does not hit bed | <input type="checkbox"/> (1) |
| Some effort against gravity; leg falls to bed by 5 seconds, but has some effort against gravity | <input type="checkbox"/> (2) |
| No effort against gravity, leg falls to bed immediately | <input type="checkbox"/> (3) |
| No movement | <input type="checkbox"/> (4) |
| Amputation or joint fusion, Explain: | <input type="checkbox"/> (UN) |
| 7 – Limb ataxia | |
| Absent | <input type="checkbox"/> (0) |
| Present in one limb | <input type="checkbox"/> (1) |
| Present in two limbs | <input type="checkbox"/> (2) |
| Amputation or joint fusion, Explain: | <input type="checkbox"/> (UN) |
| 8 – Sensory | |
| Normal; no sensory loss | <input type="checkbox"/> (0) |
| Mild-to-moderate sensory loss; patient feels pinprick less sharp or dull on the affected side; or there is a loss of superficial pain with pinprick, but patient is aware of being touched | <input type="checkbox"/> (1) |
| Severe to total sensory loss; patient is not aware of being touched in the face, arm, and leg | <input type="checkbox"/> (2) |

| 9 – Best language | |
|---|-------------------------------|
| No aphasia; normal | <input type="checkbox"/> (0) |
| Mild-to-moderate aphasia; some obvious loss of fluency or facility or comprehension, without significant limitation on ideas expressed or form of expression. Reduction of speech and/or comprehension, however, makes conversion about provided materials difficult or impossible. <i>For example, in conversation about provided materials, examiner can identify picture or naming card content from patient's response.</i> | <input type="checkbox"/> (1) |
| Severe aphasia; all communication is through fragmentary expression; great need for inference questioning and guessing by the listener. Range of information that can be exchanged is limited; listener carries burden of communication. Examiner cannot identify materials provided from patient response. | <input type="checkbox"/> (2) |
| Mute, global aphasia; no usable speech or auditory comprehension | <input type="checkbox"/> (3) |
| 10 – Dysarthria | |
| Normal | <input type="checkbox"/> (0) |
| Mild-to-moderate dysarthria; patient slurs at least some word and, at worst, can be understood with some difficulty. | <input type="checkbox"/> (1) |
| Severe dysarthria; patient's speech is so slurred as to be unintelligible in the absence or out of proportion to any dysphasia, or is mute/anarthric | <input type="checkbox"/> (2) |
| Intubated or another physical barrier, Explain: | <input type="checkbox"/> (UN) |
| 11 – Extinction and inattention (formerly neglect) | |
| No abnormality | <input type="checkbox"/> (0) |
| Visual, tactile, auditory, spatial, or personal inattention or extinction to bilateral simultaneous stimulation in one of the sensory modalities. | <input type="checkbox"/> (1) |
| Profound hemi-inattention or extinction to more than one modality; does not recognize own hand or orients to only one side of space. | <input type="checkbox"/> (2) |

Site Personnel Signature

___/___/_____
Date (DD/MMM/YYYY)

CONFORM QVSFS

Source Data Transfer Tool

Site Number: _____ Subject ID: _____

| | |
|---|---|
| Was the QVSFS assessment completed? | <input type="checkbox"/> Yes <input type="checkbox"/> No (<i>Complete protocol deviation form</i>) |
| Is this assessment performed because of a neurological event? | <input type="checkbox"/> Yes <i>If yes, assess for an adverse event</i> <input type="checkbox"/> No |
| Date of QVSFS assessment | ___ / ___ / ___ (DD/MMM/YYYY) |

| Since the last study contact (by phone or clinic) | Yes | No | Unknown |
|--|--------------------------|--------------------------|--------------------------|
| 1. Were you told by a physician that you had a stroke? | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> |
| 2. Were you ever told by a physician that you had a TIA, ministroke, or a transient ischemic attack? | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> |
| 3. Have you ever had a sudden weakness on one side of your body? | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> |
| 4. Have you ever had a sudden numbness or dead feeling on one side of your body? | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> |
| 5. Have you ever had a sudden painless loss of vision in one or both eyes? | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> |
| 6. Have you ever suddenly lost one half of your vision? | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> |
| 7. Have you ever suddenly lost the ability to understand what people are saying? | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> |
| 8. Have you ever suddenly lost the ability to express yourself verbally or in writing? | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> |

Site Personnel Signature

___ / ___ / ___
Date (DD/MMM/YYYY)

Was the mRS assessment completed?

- Yes
 No (*Complete protocol deviation form*)

Is this assessment performed because of a neurological event?

- Yes *If yes, assess for an adverse event*
 No

Date of mRS assessment

___ / ___ / ___ (DD/MMM/YYYY)

| Level of Consciousness | Response |
|--|--------------------------|
| 0 = No symptoms at all | <input type="checkbox"/> |
| 1 = No significant disability despite symptoms; able to carry out all usual duties and activities | <input type="checkbox"/> |
| 2 = Slight disability; unable to carry out all previous activities but able to look after own affairs without assistance | <input type="checkbox"/> |
| 3 = Moderate disability; requiring some help, but able to walk without assistance | <input type="checkbox"/> |
| 4 = Moderately severe disability; unable to walk without assistance, and unable to attend to own body needs without assistance | <input type="checkbox"/> |
| 5 = Severe disability; bedridden, incontinent, and requiring constant nursing care and attention | <input type="checkbox"/> |
| Score | |

Site Personnel Signature___ / ___ / ___
Date (DD/MMM/YYYY)

PROCEDURE DAY PACKET



**Yale Cardiovascular Research Group
Yale Echocardiographic Core Laboratory**

**CONFORM Pivotal Trial
TEE/TTE Sonographer Worksheet**

Please submit completed TEE/TTE Sonographer Worksheet with image uploads

Site ID: _____

Subject ID: _____

Echo Study Date: ____/____/____
 dd mon yyyy

Modality: TEE TTE

Procedure Type:

- | | | |
|---|-----------------------------------|---|
| <input type="checkbox"/> Diagnostic/Screening | <input type="checkbox"/> 45-Day | <input type="checkbox"/> Unscheduled |
| <input type="checkbox"/> Index Procedure | <input type="checkbox"/> 6-Month | <input type="checkbox"/> Adverse Event |
| <input type="checkbox"/> Pre-Discharge | <input type="checkbox"/> 12-Month | <input type="checkbox"/> Optional TEE at Baseline |

Ultrasound Manufacturer:

Transducer Type:

Comments _____

Site personnel completing form:

Name (print)

Sign

____/____/____
dd mon yyyy

| | |
|--|---|
| Date of procedure | ____/____/____ (DD/MMM/YYYY) |
| Randomized to | <input type="checkbox"/> CLAAS <input type="checkbox"/> Control |
| Study Procedure | <input type="checkbox"/> CLAAS <input type="checkbox"/> Watchman <input type="checkbox"/> Amulet |
| Investigator (Operating Physician) First Name | |
| Investigator (Operating Physician) Last Name | |
| Primary Imager First Name | |
| Primary Imager Last Name | |
| What loading dose was prescribed to the patient prior to the procedure? | <input type="checkbox"/> 81-100 mg Aspirin <input type="checkbox"/> 325 mg Aspirin <input type="checkbox"/> No loading dose prescribed prior to index procedure <input type="checkbox"/> Other: _____ |
| Procedure start time (24 HR) <i>(Defined as time of first sheath insertion in primary venous access site)</i> | _____ : _____ |
| Access Sheath Insertion site <i>*Access sheath refers to the investigational/control access sheath</i> | <input type="checkbox"/> Right femoral vein <input type="checkbox"/> Left femoral vein <input type="checkbox"/> Both right and left insertion sites |
| Access Sheath <i>*Access sheath refers to the investigational/control access sheath</i> | <input type="checkbox"/> Single Curve <input type="checkbox"/> Double Curve <input type="checkbox"/> Both Single Curve and Double Curve used <input type="checkbox"/> VizaraMed Multiflex Steerable Sheath <input type="checkbox"/> None of the above, other, specify: __ |
| Final Access Sheath used | _____ Fr. |
| Transseptal method | <input type="checkbox"/> Mechanical needle puncture <input type="checkbox"/> Radiofrequency needle puncture |

| | |
|--|--|
| What imaging was used to determine release criteria | <input type="checkbox"/> TEE <input type="checkbox"/> Flouro/Angio |
| Peri-device leak present? | <input type="checkbox"/> Yes, _____ mm <input type="checkbox"/> No |
| Time of Access Sheath removal (24 HR) <i>*Access sheath refers to the investigational/control access sheath</i> | _____ : _____ |
| Vascular hemostasis method <i>(Please select at least one response)</i> | <input type="checkbox"/> Vascular closure device <input type="checkbox"/> Suture-mediated <input type="checkbox"/> Manual compression |
| Low ACT during procedure | |
| High ACT during procedure | |
| Total fluoroscopy time (minutes) | |
| Total contrast used (mL) | |
| Estimated blood loss (mL) | |
| Total Heparin Used | <input type="checkbox"/> _____ ml <input type="checkbox"/> _____ Units <input type="checkbox"/> Other: _____ |
| Was protamine used? | <input type="checkbox"/> Yes <input type="checkbox"/> No |
| Time subject left catheterization lab (24 HR) | _____ : _____ |
| Were any other medical procedures performed? <i>(Check all that apply)</i> | <input type="checkbox"/> Yes <input type="checkbox"/> No |
| | <input type="checkbox"/> LAA Device, specify: _____ <input type="checkbox"/> Pericardial Drain <input type="checkbox"/> Pericardiocentesis <input type="checkbox"/> Conversion to open heart surgery <input type="checkbox"/> Sternotomy <input type="checkbox"/> Other, specify: _____ |

Complete left atrial seal?

- Yes
 No

Were there any new adverse events?

- Yes (*Complete an Adverse Event Form*)
 No

Did the subject receive the intended implant?

- Yes
 No

If No, specify why:

Additional Procedure Notes

Site Personnel Signature

___/___/___
Date (DD/MMM/YYYY)

**Implanting Investigator
Signature**

___/___/___
Date (DD/MMM/YYYY)

| | |
|---|---|
| Date of Additional Procedure | ____/____/____ (DD/MMM/YYYY) |
| Study Procedure | <input type="checkbox"/> Watchman FLX <input type="checkbox"/> Watchman FLX Pro <input type="checkbox"/> Amulet <input type="checkbox"/> Other |
| Investigator (Operating Physician) First Name | |
| Investigator (Operating Physician) Last Name | |
| Primary Imager First Name | |
| Primary Imager Last Name | |
| What loading dose was prescribed to the patient prior to the procedure? | <input type="checkbox"/> 81-100 mg Aspirin <input type="checkbox"/> 325 mg Aspirin <input type="checkbox"/> No loading dose prescribed prior to index procedure <input type="checkbox"/> Other: ____ |
| Procedure start time (24 HR) (Defined as time of first sheath insertion in primary venous access site) | ____ : ____ |
| Access Sheath Insertion site <i>*Access sheath refers to the investigational/control access sheath</i> | <input type="checkbox"/> Right femoral vein <input type="checkbox"/> Left femoral vein <input type="checkbox"/> Both right and left insertion sites |
| Access Sheath <i>*Access sheath refers to the investigational/control access sheath</i> | <input type="checkbox"/> Single Curve <input type="checkbox"/> Double Curve <input type="checkbox"/> Both Single Curve and Double Curve used <input type="checkbox"/> VizaraMed Multiflex Steerable Sheath <input type="checkbox"/> None of the above, other, specify: __ |
| Final Access Sheath used | ____ Fr. |
| Transseptal method | <input type="checkbox"/> Mechanical needle puncture <input type="checkbox"/> Radiofrequency needle puncture |

| | |
|--|---|
| Complete left atrial seal? | <input type="checkbox"/> Yes <input type="checkbox"/> No |
| What imaging was used to determine release criteria | <input type="checkbox"/> TEE <input type="checkbox"/> Flouro/Angio |
| Peri-device leak present? | <input type="checkbox"/> Yes, _____ mm <input type="checkbox"/> No |
| Time of Access Sheath removal (24 HR) <i>*Access sheath refers to the investigational/control access sheath</i> | _____ : _____ |
| Vascular hemostasis method <i>(Please select at least one response)</i> | <input type="checkbox"/> Vascular closure device <input type="checkbox"/> Suture-mediated <input type="checkbox"/> Manual compression |
| Did any device deficiencies occur? | <input type="checkbox"/> Yes Specify: <input type="checkbox"/> No |
| Were there any new adverse events? | <input type="checkbox"/> Yes <i>(Complete an Adverse Event Form)</i> <input type="checkbox"/> No |

Site Personnel Signature

___/___/___
Date (DD/MMM/YYYY)

Implanting Investigator Signature

___/___/___
Date (DD/MMM/YYYY)

Implant LAA Measurements

Source Data Transfer Tool

Site Number: _____ Subject ID: _____

Note: All three measurements must be collected for CLAAS Subjects. LAA Perpendicular Depth measurements are not required for Control patients.

Pre-Implant LAA Measurements:

| Angle | LAA Ostium Diameter (mm) | LAA Perpendicular Depth (mm) | LAA Maximum Length (mm) |
|------------|--------------------------|------------------------------|-------------------------|
| 0 Degree | | | |
| 45 Degree | | | |
| 90 Degree | | | |
| 135 Degree | | | |

Site Personnel Signature

___/___/___
Date (DD/MMM/YYYY)

This Worksheet is to be used at the Index Procedure.

| | | | |
|---|--|--------------------------------------|--|
| Was Echocardiogram/CT performed? | <input type="checkbox"/> Yes <input type="checkbox"/> No <i>(Complete protocol deviation form)</i> | | |
| Are the required images for this visit available? | <input type="checkbox"/> Yes <input type="checkbox"/> No <i>(Complete protocol deviation form)</i> | | |
| Were images uploaded into the Imaging Module? | <input type="checkbox"/> Yes <input type="checkbox"/> No | | |
| | <input type="checkbox"/> | | |
| Date echocardiogram/CT completed | ___ / ___ / ___ (DD/MMM/YYYY) | | |
| Imaging Type | <input type="checkbox"/> TTE – Transthoracic echocardiogram <input type="checkbox"/> TEE – Transesophageal echocardiogram <input type="checkbox"/> Cardiac CT <input type="checkbox"/> Cardiac MRI <input type="checkbox"/> Brain CT <input type="checkbox"/> MRI | | |
| Left atrial appendage visible | <input type="checkbox"/> Yes <input type="checkbox"/> No <input type="checkbox"/> Not Applicable | Left atrial appendage ostium > 40 mm | <input type="checkbox"/> Yes <i>(Review for I&E!)</i> <input type="checkbox"/> No |
| | | Left atrial appendage ostium < 10 mm | <input type="checkbox"/> Yes <i>(Review for I&E!)</i> <input type="checkbox"/> No |
| Can the LAA accommodate both a CLAAS or Control LAAO device? | <input type="checkbox"/> Yes <input type="checkbox"/> No <i>(Review for I&E!)</i> | | |
| If available, confirm if the following was noted on echo/CT: | | | |
| Dense spontaneous echo contrast consistent with thrombus? | <input type="checkbox"/> Yes <i>(Review for I&E!)</i> <input type="checkbox"/> No <input type="checkbox"/> Not Available | | |

| | | | |
|---|---|---|--|
| <p>Intra-cardiac thrombus</p> | <p><input type="checkbox"/> Yes (<i>Review for I&E!</i>) <input type="checkbox"/> No <input type="checkbox"/> Not Available</p> | <p>If yes, confirm location</p> | <p><input type="checkbox"/> Left atrium <input type="checkbox"/> Left atrial appendage <input type="checkbox"/> Left ventricle <input type="checkbox"/> Right atrium <input type="checkbox"/> Right ventricle <input type="checkbox"/> Other, specify: _____</p> |
| <p>Intra-cardiac vegetation</p> | <p><input type="checkbox"/> Yes <input type="checkbox"/> No <input type="checkbox"/> Not Available</p> | <p>If yes, confirm location</p> | <p><input type="checkbox"/> Left atrium <input type="checkbox"/> Left atrial appendage <input type="checkbox"/> Left ventricle <input type="checkbox"/> Right atrium <input type="checkbox"/> Right ventricle <input type="checkbox"/> Other, specify: _____</p> |
| <p>Patent foramen ovale warranting closure?</p> | <p><input type="checkbox"/> Yes <input type="checkbox"/> No <input type="checkbox"/> Not Available</p> | <p>If yes, is this a high risk PFO?</p> | <p><input type="checkbox"/> Yes (Review for I&E!) <input type="checkbox"/> No</p> |
| <p>Atrial septal defect?</p> | <p><input type="checkbox"/> Yes <input type="checkbox"/> No <input type="checkbox"/> Not Available</p> | <p>If yes, specify</p> | <p><input type="checkbox"/> Right to left shunt present <input type="checkbox"/> Left to right shunt present <input type="checkbox"/> Bidirectional shunt present <input type="checkbox"/> Unable to determine</p> |
| | | <p>If yes, does defect warrant closure?</p> | <p><input type="checkbox"/> Yes <input type="checkbox"/> No</p> |

| | | | |
|---|--|---|--|
| <p>Pericardial effusion present?</p> <p><input type="checkbox"/> Yes <input type="checkbox"/> No <input type="checkbox"/> Not Available</p> | | <p>If yes, select type</p> | <p><input type="checkbox"/> Circumferential <input type="checkbox"/> Loculated</p> |
| | | <p>If yes, select size</p> | <p><input type="checkbox"/> Trivial <input type="checkbox"/> Small (<1 cm) <input type="checkbox"/> Moderate (1-2 cm) <input type="checkbox"/> Large (>2 cm and <5cm) <input type="checkbox"/> Large (> 5cm) (Review for I&E!)</p> |
| | | <p>If yes, Do any of the following apply? (Review for I&E!)</p> | <p><input type="checkbox"/> Symptomatic <input type="checkbox"/> Sign or symptom of acute or chronic pericarditis <input type="checkbox"/> Evidence of tamponade physiology</p> |

Site Personnel Signature

___/___/___
Date (DD/MMM/YYYY)

Note: This Device Deficiency form is for CLAAS only. To report a deficiency or malfunction for the CONTROL procedures, please follow the manufacturer's instructions.

| | | |
|--|--|---|
| Date of Device Deficiency | ___ / ___ / _____ (DD/MMM/YYYY) | |
| Component (select one) | <input type="checkbox"/> CLAAS Implant Regular 27mm <input type="checkbox"/> CLAAS Implant Large 35mm <input type="checkbox"/> Access Sheath Regular (27mm) Single Curve <input type="checkbox"/> Access Sheath Regular (27mm) Double Curve <input type="checkbox"/> Access Sheath Large (35mm) Single Curve <input type="checkbox"/> Access Sheath Large (35mm) Double Curve <input type="checkbox"/> Delivery Catheter Regular 27mm <input type="checkbox"/> Delivery Catheter Large 35mm | |
| Lot # | | |
| Deficiency occurred | <input type="checkbox"/> During procedure prep <input type="checkbox"/> During procedure | <input type="checkbox"/> Other, specify: _____ |
| Deficiency due to | <input type="checkbox"/> Device malfunction <input type="checkbox"/> Use error | <input type="checkbox"/> Inadequate labeling <input type="checkbox"/> Other, specify: _____ |
| Did an adverse event occur due to the deficiency? | <input type="checkbox"/> Yes (Complete an Adverse Event Form and follow reporting guidelines per protocol) <input type="checkbox"/> No | |
| Outcome of the device deficiency | <input type="checkbox"/> Used another CLAAS product <input type="checkbox"/> CLAAS device embolized | <input type="checkbox"/> Procedure terminated <input type="checkbox"/> Other, describe: _____ |
| Will the device be returned to Sponsor/Manufacturer? | <input type="checkbox"/> Yes, Please follow the device return instructions <input type="checkbox"/> No | |
| Summary of device deficiency | | |

Note: If utilizing as source (no other source exists)- form should be signed by device implanter.

Site Personnel Signature

___ / ___ / _____
Date (DD/MMM/YYYY)



CONFORM CLAAS Implant

Source Data Transfer Tool

Site Number: _____ Subject ID: _____

Note: Please keep the Investigational Product Sticker of any device opened/used. If more than 2 devices were used, please complete another form if using as source.

| | | | |
|--|---|--|---|
| CLAAS Device Size | <input type="checkbox"/> Regular (27 MM) <input type="checkbox"/> Large (35 MM) | CLAAS Device Size | <input type="checkbox"/> Regular (27 MM) <input type="checkbox"/> Large (35 MM) |
| Lot # | <i>Place product sticker here</i> | Lot # | <i>Place product sticker here</i> |
| Device Outcome | <input type="checkbox"/> Used <input type="checkbox"/> Opened, Not Used <input type="checkbox"/> Opened, Used, Disposed <input type="checkbox"/> Opened, Used, Returned <input type="checkbox"/> Opened, Not Used, Returned | Device Outcome | <input type="checkbox"/> Used <input type="checkbox"/> Opened, Not Used <input type="checkbox"/> Opened, Used, Disposed <input type="checkbox"/> Opened, Used, Returned <input type="checkbox"/> Opened, Not Used, Returned |
| Did device meet position criteria? | <input type="checkbox"/> Yes <input type="checkbox"/> No | Did device meet position criteria? | <input type="checkbox"/> Yes <input type="checkbox"/> No |
| Did device meet anchor criteria? | <input type="checkbox"/> Yes <input type="checkbox"/> No | Did device meet anchor criteria? | <input type="checkbox"/> Yes <input type="checkbox"/> No |
| Did device meet seal criteria? | <input type="checkbox"/> Yes <input type="checkbox"/> No | Did device meet seal criteria? | <input type="checkbox"/> Yes <input type="checkbox"/> No |
| Was partial resheath attempted? | <input type="checkbox"/> Yes, number of partial attempts: _____ <input type="checkbox"/> No | Was partial resheath attempted? | <input type="checkbox"/> Yes, number of partial attempts: _____ <input type="checkbox"/> No |
| Was a full resheath attempted? | <input type="checkbox"/> Yes <input type="checkbox"/> No | Was a full resheath attempted? | <input type="checkbox"/> Yes <input type="checkbox"/> No |
| Did device deficiency or device malfunction occur? | <input type="checkbox"/> Yes <i>If yes, complete a Device Deficiency Form</i> <input type="checkbox"/> No | Did device deficiency or device malfunction occur? | <input type="checkbox"/> Yes <i>If yes, complete a Device Deficiency Form</i> <input type="checkbox"/> No |

Site Personnel Signature

____/____/_____
Date (DD/MMM/YYYY)



CLAS Delivery System

Source Data Transfer Tool

Site Number: _____ Subject ID: _____

Note: Please keep the Investigational Product Sticker of any device opened/used. If more than 2 devices were used, please complete another form if utilizing as source.

| | | | |
|--|---|--|---|
| Access Sheath | <input type="checkbox"/> Regular (27 MM) Single Curve <input type="checkbox"/> Regular (27 MM) Double Curve <input type="checkbox"/> Large (35 MM) Single Curve <input type="checkbox"/> Large (35 MM) Double Curve | Access Sheath | <input type="checkbox"/> Regular (27 MM) Single Curve <input type="checkbox"/> Regular (27 MM) Double Curve <input type="checkbox"/> Large (35 MM) Single Curve <input type="checkbox"/> Large (35 MM) Double Curve |
| Lot Number | <i>Place Product Sticker here</i> | Lot Number | <i>Place Product Sticker here</i> |
| Outcome | <input type="checkbox"/> Used <input type="checkbox"/> Opened, Not Used <input type="checkbox"/> Opened, Used, Disposed <input type="checkbox"/> Opened, Used, Returned <input type="checkbox"/> Opened, Not Used, Returned | Outcome | <input type="checkbox"/> Used <input type="checkbox"/> Opened, Not Used <input type="checkbox"/> Opened, Used, Disposed <input type="checkbox"/> Opened, Used, Returned <input type="checkbox"/> Opened, Not Used, Returned |
| Did device deficiency or device malfunction occur? | <input type="checkbox"/> Yes <i>If yes, complete a Device Deficiency Form</i> <input type="checkbox"/> No | Did device deficiency or device malfunction occur? | <input type="checkbox"/> Yes <i>If yes, complete a Device Deficiency Form</i> <input type="checkbox"/> No |

_____/_____/_____
Date (DD/MMM/YYYY)

Site Personnel Signature

| | |
|--|---|
| Control Product Used | |
| <input type="checkbox"/> Amulet | <input type="checkbox"/> 16 mm <input type="checkbox"/> 18 mm <input type="checkbox"/> 20 mm <input type="checkbox"/> 22 mm <input type="checkbox"/> 25 mm <input type="checkbox"/> 28 mm <input type="checkbox"/> 31 mm <input type="checkbox"/> 34 mm |
| <input type="checkbox"/> Watchman FLX | <input type="checkbox"/> 20 mm <input type="checkbox"/> 24 mm <input type="checkbox"/> 27 mm <input type="checkbox"/> 31 mm <input type="checkbox"/> 35 mm |
| <input type="checkbox"/> Watchman FLX PRO | <input type="checkbox"/> 40 mm |
| Please confirm the primary reason for selection of the commercially available device | <input type="checkbox"/> Subject Anatomy <input type="checkbox"/> Investigator Preference <input type="checkbox"/> Other, specify: _____ |
| Device Outcome | <input type="checkbox"/> Used <input type="checkbox"/> Opened, Not Used <input type="checkbox"/> Opened, Used, Disposed <input type="checkbox"/> Opened, Used, Returned <input type="checkbox"/> Opened, Not Used, Returned |
| Did device meet release criteria per Manufacturer DFU/IFU? | <input type="checkbox"/> Yes <input type="checkbox"/> No |
| Was partial recapture attempted? | <input type="checkbox"/> Yes, number of partial attempts: _____ <input type="checkbox"/> No |
| Was full recapture attempted? | <input type="checkbox"/> Yes <input type="checkbox"/> No |
| Did device deficiency or device malfunction occur? | <input type="checkbox"/> Yes <i>If yes, provide a brief description below</i> <input type="checkbox"/> No |

Site Number: _____ Subject ID: _____

Device deficiency or malfunction
description:

Device Deficiency due to:

- Device Malfunction
- User error
- Inadequate labeling
- Other

Note: If utilizing as source (no other source exists)- form should be signed by device implanter.

Site Personnel Signature

___/___/_____
Date (DD/MMM/YYYY)

Site Number: _____ Subject ID: _____

| | |
|--|---|
| <p>Did Subject meet eligibility criteria before Procedure Day ?</p> | <p><input type="checkbox"/> Yes <input type="checkbox"/> No (Complete the Study Exit form accordingly. Classify subject as “Screen Failure” -- Subject did not meet I/E criteria prior to index procedure.)</p> |
| <p>Did Subject undergo Procedure TEE ?</p> | <p><input type="checkbox"/> Yes <input type="checkbox"/> No (Complete the Study Exit form accordingly. If the subject did not undergo the procedural TEE, the subject should be exited and classified as a “Screen Failure – Subject did not meet I/E criteria prior to index procedure” (or “Withdrawn” if the subject withdrew).</p> |
| <p>Did Subject continue to meet eligibility criteria after the Procedural TEE ?</p> | <p><input type="checkbox"/> Yes <input type="checkbox"/> No (Complete the study exit form accordingly. If the subject did not continue to meet eligibility criteria after the procedural TEE, subject should complete all visits through 45-Days with only Visit Information and QVSFS forms required. After completion of the 45-Day Visit, the subject should be exited and classified as “Screen Failure – Subject did not meet I/E criteria after the Index Procedure TEE was performed and prior to the Access Sheath crossed the body.)</p> |
| <p>Did any component of the investigational or control device (e.g. access sheath) enter the subject’s body ?</p> | <p><input type="checkbox"/> Yes <input type="checkbox"/> No (Complete the study exit form accordingly. If the subject underwent procedural TEE but no component of the investigational or control device entered the subject’s body, subject should complete all visits through 45-Days with only Visit Information and QVSFS forms required. After completion of the 45-Day Visit, the subject should be exited and classified as “Screen Failure – Subject did not meet I/E criteria after the Index Procedure TEE was performed and prior to the Access Sheath crossed the body.)</p> |
| <p>Did the subject receive an LAAO implant?</p> | <p><input type="checkbox"/> Yes (If the subject underwent procedural TEE, AND a component of the investigational or control device entered the subject’s body, AND received an implant, the subject should be followed for the full 5-Year Protocol.) <input type="checkbox"/> No (Complete the study exit form accordingly. If the subject underwent procedural TEE, AND a component of the investigational or control device entered the subject’s body, BUT the subject DID NOT RECEIVE AN IMPLANT, the subject should complete all primary safety and efficacy endpoints via phone call/telehealth (including Pre-Discharge, 7-Day, 45-Day, 6-Month, 12-Month, and 18-Month Visits. Once the subject completes the 18-Month Visit, the subject should be exited as “Withdrawn – No Implant received at index procedure (after IMPLANT imaging, Access Sheath crossed the body)”)</p> |
| <p>Did the subject receive the intended LAAO implant (e.g., the device they were randomized to)?</p> | <p><input type="checkbox"/> Yes <input type="checkbox"/> No (If the subject underwent procedural TEE, AND a component of the investigational or control device entered the subject’s body, AND received an implant, <u>even if it was not the intended implant</u> (e.g., a patient randomized to CLAAS was not able to get the CLAAS device and received a Control device), the subject should be followed for the full 5-Year Protocol.) Protocol Deviation should be entered for “Procedure /Assessment incomplete or not done” and, Additional Description of Deviation text box should include “Randomized to CLAAS – received CONTROL”</p> |

Site Personnel Signature

____ / ____ / ____
Date (DD/MMM/YYYY)

Echocardiographic Exclusion Criteria

REMINDER: Procedural ultrasound imaging will be performed by a qualified physician who is *not* the implanting physician.

Potential subjects will be excluded if **ANY** of the following conditions apply

| Exclusion Criteria | Yes | No |
|---|--------------------------|--------------------------|
| 1. Left atrial appendage cannot accommodate either a commercially available device of the CLAAS device per manufacturer IFU (e.g., the anatomy and sizing must be appropriate for both devices in order to be enrolled in the trial)? | <input type="checkbox"/> | <input type="checkbox"/> |
| 2. Intracardiac thrombus or dense spontaneous echo contrast consistent with thrombus, as visualized by TEE prior to implant? | <input type="checkbox"/> | <input type="checkbox"/> |
| 3. Left ventricular ejection fraction (LVEF) < 30%? | <input type="checkbox"/> | <input type="checkbox"/> |
| 4. Existing circumferential pericardial effusion > 10 mm or symptomatic pericardial effusion, signs, or symptoms of acute or chronic pericarditis, or evidence of tamponade physiology? | <input type="checkbox"/> | <input type="checkbox"/> |
| 5. Atrial septal defect that warrants closure? | <input type="checkbox"/> | <input type="checkbox"/> |
| 6. High risk patent foramen ovale (PFO), defined as an atrial septal aneurysm (exclusion > 15 mm or length > 15 mm) or large shunt (early [within 3 beats] and/or substantial passage of bubbles, e.g. > 20)? | <input type="checkbox"/> | <input type="checkbox"/> |
| 7. Moderate or severe mitral valve stenosis (mitral valve area < 1.5 cm ²)? | <input type="checkbox"/> | <input type="checkbox"/> |
| 8. Complex atheroma with mobile plaque of the descending aorta and/or aortic arch? | <input type="checkbox"/> | <input type="checkbox"/> |
| 9. Evidence of cardiac tumor? | <input type="checkbox"/> | <input type="checkbox"/> |

If utilizing as source (no other source exists)- form should be signed by device implanter or echocardiographer present at implant.

Site Personnel Signature

___/___/_____
Date (DD/MMM/YYYY)

* If any of the listed exclusions are marked as YES, the subject shall be considered a Screen Failure and will be followed for 45 days to evaluate safety.

Original source should be obtained from a direct laboratory report from Subject Medical Record. Laboratory results to be reviewed by delegated investigator (either directly on lab report or as Source here) as relates to subject safety and general INC/EXC Criteria.

Pre-procedure oral anticoagulation should be managed as per site protocol. Warfarin should be discontinued in accordance with site standard of care practices including INR levels on the day of the procedure. We are not collecting day of procedure INR levels.

Laboratory Collection at Procedure required **within 48 hours of implant.**

Date of Hematology: ____ / ____ / ____ (DD/MMM/YYYY)

Not Done (ENTER PD)

| <i>Laboratory Assessment</i> | <i>Results Value/ unit</i> | <i>Clinically Significant</i> |
|------------------------------|----------------------------|---|
| Hemoglobin | | <input type="checkbox"/> Yes <input type="checkbox"/> No |
| Hematocrit | | <input type="checkbox"/> Yes <input type="checkbox"/> No |
| Platelet Count | | <input type="checkbox"/> Yes <input type="checkbox"/> No |

If utilizing this form as source (i.e., no other source exists), this form should be signed by Site Investigator.

Site Personnel Signature

____ / ____ / ____
Date (DD/MMM/YYYY)

PRE-DISCHARGE

CONFORM Visit Information

Source Data Transfer Tool

Site Number: _____ Subject ID: _____

| | | |
|--|---|---|
| Visit Timepoint | <input type="checkbox"/> Pre-Discharge <input type="checkbox"/> Day 7 <input type="checkbox"/> Day 45 <input type="checkbox"/> 6 Months <input type="checkbox"/> 12 Months <input type="checkbox"/> 18 Months | <input type="checkbox"/> 2 Year <input type="checkbox"/> 3 Year <input type="checkbox"/> 4 Year <input type="checkbox"/> 5 Year <input type="checkbox"/> Not related to a study visit <input type="checkbox"/> Unscheduled visit |
| Was visit completed? | <input type="checkbox"/> Yes <input type="checkbox"/> No | |
| Visit Date | ____ / ____ / ____ (DD/MMM/YYYY) | |
| Visit Type | <input type="checkbox"/> Office/clinic visit <input type="checkbox"/> Telephone contact <input type="checkbox"/> Video link | |
| Were there any new or changes to existing Adverse Events? <i>If yes, please complete or update an Adverse Event CRF</i> | <input type="checkbox"/> Yes Was the event a suspected stroke or systemic embolism? <input type="checkbox"/> Yes <input type="checkbox"/> No <input type="checkbox"/> No | |
| Did the subject have any ER visits or hospitalizations since the last visit? | <input type="checkbox"/> Yes <input type="checkbox"/> No | |
| Were there any changes in patient medical history that are cardiovascular in etiology? | <input type="checkbox"/> Yes <i>If yes, specify: _____</i> <input type="checkbox"/> No | |
| Were there any new changes to existing Concomitant Medications? | <input type="checkbox"/> Yes <i>(If yes, please add new or update Concomitant Medication CRF)</i> <input type="checkbox"/> No | |
| Was visit imaging done? | <input type="checkbox"/> Yes Are required images for this visit available? <input type="checkbox"/> Yes <input type="checkbox"/> No <input type="checkbox"/> No <input type="checkbox"/> N/A Imaging not required per protocol | |

Site Personnel Signature

____ / ____ / ____
Date (DD/MMM/YYYY)

| | |
|-----------------------------------|---|
| Were vital signs performed? | <input type="checkbox"/> Yes <input type="checkbox"/> No |
| Date of vital sign measurements | ___ / ___ / ___ (DD/MMM/YYYY) |
| Height (xxx.xx) | _____ (cm / in) (circle one) |
| Weight (xxx.x) | _____ (kg / lb) (circle one) |
| BMI (xx.x) | _____ (kg/m ²) |
| Systolic Blood Pressure (xxx) | _____ (mmHg) |
| Diastolic Blood Pressure (xxx) | _____ (mmHg) |
| Heart Rate (xxx) | _____ (bpm) |

Site Personnel Signature

___ / ___ / ___
Date (DD/MMM/YYYY)

TTE is required to surveil for pericardial effusion. The study must be performed a minimum of 3 hours after discharge from cardiac catheterization laboratory.

| | |
|---|--|
| Was Echocardiogram/CT performed? | <input type="checkbox"/> Yes <input type="checkbox"/> No <i>(Complete protocol deviation form)</i> |
| Are the required images for this visit available? | <input type="checkbox"/> Yes <input type="checkbox"/> No <i>(Complete protocol deviation form)</i> |
| Was imaging uploaded into the Imaging Module? | <input type="checkbox"/> Yes <input type="checkbox"/> No |
| Date echocardiogram/CT completed | ___ / ___ / ___ (DD/MMM/YYYY) |
| What time was pre-discharge TTE performed? | _____ : _____ |
| Imaging Type | <input type="checkbox"/> TTE – Transthoracic echocardiogram <input type="checkbox"/> TEE – Transesophageal echocardiogram <input type="checkbox"/> Cardiac CT <input type="checkbox"/> Cardiac MRI <input type="checkbox"/> Brain CT <input type="checkbox"/> MRI |

If available, confirm if the following was noted on echo/CT:

| | |
|---|--|
| Left atrial appendage visible | <input type="checkbox"/> Yes <input type="checkbox"/> No <input type="checkbox"/> Not Available |
| Dense spontaneous echo contrast consistent with thrombus? | <input type="checkbox"/> Yes <input type="checkbox"/> No <input type="checkbox"/> Not Available |
| Intra-cardiac thrombus | <input type="checkbox"/> Yes <i>(Complete AE form)</i> <input type="checkbox"/> No <input type="checkbox"/> Not available |
| | If yes, confirm location <input type="checkbox"/> Left atrium <input type="checkbox"/> Left atrial appendage <input type="checkbox"/> Left ventricle <input type="checkbox"/> Right atrium <input type="checkbox"/> Right ventricle <input type="checkbox"/> Other, specify: _____ |

| | | | |
|--|--|--------------------------------------|--|
| Intra-cardiac vegetation | <input type="checkbox"/> Yes <input type="checkbox"/> No <input type="checkbox"/> Not available | If yes, confirm location | <input type="checkbox"/> Left atrium <input type="checkbox"/> Left atrial appendage <input type="checkbox"/> Left ventricle <input type="checkbox"/> Right atrium <input type="checkbox"/> Right ventricle <input type="checkbox"/> Other, specify: _____ |
| Patent foramen ovale warranting closure? | <input type="checkbox"/> Yes <input type="checkbox"/> No <input type="checkbox"/> Not Available | If yes, is this a high risk PFO? | <input type="checkbox"/> Yes (<i>Complete AE Form</i>) <input type="checkbox"/> No |
| Atrial septal defect? | <input type="checkbox"/> Yes <input type="checkbox"/> No <input type="checkbox"/> Not available | If yes, specify | <input type="checkbox"/> Right to left shunt present <input type="checkbox"/> Left to right shunt present <input type="checkbox"/> Bidirectional shunt <input type="checkbox"/> Unable to determine |
| | | If yes, does defect warrant closure? | <input type="checkbox"/> Yes (<i>Complete AE form</i>) <input type="checkbox"/> No |
| Pericardial effusion present? | <input type="checkbox"/> Yes <input type="checkbox"/> No <input type="checkbox"/> Not available | | |
| | If yes, select type | | <input type="checkbox"/> Circumferential <input type="checkbox"/> Loculated |
| | If yes, select size <i>(Pericardial effusion deemed as trivial or small does not meet adverse event reporting criteria)</i> | | <input type="checkbox"/> Trivial <input type="checkbox"/> Small (<1 cm) <input type="checkbox"/> Moderate (1-2 cm) <input type="checkbox"/> Large (>2 cm and <5cm) <input type="checkbox"/> Large (>5 cm) (Review for I&E!) |
| | If yes, Do any of the following apply? <i>(Check all that apply)</i> | | <input type="checkbox"/> Symptomatic <input type="checkbox"/> Sign or symptom of acute or chronic pericarditis <input type="checkbox"/> Evidence of tamponade physiology |

Site Personnel Signature

___/___/___
Date (DD/MMM/YYYY)

Was the NIHSS assessment completed?

- Yes
 No (*Complete Protocol Deviation form*)

Date of NIHSS assessment ___ / ___ / ___ (DD/MMM/YYYY)

| 1(a) – Level of consciousness | |
|---|------------------------------|
| Alert, keenly responsive | <input type="checkbox"/> (0) |
| Not alert; but arousable by minor stimulation to obey, answer or respond | <input type="checkbox"/> (1) |
| Not alert' requires repeated stimulation to attend, or is obtunded and requires strong or painful stimulation to make movements (not stereotyped) | <input type="checkbox"/> (2) |
| Responds only with reflex motor or autonomic unresponsive, flaccid or areflexic | <input type="checkbox"/> (3) |
| 1(b) – Level of consciousness questions | |
| Answers both questions correctly | <input type="checkbox"/> (0) |
| Answers one question correctly | <input type="checkbox"/> (1) |
| Answers neither question correctly | <input type="checkbox"/> (2) |
| 1(c) – Level of consciousness command | |
| Performs both tasks correctly | <input type="checkbox"/> (0) |
| Performs one task correctly | <input type="checkbox"/> (1) |
| Performs neither task correctly | <input type="checkbox"/> (2) |
| 2 – Best gaze | |
| Normal | <input type="checkbox"/> (0) |
| Partial gaze palsy; gaze is abnormal in one or both eyes, but forced deviation or total gaze paresis is not present | <input type="checkbox"/> (1) |
| Forced deviation, or total gaze paresis not overcome by the oculoccephalic maneuver | <input type="checkbox"/> (2) |
| 3 – Visual | |
| No visual loss | <input type="checkbox"/> (0) |
| Partial hemianopia | <input type="checkbox"/> (1) |
| Complete hemianopia | <input type="checkbox"/> (2) |
| Bilateral hemianopia (blind including cortical blindness) | <input type="checkbox"/> (3) |

| 4 – Facial palsy | |
|---|-------------------------------|
| Normal symmetrical movements | <input type="checkbox"/> (0) |
| Minor paralysis (flattened nasolabial fold, asymmetry on smiling) | <input type="checkbox"/> (1) |
| Partial paralysis (total or near-total paralysis of lower face) | <input type="checkbox"/> (2) |
| Complete paralysis of one or both sides (absence of facial movement in the upper and lower face) | <input type="checkbox"/> (3) |
| 5(a) – Motor arm - left | |
| No drift, limb holds 90 (or 45) degrees for 10 full seconds | <input type="checkbox"/> (0) |
| Drift; limb holds 90 (or 45) degrees, but drifts down before full 10 seconds; does not hit bed or other support | <input type="checkbox"/> (1) |
| Some effort against gravity; limb cannot get to or maintain (if cued) 90 (or 45) degrees; drifts down to bed, but has some effort against gravity | <input type="checkbox"/> (2) |
| No effort against gravity, limb falls | <input type="checkbox"/> (3) |
| No movement | <input type="checkbox"/> (4) |
| Amputation or joint fusion, Explain | <input type="checkbox"/> (UN) |
| 5(b) – Motor arm - right | |
| No drift, limb holds 90 (or 45) degrees for 10 full seconds | <input type="checkbox"/> (0) |
| Drift; limb holds 90 (or 45) degrees, but drifts down before full 10 seconds; does not hit bed or other support | <input type="checkbox"/> (1) |
| Some effort against gravity; limb cannot get to or maintain (if cued) 90 (or 45) degrees; drifts down to bed, but has some effort against gravity | <input type="checkbox"/> (2) |
| No effort against gravity, limb falls | <input type="checkbox"/> (3) |
| No movement | <input type="checkbox"/> (4) |
| Amputation or joint fusion, Explain: | <input type="checkbox"/> (UN) |

| 6(a) Motor leg - left | |
|--|-------------------------------|
| No drift, leg holds 30-degree position for full 5 seconds | <input type="checkbox"/> (0) |
| Drift: leg falls by the end of the 5-second period but does not hit bed | <input type="checkbox"/> (1) |
| Some effort against gravity; leg falls to bed by 5 seconds, but has some effort against gravity | <input type="checkbox"/> (2) |
| No effort against gravity, leg falls to bed immediately | <input type="checkbox"/> (3) |
| No movement | <input type="checkbox"/> (4) |
| Amputation or joint fusion, Explain: | <input type="checkbox"/> (UN) |
| 6(a) Motor leg - right | |
| No drift, leg holds 30-degree position for full 5 seconds | <input type="checkbox"/> (0) |
| Drift: leg falls by the end of the 5-second period but does not hit bed | <input type="checkbox"/> (1) |
| Some effort against gravity; leg falls to bed by 5 seconds, but has some effort against gravity | <input type="checkbox"/> (2) |
| No effort against gravity, leg falls to bed immediately | <input type="checkbox"/> (3) |
| No movement | <input type="checkbox"/> (4) |
| Amputation or joint fusion, Explain: | <input type="checkbox"/> (UN) |
| 7 – Limb ataxia | |
| Absent | <input type="checkbox"/> (0) |
| Present in one limb | <input type="checkbox"/> (1) |
| Present in two limbs | <input type="checkbox"/> (2) |
| Amputation or joint fusion, Explain: | <input type="checkbox"/> (UN) |
| 8 – Sensory | |
| Normal; no sensory loss | <input type="checkbox"/> (0) |
| Mild-to-moderate sensory loss; patient feels pinprick less sharp or dull on the affected side; or there is a loss of superficial pain with pinprick, but patient is aware of being touched | <input type="checkbox"/> (1) |
| Severe to total sensory loss; patient is not aware of being touched in the face, arm, and leg | <input type="checkbox"/> (2) |

| 9 – Best language | |
|---|-------------------------------|
| No aphasia; normal | <input type="checkbox"/> (0) |
| Mild-to-moderate aphasia; some obvious loss of fluency or facility or comprehension, without significant limitation on ideas expressed or form of expression. Reduction of speech and/or comprehension, however, makes conversion about provided materials difficult or impossible. <i>For example, in conversation about provided materials, examiner can identify picture or naming card content from patient's response.</i> | <input type="checkbox"/> (1) |
| Severe aphasia; all communication is through fragmentary expression; great need for inference questioning and guessing by the listener. Range of information that can be exchanged is limited; listener carries burden of communication. Examiner cannot identify materials provided form patient response. | <input type="checkbox"/> (2) |
| Mute, global aphasia; no usable speech or auditory comprehension | <input type="checkbox"/> (3) |
| 10 – Dysarthria | |
| Normal | <input type="checkbox"/> (0) |
| Mild-to-moderate dysarthria; patient slurs at least some word and, at worst, can be understood with some difficult. | <input type="checkbox"/> (1) |
| Severe dysarthria; patient's speech is so slurred as to be unintelligible in the absence or out of proportion to any dysphasia, or is mute/anarthric | <input type="checkbox"/> (2) |
| Intubated or another physical barrier, Explain: | <input type="checkbox"/> (UN) |
| 11 – Extinction and inattention (formerly neglect) | |
| No abnormality | <input type="checkbox"/> (0) |
| Visual, tactile, auditory, spatial, or personal inattention or extinction to bilateral simultaneous stimulation in one of the sensory modalities. | <input type="checkbox"/> (1) |
| Profound hemi-inattention or extinction to more than one modality; does not recognize own hand or orients to only one side of space. | <input type="checkbox"/> (2) |

Site Personnel Signature

___/___/_____
Date (DD/MMM/YYYY)

Was the mRS assessment completed?

- Yes
 No (*Complete protocol deviation form*)

Is this assessment performed because of a neurological event?

- Yes *If yes, assess for an adverse event*
 No

Date of mRS assessment

___ / ___ / ___ (DD/MMM/YYYY)

| Level of Consciousness | Response |
|--|--------------------------|
| 0 = No symptoms at all | <input type="checkbox"/> |
| 1 = No significant disability despite symptoms; able to carry out all usual duties and activities | <input type="checkbox"/> |
| 2 = Slight disability; unable to carry out all previous activities but able to look after own affairs without assistance | <input type="checkbox"/> |
| 3 = Moderate disability; requiring some help, but able to walk without assistance | <input type="checkbox"/> |
| 4 = Moderately severe disability; unable to walk without assistance, and unable to attend to own body needs without assistance | <input type="checkbox"/> |
| 5 = Severe disability; bedridden, incontinent, and requiring constant nursing care and attention | <input type="checkbox"/> |
| Score | |

Site Personnel Signature___ / ___ / ___
Date (DD/MMM/YYYY)

7 DAY

CONFORM Visit Information

Source Data Transfer Tool

Site Number: _____ Subject ID: _____

| | | |
|--|---|---|
| Visit Timepoint | <input type="checkbox"/> Pre-Discharge <input type="checkbox"/> Day 7 <input type="checkbox"/> Day 45 <input type="checkbox"/> 6 Months <input type="checkbox"/> 12 Months <input type="checkbox"/> 18 Months | <input type="checkbox"/> 2 Year <input type="checkbox"/> 3 Year <input type="checkbox"/> 4 Year <input type="checkbox"/> 5 Year <input type="checkbox"/> Not related to a study visit <input type="checkbox"/> Unscheduled visit |
| Was visit completed? | <input type="checkbox"/> Yes <input type="checkbox"/> No | |
| Visit Date | ____ / ____ / ____ (DD/MMM/YYYY) | |
| Visit Type | <input type="checkbox"/> Office/clinic visit <input type="checkbox"/> Telephone contact <input type="checkbox"/> Video link | |
| Were there any new or changes to existing Adverse Events? <i>If yes, please complete or update an Adverse Event CRF</i> | <input type="checkbox"/> Yes Was the event a suspected stroke or systemic embolism? <input type="checkbox"/> Yes <input type="checkbox"/> No <input type="checkbox"/> No | |
| Did the subject have any ER visits or hospitalizations since the last visit? | <input type="checkbox"/> Yes <input type="checkbox"/> No | |
| Were there any changes in patient medical history that are cardiovascular in etiology? | <input type="checkbox"/> Yes <i>If yes, specify: _____</i> <input type="checkbox"/> No | |
| Were there any new changes to existing Concomitant Medications? | <input type="checkbox"/> Yes <i>(If yes, please add new or update Concomitant Medication CRF)</i> <input type="checkbox"/> No | |
| Was visit imaging done? | <input type="checkbox"/> Yes Are required images for this visit available? <input type="checkbox"/> Yes <input type="checkbox"/> No <input type="checkbox"/> No <input type="checkbox"/> N/A Imaging not required per protocol | |

Site Personnel Signature

____ / ____ / ____
Date (DD/MMM/YYYY)

CONFORM QVSFS

Source Data Transfer Tool

Site Number: _____ Subject ID: _____

| | |
|---|--|
| Was the QVSFS assessment completed? | <input type="checkbox"/> Yes <input type="checkbox"/> No <i>(Complete protocol deviation form)</i> |
| Is this assessment performed because of a neurological event? | <input type="checkbox"/> Yes <i>If yes, assess for an adverse event</i> <input type="checkbox"/> No |
| Date of QVSFS assessment | ___ / ___ / ___ (DD/MMM/YYYY) |

| Since the last study contact (by phone or clinic) | Yes | No | Unknown |
|--|--------------------------|--------------------------|--------------------------|
| 1. Were you told by a physician that you had a stroke? | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> |
| 2. Were you ever told by a physician that you had a TIA, ministroke, or a transient ischemic attack? | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> |
| 3. Have you ever had a sudden weakness on one side of your body? | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> |
| 4. Have you ever had a sudden numbness or dead feeling on one side of your body? | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> |
| 5. Have you ever had a sudden painless loss of vision in one or both eyes? | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> |
| 6. Have you ever suddenly lost one half of your vision? | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> |
| 7. Have you ever suddenly lost the ability to understand what people are saying? | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> |
| 8. Have you ever suddenly lost the ability to express yourself verbally or in writing? | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> |

Site Personnel Signature

___ / ___ / ___
Date (DD/MMM/YYYY)

45 DAY

CONFORM Visit Information

Source Data Transfer Tool

Site Number: _____ Subject ID: _____

| | | |
|--|---|---|
| Visit Timepoint | <input type="checkbox"/> Pre-Discharge <input type="checkbox"/> Day 7 <input type="checkbox"/> Day 45 <input type="checkbox"/> 6 Months <input type="checkbox"/> 12 Months <input type="checkbox"/> 18 Months | <input type="checkbox"/> 2 Year <input type="checkbox"/> 3 Year <input type="checkbox"/> 4 Year <input type="checkbox"/> 5 Year <input type="checkbox"/> Not related to a study visit <input type="checkbox"/> Unscheduled visit |
| Was visit completed? | <input type="checkbox"/> Yes <input type="checkbox"/> No | |
| Visit Date | ____ / ____ / ____ (DD/MMM/YYYY) | |
| Visit Type | <input type="checkbox"/> Office/clinic visit <input type="checkbox"/> Telephone contact <input type="checkbox"/> Video link | |
| Were there any new or changes to existing Adverse Events? <i>If yes, please complete or update an Adverse Event CRF</i> | <input type="checkbox"/> Yes Was the event a suspected stroke or systemic embolism? <input type="checkbox"/> Yes <input type="checkbox"/> No <input type="checkbox"/> No | |
| Did the subject have any ER visits or hospitalizations since the last visit? | <input type="checkbox"/> Yes <input type="checkbox"/> No | |
| Were there any changes in patient medical history that are cardiovascular in etiology? | <input type="checkbox"/> Yes <i>If yes, specify: _____</i> <input type="checkbox"/> No | |
| Were there any new changes to existing Concomitant Medications? | <input type="checkbox"/> Yes <i>(If yes, please add new or update Concomitant Medication CRF)</i> <input type="checkbox"/> No | |
| Was visit imaging done? | <input type="checkbox"/> Yes Are required images for this visit available? <input type="checkbox"/> Yes <input type="checkbox"/> No <input type="checkbox"/> No <input type="checkbox"/> N/A Imaging not required per protocol | |

Site Personnel Signature

____ / ____ / ____
Date (DD/MMM/YYYY)

| | |
|-----------------------------------|---|
| Were vital signs performed? | <input type="checkbox"/> Yes <input type="checkbox"/> No |
| Date of vital sign measurements | ___ / ___ / ___ (DD/MMM/YYYY) |
| Height (xxx.xx) | _____ (cm / in) (circle one) |
| Weight (xxx.x) | _____ (kg / lb) (circle one) |
| BMI (xx.x) | _____ (kg/m ²) |
| Systolic Blood Pressure (xxx) | _____ (mmHg) |
| Diastolic Blood Pressure (xxx) | _____ (mmHg) |
| Heart Rate (xxx) | _____ (bpm) |

Site Personnel Signature

___ / ___ / ___
Date (DD/MMM/YYYY)

For use with Follow Up Visits as needed (45-Day, 12 Months, and Unscheduled).

Reminders:

- **At 45 Days and 12 Months:**
- TEE or CT is mandatory per protocol at 45 Days and 12 Months for Implanted Subjects
- If a CT is completed and shows findings (i.e., leak or thrombus), a TEE is required to confirm the finding as soon as possible

| | |
|---|--|
| Was Echocardiogram/CT performed? | <input type="checkbox"/> Yes <input type="checkbox"/> No <i>(Complete protocol deviation form)</i> |
| Are the required images for this visit available? | <input type="checkbox"/> Yes <input type="checkbox"/> No <i>(Complete protocol deviation form)</i> |
| Were images uploaded into the Imaging Module? | <input type="checkbox"/> Yes <input type="checkbox"/> No |
| Time period of Imaging | <input type="checkbox"/> 45 Day <input type="checkbox"/> 12 Months <input type="checkbox"/> Unscheduled, specify: _____ |
| Date echocardiogram/CT completed | ___ / ___ / ___ (DD/MMM/YYYY) |
| Imaging Type | <input type="checkbox"/> TTE – Transthoracic echocardiogram <input type="checkbox"/> TEE – Transesophageal echocardiogram <input type="checkbox"/> Cardiac CT <input type="checkbox"/> Cardiac MRI <input type="checkbox"/> Brain CT <input type="checkbox"/> MRI |
| Left atrial appendage visible | <input type="checkbox"/> Yes <input type="checkbox"/> No <input type="checkbox"/> Not Available |

If available, confirm if the following was noted on echo/CT:

| | | | | |
|---|---|--|--------------------------|--|
| Dense spontaneous echo contrast consistent with Thrombus? | <input type="checkbox"/> Yes <input type="checkbox"/> No <input type="checkbox"/> Not Available | | | |
| Intra-cardiac thrombus | <table border="1"> <tr> <td><input type="checkbox"/> Yes <i>(Complete AE form)</i> <input type="checkbox"/> No <input type="checkbox"/> Not available</td> <td>If yes, confirm location</td> <td> <input type="checkbox"/> Left atrium <input type="checkbox"/> Left atrial appendage <input type="checkbox"/> Left ventricle <input type="checkbox"/> Right atrium <input type="checkbox"/> Right ventricle <input type="checkbox"/> Other, specify: _____ </td> </tr> </table> | <input type="checkbox"/> Yes <i>(Complete AE form)</i> <input type="checkbox"/> No <input type="checkbox"/> Not available | If yes, confirm location | <input type="checkbox"/> Left atrium <input type="checkbox"/> Left atrial appendage <input type="checkbox"/> Left ventricle <input type="checkbox"/> Right atrium <input type="checkbox"/> Right ventricle <input type="checkbox"/> Other, specify: _____ |
| <input type="checkbox"/> Yes <i>(Complete AE form)</i> <input type="checkbox"/> No <input type="checkbox"/> Not available | If yes, confirm location | <input type="checkbox"/> Left atrium <input type="checkbox"/> Left atrial appendage <input type="checkbox"/> Left ventricle <input type="checkbox"/> Right atrium <input type="checkbox"/> Right ventricle <input type="checkbox"/> Other, specify: _____ | | |

| | | | |
|--|--|--|--|
| Intra-cardiac vegetation | <input type="checkbox"/> Yes <input type="checkbox"/> No <input type="checkbox"/> Not available | If yes, confirm location | <input type="checkbox"/> Left atrium <input type="checkbox"/> Left atrial appendage <input type="checkbox"/> Left ventricle <input type="checkbox"/> Right atrium <input type="checkbox"/> Right ventricle <input type="checkbox"/> Other, specify: _____ |
| Patent foramen ovale warranting closure? | <input type="checkbox"/> Yes <input type="checkbox"/> No <input type="checkbox"/> Not available | Is this a high risk PFO? | <input type="checkbox"/> Yes <input type="checkbox"/> No |
| Atrial septal defect? | <input type="checkbox"/> Yes <input type="checkbox"/> No <input type="checkbox"/> Not available | If yes, specify | <input type="checkbox"/> Right to left shunt present <input type="checkbox"/> Left to right shunt present <input type="checkbox"/> Bidirectional shunt <input type="checkbox"/> Unable to determine |
| | | If yes, does defect warrant closure? | <input type="checkbox"/> Yes <input type="checkbox"/> No |
| Left atrial appendage occlusion device position stable and position unchanged? | <input type="checkbox"/> Yes <input type="checkbox"/> No <input type="checkbox"/> Not available | | |
| Peri-device leak present? | <input type="checkbox"/> Yes <input type="checkbox"/> No <input type="checkbox"/> Not available | If yes, specify (mm) | _____ mm |
| Pericardial effusion present? | <input type="checkbox"/> Yes (<i>Assess for AE</i>) <input type="checkbox"/> No <input type="checkbox"/> Not available | | |
| | If yes, select type | <input type="checkbox"/> Circumferential <input type="checkbox"/> Loculated | |
| | If yes, select size <i>*AE is reportable for pericardial effusions Moderate or larger</i> | <input type="checkbox"/> Trivial <input type="checkbox"/> Small (<1 cm) <input type="checkbox"/> Moderate (1-2 cm) <input type="checkbox"/> Large (>2 cm and <5cm) <input type="checkbox"/> Large (> 5 cm) | |
| | If yes, do any of the following apply? | <input type="checkbox"/> Symptomatic <input type="checkbox"/> Sign or symptom of acute or chronic pericarditis <input type="checkbox"/> Evidence of tamponade physiology | |

Device embolization?

- Yes (*Complete AE form*)
 No
 Not available

Site Personnel Signature

___/___/___
Date (DD/MMM/YYYY)

CONFORM QVSFS

Source Data Transfer Tool

Site Number: _____ Subject ID: _____

| | |
|---|--|
| Was the QVSFS assessment completed? | <input type="checkbox"/> Yes <input type="checkbox"/> No <i>(Complete protocol deviation form)</i> |
| Is this assessment performed because of a neurological event? | <input type="checkbox"/> Yes <i>If yes, assess for an adverse event</i> <input type="checkbox"/> No |
| Date of QVSFS assessment | ___ / ___ / ___ (DD/MMM/YYYY) |

| Since the last study contact (by phone or clinic) | Yes | No | Unknown |
|--|--------------------------|--------------------------|--------------------------|
| 1. Were you told by a physician that you had a stroke? | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> |
| 2. Were you ever told by a physician that you had a TIA, ministroke, or a transient ischemic attack? | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> |
| 3. Have you ever had a sudden weakness on one side of your body? | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> |
| 4. Have you ever had a sudden numbness or dead feeling on one side of your body? | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> |
| 5. Have you ever had a sudden painless loss of vision in one or both eyes? | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> |
| 6. Have you ever suddenly lost one half of your vision? | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> |
| 7. Have you ever suddenly lost the ability to understand what people are saying? | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> |
| 8. Have you ever suddenly lost the ability to express yourself verbally or in writing? | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> |

 Site Personnel Signature

___ / ___ / ___
 Date (DD/MMM/YYYY)

6 MONTH

CONFORM Visit Information

Source Data Transfer Tool

Site Number: _____ Subject ID: _____

| | | |
|--|---|---|
| Visit Timepoint | <input type="checkbox"/> Pre-Discharge <input type="checkbox"/> Day 7 <input type="checkbox"/> Day 45 <input type="checkbox"/> 6 Months <input type="checkbox"/> 12 Months <input type="checkbox"/> 18 Months | <input type="checkbox"/> 2 Year <input type="checkbox"/> 3 Year <input type="checkbox"/> 4 Year <input type="checkbox"/> 5 Year <input type="checkbox"/> Not related to a study visit <input type="checkbox"/> Unscheduled visit |
| Was visit completed? | <input type="checkbox"/> Yes <input type="checkbox"/> No | |
| Visit Date | ____ / ____ / ____ (DD/MMM/YYYY) | |
| Visit Type | <input type="checkbox"/> Office/clinic visit <input type="checkbox"/> Telephone contact <input type="checkbox"/> Video link | |
| Were there any new or changes to existing Adverse Events? <i>If yes, please complete or update an Adverse Event CRF</i> | <input type="checkbox"/> Yes Was the event a suspected stroke or systemic embolism? <input type="checkbox"/> Yes <input type="checkbox"/> No <input type="checkbox"/> No | |
| Did the subject have any ER visits or hospitalizations since the last visit? | <input type="checkbox"/> Yes <input type="checkbox"/> No | |
| Were there any changes in patient medical history that are cardiovascular in etiology? | <input type="checkbox"/> Yes <i>If yes, specify: _____</i> <input type="checkbox"/> No | |
| Were there any new changes to existing Concomitant Medications? | <input type="checkbox"/> Yes <i>(If yes, please add new or update Concomitant Medication CRF)</i> <input type="checkbox"/> No | |
| Was visit imaging done? | <input type="checkbox"/> Yes Are required images for this visit available? <input type="checkbox"/> Yes <input type="checkbox"/> No <input type="checkbox"/> No <input type="checkbox"/> N/A Imaging not required per protocol | |

Site Personnel Signature

____ / ____ / ____
Date (DD/MMM/YYYY)

| | |
|---|--|
| Was the QVSFS assessment completed? | <input type="checkbox"/> Yes <input type="checkbox"/> No <i>(Complete protocol deviation form)</i> |
| Is this assessment performed because of a neurological event? | <input type="checkbox"/> Yes <i>If yes, assess for an adverse event</i> <input type="checkbox"/> No |
| Date of QVSFS assessment | ___ / ___ / ___ (DD/MMM/YYYY) |

| Since the last study contact (by phone or clinic) | Yes | No | Unknown |
|--|--------------------------|--------------------------|--------------------------|
| 1. Were you told by a physician that you had a stroke? | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> |
| 2. Were you ever told by a physician that you had a TIA, ministroke, or a transient ischemic attack? | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> |
| 3. Have you ever had a sudden weakness on one side of your body? | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> |
| 4. Have you ever had a sudden numbness or dead feeling on one side of your body? | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> |
| 5. Have you ever had a sudden painless loss of vision in one or both eyes? | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> |
| 6. Have you ever suddenly lost one half of your vision? | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> |
| 7. Have you ever suddenly lost the ability to understand what people are saying? | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> |
| 8. Have you ever suddenly lost the ability to express yourself verbally or in writing? | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> |

Site Personnel Signature

___ / ___ / ___
Date (DD/MMM/YYYY)

12 MONTH

CONFORM Visit Information

Source Data Transfer Tool

Site Number: _____ Subject ID: _____

| | | |
|--|---|---|
| Visit Timepoint | <input type="checkbox"/> Pre-Discharge <input type="checkbox"/> Day 7 <input type="checkbox"/> Day 45 <input type="checkbox"/> 6 Months <input type="checkbox"/> 12 Months <input type="checkbox"/> 18 Months | <input type="checkbox"/> 2 Year <input type="checkbox"/> 3 Year <input type="checkbox"/> 4 Year <input type="checkbox"/> 5 Year <input type="checkbox"/> Not related to a study visit <input type="checkbox"/> Unscheduled visit |
| Was visit completed? | <input type="checkbox"/> Yes <input type="checkbox"/> No | |
| Visit Date | ____ / ____ / ____ (DD/MMM/YYYY) | |
| Visit Type | <input type="checkbox"/> Office/clinic visit <input type="checkbox"/> Telephone contact <input type="checkbox"/> Video link | |
| Were there any new or changes to existing Adverse Events? <i>If yes, please complete or update an Adverse Event CRF</i> | <input type="checkbox"/> Yes Was the event a suspected stroke or systemic embolism? <input type="checkbox"/> Yes <input type="checkbox"/> No <input type="checkbox"/> No | |
| Did the subject have any ER visits or hospitalizations since the last visit? | <input type="checkbox"/> Yes <input type="checkbox"/> No | |
| Were there any changes in patient medical history that are cardiovascular in etiology? | <input type="checkbox"/> Yes <i>If yes, specify: _____</i> <input type="checkbox"/> No | |
| Were there any new changes to existing Concomitant Medications? | <input type="checkbox"/> Yes <i>(If yes, please add new or update Concomitant Medication CRF)</i> <input type="checkbox"/> No | |
| Was visit imaging done? | <input type="checkbox"/> Yes Are required images for this visit available? <input type="checkbox"/> Yes <input type="checkbox"/> No <input type="checkbox"/> No <input type="checkbox"/> N/A Imaging not required per protocol | |

Site Personnel Signature

____ / ____ / ____
Date (DD/MMM/YYYY)

| | |
|-----------------------------------|---|
| Were vital signs performed? | <input type="checkbox"/> Yes <input type="checkbox"/> No |
| Date of vital sign measurements | ___ / ___ / ___ (DD/MMM/YYYY) |
| Height (xxx.xx) | _____ (cm / in) (circle one) |
| Weight (xxx.x) | _____ (kg / lb) (circle one) |
| BMI (xx.x) | _____ (kg/m ²) |
| Systolic Blood Pressure (xxx) | _____ (mmHg) |
| Diastolic Blood Pressure (xxx) | _____ (mmHg) |
| Heart Rate (xxx) | _____ (bpm) |

Site Personnel Signature

___ / ___ / ___
Date (DD/MMM/YYYY)

For use with Follow Up Visits as needed (45-Day, 12 Months, and Unscheduled).

Reminders:

- **At 45 Days and 12 Months:**
- TEE or CT is mandatory per protocol at 45 Days and 12 Months for Implanted Subjects
- If a CT is completed and shows findings (i.e., leak or thrombus), a TEE is required to confirm the finding as soon as possible

| | |
|---|--|
| Was Echocardiogram/CT performed? | <input type="checkbox"/> Yes <input type="checkbox"/> No <i>(Complete protocol deviation form)</i> |
| Are the required images for this visit available? | <input type="checkbox"/> Yes <input type="checkbox"/> No <i>(Complete protocol deviation form)</i> |
| Were images uploaded into the Imaging Module? | <input type="checkbox"/> Yes <input type="checkbox"/> No |
| Time period of Imaging | <input type="checkbox"/> 45 Day <input type="checkbox"/> 12 Months <input type="checkbox"/> Unscheduled, specify: _____ |
| Date echocardiogram/CT completed | ___ / ___ / ___ (DD/MMM/YYYY) |
| Imaging Type | <input type="checkbox"/> TTE – Transthoracic echocardiogram <input type="checkbox"/> TEE – Transesophageal echocardiogram <input type="checkbox"/> Cardiac CT <input type="checkbox"/> Cardiac MRI <input type="checkbox"/> Brain CT <input type="checkbox"/> MRI |
| Left atrial appendage visible | <input type="checkbox"/> Yes <input type="checkbox"/> No <input type="checkbox"/> Not Available |

If available, confirm if the following was noted on echo/CT:

| | | | | |
|---|---|--|--------------------------|--|
| Dense spontaneous echo contrast consistent with Thrombus? | <input type="checkbox"/> Yes <input type="checkbox"/> No <input type="checkbox"/> Not Available | | | |
| Intra-cardiac thrombus | <table border="1"> <tr> <td><input type="checkbox"/> Yes <i>(Complete AE form)</i> <input type="checkbox"/> No <input type="checkbox"/> Not available</td> <td>If yes, confirm location</td> <td> <input type="checkbox"/> Left atrium <input type="checkbox"/> Left atrial appendage <input type="checkbox"/> Left ventricle <input type="checkbox"/> Right atrium <input type="checkbox"/> Right ventricle <input type="checkbox"/> Other, specify: _____ </td> </tr> </table> | <input type="checkbox"/> Yes <i>(Complete AE form)</i> <input type="checkbox"/> No <input type="checkbox"/> Not available | If yes, confirm location | <input type="checkbox"/> Left atrium <input type="checkbox"/> Left atrial appendage <input type="checkbox"/> Left ventricle <input type="checkbox"/> Right atrium <input type="checkbox"/> Right ventricle <input type="checkbox"/> Other, specify: _____ |
| <input type="checkbox"/> Yes <i>(Complete AE form)</i> <input type="checkbox"/> No <input type="checkbox"/> Not available | If yes, confirm location | <input type="checkbox"/> Left atrium <input type="checkbox"/> Left atrial appendage <input type="checkbox"/> Left ventricle <input type="checkbox"/> Right atrium <input type="checkbox"/> Right ventricle <input type="checkbox"/> Other, specify: _____ | | |

| | | | |
|--|--|--|--|
| Intra-cardiac vegetation | <input type="checkbox"/> Yes <input type="checkbox"/> No <input type="checkbox"/> Not available | If yes, confirm location | <input type="checkbox"/> Left atrium <input type="checkbox"/> Left atrial appendage <input type="checkbox"/> Left ventricle <input type="checkbox"/> Right atrium <input type="checkbox"/> Right ventricle <input type="checkbox"/> Other, specify: _____ |
| Patent foramen ovale warranting closure? | <input type="checkbox"/> Yes <input type="checkbox"/> No <input type="checkbox"/> Not available | Is this a high risk PFO? | <input type="checkbox"/> Yes <input type="checkbox"/> No |
| Atrial septal defect? | <input type="checkbox"/> Yes <input type="checkbox"/> No <input type="checkbox"/> Not available | If yes, specify | <input type="checkbox"/> Right to left shunt present <input type="checkbox"/> Left to right shunt present <input type="checkbox"/> Bidirectional shunt <input type="checkbox"/> Unable to determine |
| | | If yes, does defect warrant closure? | <input type="checkbox"/> Yes <input type="checkbox"/> No |
| Left atrial appendage occlusion device position stable and position unchanged? | <input type="checkbox"/> Yes <input type="checkbox"/> No <input type="checkbox"/> Not available | | |
| Peri-device leak present? | <input type="checkbox"/> Yes <input type="checkbox"/> No <input type="checkbox"/> Not available | If yes, specify (mm) | _____ mm |
| Pericardial effusion present? | <input type="checkbox"/> Yes (<i>Assess for AE</i>) <input type="checkbox"/> No <input type="checkbox"/> Not available | | |
| | If yes, select type | <input type="checkbox"/> Circumferential <input type="checkbox"/> Loculated | |
| | If yes, select size <i>*AE is reportable for pericardial effusions Moderate or larger</i> | <input type="checkbox"/> Trivial <input type="checkbox"/> Small (<1 cm) <input type="checkbox"/> Moderate (1-2 cm) <input type="checkbox"/> Large (>2 cm and <5cm) <input type="checkbox"/> Large (> 5 cm) | |
| | If yes, do any of the following apply? | <input type="checkbox"/> Symptomatic <input type="checkbox"/> Sign or symptom of acute or chronic pericarditis <input type="checkbox"/> Evidence of tamponade physiology | |

Device embolization?

- Yes (*Complete AE form*)
 No
 Not available

Site Personnel Signature

____/____/_____
Date (DD/MMM/YYYY)

| | |
|---|--|
| Was the QVSFS assessment completed? | <input type="checkbox"/> Yes <input type="checkbox"/> No <i>(Complete protocol deviation form)</i> |
| Is this assessment performed because of a neurological event? | <input type="checkbox"/> Yes <i>If yes, assess for an adverse event</i> <input type="checkbox"/> No |
| Date of QVSFS assessment | ___ / ___ / ___ (DD/MMM/YYYY) |

| Since the last study contact (by phone or clinic) | Yes | No | Unknown |
|--|--------------------------|--------------------------|--------------------------|
| 1. Were you told by a physician that you had a stroke? | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> |
| 2. Were you ever told by a physician that you had a TIA, ministroke, or a transient ischemic attack? | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> |
| 3. Have you ever had a sudden weakness on one side of your body? | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> |
| 4. Have you ever had a sudden numbness or dead feeling on one side of your body? | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> |
| 5. Have you ever had a sudden painless loss of vision in one or both eyes? | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> |
| 6. Have you ever suddenly lost one half of your vision? | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> |
| 7. Have you ever suddenly lost the ability to understand what people are saying? | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> |
| 8. Have you ever suddenly lost the ability to express yourself verbally or in writing? | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> |

Site Personnel Signature

___ / ___ / ___
Date (DD/MMM/YYYY)

18 MONTH

CONFORM Visit Information

Source Data Transfer Tool

Site Number: _____ Subject ID: _____

| | | |
|--|---|---|
| Visit Timepoint | <input type="checkbox"/> Pre-Discharge <input type="checkbox"/> Day 7 <input type="checkbox"/> Day 45 <input type="checkbox"/> 6 Months <input type="checkbox"/> 12 Months <input type="checkbox"/> 18 Months | <input type="checkbox"/> 2 Year <input type="checkbox"/> 3 Year <input type="checkbox"/> 4 Year <input type="checkbox"/> 5 Year <input type="checkbox"/> Not related to a study visit <input type="checkbox"/> Unscheduled visit |
| Was visit completed? | <input type="checkbox"/> Yes <input type="checkbox"/> No | |
| Visit Date | ____ / ____ / ____ (DD/MMM/YYYY) | |
| Visit Type | <input type="checkbox"/> Office/clinic visit <input type="checkbox"/> Telephone contact <input type="checkbox"/> Video link | |
| Were there any new or changes to existing Adverse Events? <i>If yes, please complete or update an Adverse Event CRF</i> | <input type="checkbox"/> Yes Was the event a suspected stroke or systemic embolism? <input type="checkbox"/> Yes <input type="checkbox"/> No <input type="checkbox"/> No | |
| Did the subject have any ER visits or hospitalizations since the last visit? | <input type="checkbox"/> Yes <input type="checkbox"/> No | |
| Were there any changes in patient medical history that are cardiovascular in etiology? | <input type="checkbox"/> Yes <i>If yes, specify: _____</i> <input type="checkbox"/> No | |
| Were there any new changes to existing Concomitant Medications? | <input type="checkbox"/> Yes <i>(If yes, please add new or update Concomitant Medication CRF)</i> <input type="checkbox"/> No | |
| Was visit imaging done? | <input type="checkbox"/> Yes Are required images for this visit available? <input type="checkbox"/> Yes <input type="checkbox"/> No <input type="checkbox"/> No <input type="checkbox"/> N/A Imaging not required per protocol | |

Site Personnel Signature

____ / ____ / ____
Date (DD/MMM/YYYY)

Was the NIHSS assessment completed?

- Yes
 No (*Complete Protocol Deviation form*)

Date of NIHSS assessment

___ / ___ / ___ (DD/MM/YYYY)

| 1(a) – Level of consciousness | |
|---|------------------------------|
| Alert, keenly responsive | <input type="checkbox"/> (0) |
| Not alert; but arousable by minor stimulation to obey, answer or respond | <input type="checkbox"/> (1) |
| Not alert' requires repeated stimulation to attend, or is obtunded and requires strong or painful stimulation to make movements (not stereotyped) | <input type="checkbox"/> (2) |
| Responds only with reflex motor or autonomic unresponsive, flaccid or areflexic | <input type="checkbox"/> (3) |
| 1(b) – Level of consciousness questions | |
| Answers both questions correctly | <input type="checkbox"/> (0) |
| Answers one question correctly | <input type="checkbox"/> (1) |
| Answers neither question correctly | <input type="checkbox"/> (2) |
| 1(c) – Level of consciousness command | |
| Performs both tasks correctly | <input type="checkbox"/> (0) |
| Performs one task correctly | <input type="checkbox"/> (1) |
| Performs neither task correctly | <input type="checkbox"/> (2) |
| 2 – Best gaze | |
| Normal | <input type="checkbox"/> (0) |
| Partial gaze palsy; gaze is abnormal in one or both eyes, but forced deviation or total gaze paresis is not present | <input type="checkbox"/> (1) |
| Forced deviation, or total gaze paresis not overcome by the oculoccephalic maneuver | <input type="checkbox"/> (2) |
| 3 – Visual | |
| No visual loss | <input type="checkbox"/> (0) |
| Partial hemianopia | <input type="checkbox"/> (1) |
| Complete hemianopia | <input type="checkbox"/> (2) |
| Bilateral hemianopia (blind including cortical blindness) | <input type="checkbox"/> (3) |

| 4 – Facial palsy | |
|---|-------------------------------|
| Normal symmetrical movements | <input type="checkbox"/> (0) |
| Minor paralysis (flattened nasolabial fold, asymmetry on smiling) | <input type="checkbox"/> (1) |
| Partial paralysis (total or near-total paralysis of lower face) | <input type="checkbox"/> (2) |
| Complete paralysis of one or both sides (absence of facial movement in the upper and lower face) | <input type="checkbox"/> (3) |
| 5(a) – Motor arm - left | |
| No drift, limb holds 90 (or 45) degrees for 10 full seconds | <input type="checkbox"/> (0) |
| Drift; limb holds 90 (or 45) degrees, but drifts down before full 10 seconds; does not hit bed or other support | <input type="checkbox"/> (1) |
| Some effort against gravity; limb cannot get to or maintain (if cued) 90 (or 45) degrees; drifts down to bed, but has some effort against gravity | <input type="checkbox"/> (2) |
| No effort against gravity, limb falls | <input type="checkbox"/> (3) |
| No movement | <input type="checkbox"/> (4) |
| Amputation or joint fusion, Explain | <input type="checkbox"/> (UN) |
| 5(b) – Motor arm - right | |
| No drift, limb holds 90 (or 45) degrees for 10 full seconds | <input type="checkbox"/> (0) |
| Drift; limb holds 90 (or 45) degrees, but drifts down before full 10 seconds; does not hit bed or other support | <input type="checkbox"/> (1) |
| Some effort against gravity; limb cannot get to or maintain (if cued) 90 (or 45) degrees; drifts down to bed, but has some effort against gravity | <input type="checkbox"/> (2) |
| No effort against gravity, limb falls | <input type="checkbox"/> (3) |
| No movement | <input type="checkbox"/> (4) |
| Amputation or joint fusion, Explain: | <input type="checkbox"/> (UN) |

| 6(a) Motor leg - left | |
|--|-------------------------------|
| No drift, leg holds 30-degree position for full 5 seconds | <input type="checkbox"/> (0) |
| Drift: leg falls by the end of the 5-second period but does not hit bed | <input type="checkbox"/> (1) |
| Some effort against gravity; leg falls to bed by 5 seconds, but has some effort against gravity | <input type="checkbox"/> (2) |
| No effort against gravity, leg falls to bed immediately | <input type="checkbox"/> (3) |
| No movement | <input type="checkbox"/> (4) |
| Amputation or joint fusion, Explain: | <input type="checkbox"/> (UN) |
| 6(a) Motor leg - right | |
| No drift, leg holds 30-degree position for full 5 seconds | <input type="checkbox"/> (0) |
| Drift: leg falls by the end of the 5-second period but does not hit bed | <input type="checkbox"/> (1) |
| Some effort against gravity; leg falls to bed by 5 seconds, but has some effort against gravity | <input type="checkbox"/> (2) |
| No effort against gravity, leg falls to bed immediately | <input type="checkbox"/> (3) |
| No movement | <input type="checkbox"/> (4) |
| Amputation or joint fusion, Explain: | <input type="checkbox"/> (UN) |
| 7 – Limb ataxia | |
| Absent | <input type="checkbox"/> (0) |
| Present in one limb | <input type="checkbox"/> (1) |
| Present in two limbs | <input type="checkbox"/> (2) |
| Amputation or joint fusion, Explain: | <input type="checkbox"/> (UN) |
| 8 – Sensory | |
| Normal; no sensory loss | <input type="checkbox"/> (0) |
| Mild-to-moderate sensory loss; patient feels pinprick less sharp or dull on the affected side; or there is a loss of superficial pain with pinprick, but patient is aware of being touched | <input type="checkbox"/> (1) |
| Severe to total sensory loss; patient is not aware of being touched in the face, arm, and leg | <input type="checkbox"/> (2) |

| 9 – Best language | |
|---|-------------------------------|
| No aphasia; normal | <input type="checkbox"/> (0) |
| Mild-to-moderate aphasia; some obvious loss of fluency or facility or comprehension, without significant limitation on ideas expressed or form of expression. Reduction of speech and/or comprehension, however, makes conversion about provided materials difficult or impossible. <i>For example, in conversation about provided materials, examiner can identify picture or naming card content from patient's response.</i> | <input type="checkbox"/> (1) |
| Severe aphasia; all communication is through fragmentary expression; great need for inference questioning and guessing by the listener. Range of information that can be exchanged is limited; listener carries burden of communication. Examiner cannot identify materials provided form patient response. | <input type="checkbox"/> (2) |
| Mute, global aphasia; no usable speech or auditory comprehension | <input type="checkbox"/> (3) |
| 10 – Dysarthria | |
| Normal | <input type="checkbox"/> (0) |
| Mild-to-moderate dysarthria; patient slurs at least some word and, at worst, can be understood with some difficult. | <input type="checkbox"/> (1) |
| Severe dysarthria; patient's speech is so slurred as to be unintelligible in the absence or out of proportion to any dysphasia, or is mute/anarthric | <input type="checkbox"/> (2) |
| Intubated or another physical barrier, Explain: | <input type="checkbox"/> (UN) |
| 11 – Extinction and inattention (formerly neglect) | |
| No abnormality | <input type="checkbox"/> (0) |
| Visual, tactile, auditory, spatial, or personal inattention or extinction to bilateral simultaneous stimulation in one of the sensory modalities. | <input type="checkbox"/> (1) |
| Profound hemi-inattention or extinction to more than one modality; does not recognize own hand or orients to only one side of space. | <input type="checkbox"/> (2) |

Site Personnel Signature

___/___/_____
Date (DD/MMM/YYYY)

CONFORM QVSFS

Source Data Transfer Tool

Site Number: _____ Subject ID: _____

| | |
|---|--|
| Was the QVSFS assessment completed? | <input type="checkbox"/> Yes <input type="checkbox"/> No <i>(Complete protocol deviation form)</i> |
| Is this assessment performed because of a neurological event? | <input type="checkbox"/> Yes <i>If yes, assess for an adverse event</i> <input type="checkbox"/> No |
| Date of QVSFS assessment | ___ / ___ / ___ (DD/MMM/YYYY) |

| Since the last study contact (by phone or clinic) | Yes | No | Unknown |
|--|--------------------------|--------------------------|--------------------------|
| 1. Were you told by a physician that you had a stroke? | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> |
| 2. Were you ever told by a physician that you had a TIA, ministroke, or a transient ischemic attack? | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> |
| 3. Have you ever had a sudden weakness on one side of your body? | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> |
| 4. Have you ever had a sudden numbness or dead feeling on one side of your body? | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> |
| 5. Have you ever had a sudden painless loss of vision in one or both eyes? | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> |
| 6. Have you ever suddenly lost one half of your vision? | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> |
| 7. Have you ever suddenly lost the ability to understand what people are saying? | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> |
| 8. Have you ever suddenly lost the ability to express yourself verbally or in writing? | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> |

Site Personnel Signature

___ / ___ / ___
Date (DD/MMM/YYYY)

Was the mRS assessment completed?

- Yes
 No (*Complete protocol deviation form*)

Is this assessment performed because of a neurological event?

- Yes *If yes, assess for an adverse event*
 No

Date of mRS assessment

___ / ___ / ___ (DD/MMM/YYYY)

| Level of Consciousness | Response |
|--|--------------------------|
| 0 = No symptoms at all | <input type="checkbox"/> |
| 1 = No significant disability despite symptoms; able to carry out all usual duties and activities | <input type="checkbox"/> |
| 2 = Slight disability; unable to carry out all previous activities but able to look after own affairs without assistance | <input type="checkbox"/> |
| 3 = Moderate disability; requiring some help, but able to walk without assistance | <input type="checkbox"/> |
| 4 = Moderately severe disability; unable to walk without assistance, and unable to attend to own body needs without assistance | <input type="checkbox"/> |
| 5 = Severe disability; bedridden, incontinent, and requiring constant nursing care and attention | <input type="checkbox"/> |
| Score | |

Site Personnel Signature___ / ___ / ___
Date (DD/MMM/YYYY)

**2 YEAR THROUGH
5 YEAR**

CONFORM Visit Information

Source Data Transfer Tool

Site Number: _____ Subject ID: _____

| | | |
|--|---|---|
| Visit Timepoint | <input type="checkbox"/> Pre-Discharge <input type="checkbox"/> Day 7 <input type="checkbox"/> Day 45 <input type="checkbox"/> 6 Months <input type="checkbox"/> 12 Months <input type="checkbox"/> 18 Months | <input type="checkbox"/> 2 Year <input type="checkbox"/> 3 Year <input type="checkbox"/> 4 Year <input type="checkbox"/> 5 Year <input type="checkbox"/> Not related to a study visit <input type="checkbox"/> Unscheduled visit |
| Was visit completed? | <input type="checkbox"/> Yes <input type="checkbox"/> No | |
| Visit Date | ____ / ____ / ____ (DD/MMM/YYYY) | |
| Visit Type | <input type="checkbox"/> Office/clinic visit <input type="checkbox"/> Telephone contact <input type="checkbox"/> Video link | |
| Were there any new or changes to existing Adverse Events? <i>If yes, please complete or update an Adverse Event CRF</i> | <input type="checkbox"/> Yes Was the event a suspected stroke or systemic embolism? <input type="checkbox"/> Yes <input type="checkbox"/> No <input type="checkbox"/> No | |
| Did the subject have any ER visits or hospitalizations since the last visit? | <input type="checkbox"/> Yes <input type="checkbox"/> No | |
| Were there any changes in patient medical history that are cardiovascular in etiology? | <input type="checkbox"/> Yes <i>If yes, specify: _____</i> <input type="checkbox"/> No | |
| Were there any new changes to existing Concomitant Medications? | <input type="checkbox"/> Yes <i>(If yes, please add new or update Concomitant Medication CRF)</i> <input type="checkbox"/> No | |
| Was visit imaging done? | <input type="checkbox"/> Yes Are required images for this visit available? <input type="checkbox"/> Yes <input type="checkbox"/> No <input type="checkbox"/> No <input type="checkbox"/> N/A Imaging not required per protocol | |

Site Personnel Signature

____ / ____ / ____
Date (DD/MMM/YYYY)

CONFORM QVSFS

Source Data Transfer Tool

Site Number: _____ Subject ID: _____

| | |
|---|--|
| Was the QVSFS assessment completed? | <input type="checkbox"/> Yes <input type="checkbox"/> No <i>(Complete protocol deviation form)</i> |
| Is this assessment performed because of a neurological event? | <input type="checkbox"/> Yes <i>If yes, assess for an adverse event</i> <input type="checkbox"/> No |
| Date of QVSFS assessment | ___ / ___ / ___ (DD/MMM/YYYY) |

| Since the last study contact (by phone or clinic) | Yes | No | Unknown |
|--|--------------------------|--------------------------|--------------------------|
| 1. Were you told by a physician that you had a stroke? | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> |
| 2. Were you ever told by a physician that you had a TIA, ministroke, or a transient ischemic attack? | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> |
| 3. Have you ever had a sudden weakness on one side of your body? | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> |
| 4. Have you ever had a sudden numbness or dead feeling on one side of your body? | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> |
| 5. Have you ever had a sudden painless loss of vision in one or both eyes? | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> |
| 6. Have you ever suddenly lost one half of your vision? | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> |
| 7. Have you ever suddenly lost the ability to understand what people are saying? | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> |
| 8. Have you ever suddenly lost the ability to express yourself verbally or in writing? | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> |

Site Personnel Signature

___ / ___ / ___
Date (DD/MMM/YYYY)

STUDY EXIT

Date of study exit

___ / ___ / ___ (DD/MMM/YYYY)

Screen Failure

- Subject did not meet I/E criteria prior to index procedure (if subject was randomized, please do not select)
- Subject did not meet I/E criteria after the Index Procedure TEE was performed and prior to the Access Sheath crossed the body
- Other Inc/Exc/Screening assessment failure; **Describe below:**

Subject Classification

Withdrawn

- No Implant received at index procedure (after IMPLANT imaging, Access Sheath crossed the body)
- Subject withdrew consent
- Subject lost to follow up
- Investigator decision to withdraw subject
- Site terminated by Sponsor
- Sponsor terminated the Study
- Subject withdrew due to COVID -19 diagnosis
- Subject withdrew due to COVID -19 safety concerns
- Other; **Describe below**

Subject Classification

Subject Death

- Completed Study** – Subject implanted and completed follow up through 5 years

Subject Classification

Subject Classification

___ / ___ / ___

Date (DD/MMM/YYYY)

Site Personnel Signature

