

# California CABG Outcomes Reporting Program Data Abstractor Training Manual

## Version 8.4



## **Document Revision History**

| Date      | Version   | on Description  |  |
|-----------|---|---|--|
| 3/2/2020  | 8.1   | Training Manual Release                                   |  |
| 4/8/2020  | 8.2   | Added COVID-19 Data Element and valid values for Country. |  |
| 5/11/2020 | 8.3   | Removed Patient address data elements.                    |  |
| 6/9/2020  | Updated definitions to be in synch with the STS Training Manual release <a href="https://www.sts.org/sites/default/files/Training%20Manual%20V_20 2%20July%202020.pdf">https://www.sts.org/sites/default/files/Training%20Manual%20V_20 2%20July%202020.pdf</a> |   |  |
| 7/28/2020 | 8.4   | Finalized.  |  |

#### **CCORP**

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Data Elements in Export Order

Effective with July 1, 2020 Discharges

### **Overview: DATA ELEMENT EXPORT ORDER**

| Da  | nta Element   | Classification       | Origin             |
|-----|---|----------------------|--------------------|
| 1.  | Medical Record Number   | Demographics         | STS                |
| 2.  | Type of CABG  | Operative            | Non-STS            |
| 3.  | Date of Surgery   | Hospitalization      | STS                |
| 4.  | Date of Birth   | Demographics         | STS                |
| 5.  | Patient Age   | Demographics         | STS                |
| 6.  | Sex   | Demographics         | STS                |
| 7.  | Primary Payor   | Hospitalization      | STS                |
| 8.  | Secondary (Supplemental) Payor  | Hospitalization      | STS                |
| 9.  | Race Documented   | Demographics         | STS                |
| 10. | Race – White  | Demographics         | STS                |
| 11. | Race – Black/African American   | Demographics         | STS                |
| 12. | Race – Asian  | Demographics         | STS                |
| 13. | Race – American Indian/ Alaskan Native                                    | Demographics         | STS                |
| 14. | Race – Native Hawaiian/ Pacific Islander                                  | Demographics         | STS                |
| 15. | Race - Other  | Demographics         | STS                |
| 16. | Hispanic or Latino or Spanish Ethnicity                                   | Demographics         | STS                |
| 17. | Hospital Discharge Date   | Discharge/ Mortality | STS                |
| 18. | Status at Hospital Discharge  | Discharge/ Mortality | STS                |
| 19. | Patient Transfer to Another Acute Care Hospital                           | Discharge/ Mortality | STS                |
| 20. | Patient Transfer to Acute Care Hospital-<br>Date                          | Discharge/ Mortality | STS                |
| 21. | Mortality Date  | Discharge/ Mortality | STS                |
| 22. | Mort – Status at 30 Days After Surgery (either discharged or in-hospital) | Discharge/ Mortality | STS                |
|     | Responsible Surgeon Name (3 separate fields)                              | Operative            | Non-STS            |
|     | a. Surgeon Last Name  | Operative            | Non-STS            |
|     | o. <u>Surgeon First Name</u><br>c. <u>Surgeon Middle Initial</u>          | Operative Operative  | Non-STS<br>Non-STS |
|     | Responsible Surgeon CA License Number                                     | Operative            | Non-STS            |
| 25. | Height (cm)   | Risk Factors         | STS                |
| 26. | Weight (kg)   | Risk Factors         | STS                |
| 27. | <u>Diabetes</u>   | Risk Factors         | STS                |
| 28. | <u>Diabetes Control</u>   | Risk Factors         | STS                |
| 29. | Dialysis  | Risk Factors         | STS                |

| Data Element                            | Classification                 | Origin |
|---|--------------------------------|--------|
| 30. <u>Hypertension</u>                 | Risk Factors                   | STS    |
| 31. Endocarditis                        | Risk Factors                   | STS    |
| 32. Infectious Endocarditis Type        | Risk Factors                   | STS    |
| 33. <u>Chronic Lung Disease</u>         | Risk Factors                   | STS    |
| 34. <u>Pneumonia</u>                    | Risk Factors                   | STS    |
| 35. <u>Liver Disease</u>                | Risk Factors                   | STS    |
| 36. <u>Immunocompromised Present</u>    | Risk Factors                   | STS    |
| 37. <u>COVID-19</u>                     | Risk Factors                   | STS    |
| 38. <u>Cancer within 5 Years</u>        | Risk Factors                   | STS    |
| 39. <u>Peripheral Artery Disease</u>    | Risk Factors                   | STS    |
| 40. <u>Cerebrovascular Disease</u>      | Risk Factors                   | STS    |
| 41. Prior CVA                           | Risk Factors                   | STS    |
| 42. <u>Prior CVA When</u>               | Risk Factors                   | STS    |
| 43. <u>CVD TIA</u>                      | Risk Factors                   | STS    |
| 44. <u>CVD – Carotid Stenosis</u>       | Risk Factors                   | STS    |
| 45. <u>CVD Carotid Stenosis – Right</u> | Risk Factors                   | STS    |
| 46. <u>CVD Carotid Stenosis – Left</u>  | Risk Factors                   | STS    |
| 47. CVD Prior Carotid Surgery           | Risk Factors                   | STS    |
| 48. <u>Last Creatinine Level</u>        | Risk Factors                   | STS    |
| 49. <u>Total Albumin</u>                | Risk Factors                   | STS    |
| 50. <u>Total Bilirubin</u>              | Risk Factors                   | STS    |
| 51. <u>INR</u>                          | Risk Factors                   | STS    |
| 52. <u>Sodium</u>                       | Risk Factors                   | STS    |
| 53. <u>Previous CABG</u>                | Previous Cardiac Interventions | STS    |
| 54. <u>Previous Valve</u>               | Previous Cardiac Interventions | STS    |
| 55. <u>Previous PCI</u>                 | Previous Cardiac Interventions | STS    |
| 56. <u>Previous PCI – Interval</u>      | Previous Cardiac Interventions | STS    |
| 57. <u>Prior MI</u>                     | Preoperative Cardiac Status    | STS    |
| 58. <u>MI - When</u>                    | Preoperative Cardiac Status    | STS    |
| 59. <u>Heart Failure</u>                | Preoperative Cardiac Status    | STS    |
| 60. <u>Heart Failure Timing</u>         | Preoperative Cardiac Status    | STS    |
| 61. <u>Classification – NYHA</u>        | Preoperative Cardiac Status    | STS    |
| 62. <u>Cardiogenic Shock</u>            | Preoperative Cardiac Status    | STS    |
| 63. <u>Resuscitation</u>                | Preoperative Cardiac Status    | STS    |
| 64. <u>Cardiac Arrhythmia</u>           | Preoperative Cardiac Status    | STS    |

| Data Element   | Classification              | Origin |
|--|-----------------------------|--------|
| 65. <u>Cardiac Arrhythmia – Vtach/VFib</u>                         | Preoperative Cardiac Status | STS    |
| 66. <u>Cardiac Arrhythmia - AFlutter</u>                           | Preoperative Cardiac Status | STS    |
| 67. <u>Cardiac Arrhythmia – Third Degree Heart</u><br><u>Block</u> | Preoperative Cardiac Status | STS    |
| 68. Cardiac Arrhythmia – Atrial Fibrillation                       | Preoperative Cardiac Status | STS    |
| 69. Atrial Fibrillation- Type                                      | Preoperative Cardiac Status | STS    |
| 70. Warfarin Use (within 5 days)                                   | Preoperative Medications    | STS    |
| 71. Coronary Anatomy/Disease Known                                 | Hemodynamics / Cath / Echo  | STS    |
| 72. <u>Number Diseased Vessels</u>                                 | Hemodynamics / Cath / Echo  | STS    |
| 73. Left Main Stenosis>= 50% Known                                 | Hemodynamics / Cath / Echo  | STS    |
| 74. Hemo Data EF Done  | Hemodynamics / Cath / Echo  | STS    |
| 75. Hemo Data EF   | Hemodynamics / Cath / Echo  | STS    |
| 76. PA Systolic Pressure Measured                                  | Hemodynamics / Cath / Echo  | STS    |
| 77. PA Systolic Pressure   | Hemodynamics / Cath / Echo  | STS    |
| 78. Mitral Valve Regurgitation                                     | Hemodynamics / Cath / Echo  | STS    |
| 79. Mitral Regurgitation   | Hemodynamics / Cath / Echo  | STS    |
| 80. <u>Incidence</u>   | Operative                   | STS    |
| 81. Status   | Operative                   | STS    |
| 82. <u>Urgent/ Emergent/ Emergent Salvage</u> Reason               | Operative                   | STS    |
| 83. Perfusion Strategy   | Operative                   | STS    |
| 84. CPB Utilization Combination Plan                               | Operative                   | STS    |
| 85. Internal Mammary Artery Used                                   | Coronary Bypass             | STS    |
| 86. Reason for No IMA  | Coronary Bypass             | STS    |
| 87. <u>Valve</u>   | Operative                   | STS    |
| 88. <u>Aortic Valve</u>  | Operative                   | STS    |
| 89. Aortic Valve Procedure   | Valve Surgery               | STS    |
| 90. Mitral Valve   | Operative                   | STS    |
| 91. Mitral Valve Procedure   | Valve Surgery               | STS    |
| 92. <u>Tricuspid Valve</u>   | Operative                   | STS    |
| 93. <u>Pulmonic Valve</u>  | Operative                   | STS    |
| 94. Reoperation for Bleed/Tamponade                                | Postoperative Events        | STS    |
| 95. <u>Unplanned Coronary Artery Intervention</u>                  | Postoperative Events        | STS    |
| 96. <u>Unplanned Coronary Artery Intervention-Vessel</u>           | Postoperative Events        | STS    |
| 97. Deep Sternal   | Postoperative Events        | STS    |

| Data Element                             | Classification             | Origin  |
|--|----------------------------|---------|
| 98. <u>Neuro - Stroke Permanent</u>      | Postoperative Events       | STS     |
| 99. Pulm – Ventilation Prolonged         | Postoperative Events       | STS     |
| 100. <u>Renal – Renal Failure</u>        | Postoperative Events       | STS     |
| 101. Renal – Dialysis Requireme          | nt Postoperative Events    | STS     |
| 102. <u>Other – A Fib</u>                | Postoperative Events       | STS     |
| 103. <u>Facility Identification Numb</u> | <u>per</u> Hospitalization | Non-STS |

| D  | ata Element                                    | Valid Values  | Definition   |
|----|--|---|--|
| 1. | Medical Record<br>Number<br>STS Sequence #: 85 | Alphanumeric  | Indicate the patient's medical record number at the hospital where surgery occurred. This field should be collected in compliance with state/local privacy laws.   |
| 2. | Type of CABG<br>CCORP-specific<br>variable     | 1 = Isolated<br>3 = CABG + Valve<br>4= Other Non-isolated<br>CABG   | Indicate whether the surgery was considered an isolated CABG, CABG + Valve, or all other CABG.  Other Non-isolated must include a CABG (not isolated valve).  CCORP Clarification/Comments: *See reference on pages 53-55.   |
| 3. | Date of Surgery<br>STS Sequence #: 310         | Numeric: mmddyyyy   | Indicate the date of coronary artery bypass graft procedure  CCORP Clarification/Comments: The date the patient enters the operating room for surgery.   |
| 4. | Date of Birth<br>STS Sequence #: 65            | Numeric: mmddyyyy   | Indicate the patient's date of birth using 4-digit format for year. This field should be collected in compliance with state/local privacy laws.  |
| 5. | Patient Age<br>STS Sequence #: 70              | Numeric   | Indicate the patient's age in years, at time of surgery. This should be calculated from the date of birth and the date of surgery, according to the convention used in the USA (the number of birthdate anniversaries reached by the date of surgery). Do not submit CABG for patients <18 years old.  |
| 6. | Sex<br>STS Sequence #: 75                      | 1 = Male<br>2 = Female  | Indicate the patient's sex at birth as either male or female.  CCORP Clarification/Comments: Patients who have undergone gender reassignment surgery maintain the risk associated with their chromosomal gender. Code gender at birth.   |
| 7. | Primary Payor<br>STS Sequence #: 291           | 1 = None/Self 2 = Medicare (includes commercially managed options) 3 = Medicaid (includes commercially managed options) 9 = Commercial Health Insurance 10 = Health Maintenance Organization 4 = Military 11 = Non-U.S. Plan 13 = Other | Indicate the primary insurance payor at time of arrival.  STS Intent/Clarification: When there is more than one payor, the primary payor pays first. The patient admitted after a car accident may have the primary insurance listed as the auto insurance policy with his health care policy as his secondary insurance. In this scenario, the intent is to capture the patient's normal health care policy, do not capture the auto insurance policy as primary payor.  General Information: Payor Description  None / Self – the patient has no insurance, or the patient is self pay. Code Christian Healthcare Ministries and Medi-Share Christian Health Care in this selection.  Medicare – Includes commercially managed options  Medicare Part A – is hospital insurance and covers inpatient hospital stays, skilled nursing facility, hospice care and some home health care. Some patients may only have Medicare A and this is not included in Fee-for-Service.  Medicare Part B – is medical insurance; payment for Pro-fee or the coverage for physician services (therefore it is coded as Fee-for-Service), outpatient care, medical supplies, and preventive services. |

| Data Element | Valid Values | Definition   |
|--------------|--------------|--|
|              |              | Medicare Part C / Medicare Advantage Plan – is still a Medicare program which is managed by an insurance company, most have additional benefits – vision, and/or dental. Medicare Advantage Plan covers most Medicare benefits and usually require patients to see specific providers in their network.  All Medicare Advantage / Managed Care plans (ie. Humana HMO Medicare) are captured in the payor category as Medicare only. For example, if the patient has Medicare HMO, code as primary payor Medicare, there is no secondary payor in this scenario. Medicare Part D is prescription drug coverage. Medicare Part D is optional, and it's available only through private insurance companies that contract with Medicare (Medicare Advantage or Managed Care plans). Medicare Supplement plans are not part of Medicare – this is a separate private health insurance plan. |
|              |              | Medicare Advantage Plan Types: HMO   |
|              |              | PPO Private Fee-for-Service  |
|              |              | Special needs plan   |
|              |              | Medicare Medical Savings Account plan  |
|              |              | Medicaid - Medicaid [Medi-Cal in California] in the United States is a federal and state program that helps with medical costs for some people with limited income and resources. Medicaid also offers benefits not normally covered by Medicare, including nursing home care and personal care services. All Medicaid Commercial / Managed Care plans (ie. Humana Medicare, Star Molina Medicaid) are captured in the payor category as Medicaid only.  Commercial Health Insurance - Commercial health insurance is health insurance provided and  |
|              |              | administered by non-governmental entities. It covers medical expenses and disability income for the insured. Commercial insurance includes Medicare Supplement plans such as Medigap or AARP etc. It is a private insurance policy that can help pay for some of the health care cost Medicare doesn't cover, such as co-payments, coinsurance, and deductibles. <b>This is not part of Medicare</b> – this is a separate private health insurance plan. Point-of-service plan (POS) and Preferred Provider Organization (PPO) plans not associated with Medicare Advantage plans will be captured here.   |
|              |              | Health Maintenance Organization (HMO) - An HMO gives you access to certain doctors and hospitals within its network. A network is made up of providers that have agreed to lower their rates for plan members and meet quality standards. But unlike PPO plans, care under an HMO plan is covered only if you see a provider within that HMO's network. There are few opportunities to see a non-network   |

| D  | ata Element  | Valid Values  | Definition  |
|----|--|---|---|
|    |  |   | provider. There are also typically more restrictions for coverage than other plans, such as allowing only a certain number of visits, tests or treatments.  Military – US Military provides insurance. Typically reported as VA insurance or Tricare.  Non-U.S. Plan – Insurance covered by a non-U.S. source.  Other – All other insurance not listed in the above selections such as Indian Health Services,  Correctional Facility, State Specific plans, other government insurance, charitable care or foundation funding. |
| 8. | Secondary<br>(Supplemental) Payor<br>STS Sequence #: 298 | 1 = None/Self 2 = Medicare (includes commercially managed options) 3 = Medicaid (includes commercially managed options) 9 = Commercial Health Insurance 10 = Health Maintenance Organization 4 = Military 11 = Non-U.S. Plan 13 = Other | Indicate which if any secondary insurance payor the patient had at time of arrival.  Intent/Clarification: When there is more than one payor, the secondary payor pays after the primary payor.   |
| 9. | Race Documented<br>STS Sequence # 150                    | 1 = Yes;<br>2 = No;<br>3 = Patient Declined to<br>Disclose  | Indicate whether the race is documented. Intent/Clarification: Race should be self-reported by the patient/family. Do not assign race or make assumptions if race is not documented.  Yes No Patient Declined to Disclose- Indicate if the patient declined to provide race or if race was not documented.  |
| 10 | . Race – White<br>STS Sequence #: 151                    | 1 = Yes<br>2 = No   | Indicate whether the patient's race, as determined by the patient or family, includes White. "White" refers to a person having origins in any of the original peoples of Europe, the Middle East, or North Africa. It includes people who indicated their race(s) as "White" or reported entries such as Irish, German, Italian, Lebanese, Arab, Moroccan, or Caucasian.  |

| Data Element  | Valid Values                            | Definition   |
|---|---|--|
| 11. Race – Black/African<br>American<br>STS Sequence #: 151               | 1 = Yes<br>2 = No                       | Indicate whether the patient's race, as determined by the patient or family, includes Black/African-American. "Black or African-American" refers to a person having origins in any of the black racial groups of Africa. It includes people who indicated their race(s) as "Black, African Am., or Negro" or reported entries such as African American, Kenyan, Nigerian, or Haitian.  |
| 12. Race – Asian<br>STS Sequence #: 151                                   | 1 = Yes<br>2 = No                       | Indicate whether the patient's race, as determined by the patient or family, includes Asian. "Asian" refers to a person having origins in any of the original peoples of the Far East, Southeast Asia, or the Indian subcontinent, including, for example, Cambodia, China, India, Japan, Korea, Malaysia, Pakistan, the Philippine Islands, Thailand, and Vietnam. It includes people who indicated their race(s) as "Asian" or reported entries such as "Asian Indian", "Chinese", "Filipino", "Korean", "Japanese", "Vietnamese", and "Other Asian" or provided other detailed Asian responses.   |
| 13. Race – American<br>Indian/ Alaskan<br>Native<br>STS Sequence #:151    | 1 = Yes<br>2 = No                       | Indicate whether the patient's race, as determined by the patient or family, includes American Indian/Alaskan Native. "American Indian or Alaska Native" refers to a person having origins in any of the original peoples of North and South America (including Central America) and who maintains tribal affiliation or community attachment. This category includes people who indicated their race(s) as "American Indian or Alaska Native" or reported their enrolled or principal tribe, such as Navajo, Blackfeet, Inupiat, Yup'ik, or Central American Indian groups or South American Indian groups. This includes all in North American native peoples such as American Indian/Alaskan Native, Inuit. |
| 14. Race – Native<br>Hawaiian/ Pacific<br>Islander<br>STS Sequence #: 151 | 1 = Yes<br>2 = No                       | Indicate whether the patient's race, as determined by the patient or family, includes Native Hawaiian / Pacific Islander. "Native Hawaiian or Other Pacific Islander" refers to a person having origins in any of the original peoples of Hawaii, Guam, Samoa, or other Pacific Islands. It includes people who indicated their race(s) as "Pacific Islander" or reported entries such as "Native Hawaiian", "Guamanian or Chamorro", "Samoan", and "Other Pacific Islander" or provided other detailed Pacific Islander responses.  |
| 15. Race – Other<br>STS Sequence #: 151                                   | 1 = Yes<br>2 = No                       | Indicate whether the patient's race, as determined by the patient or family, includes any other race.  "Some Other Race" includes all other responses not included in the White, Black or African American, American Indian or Alaska Native, Asian, and Native Hawaiian or Other Pacific Islander race categories described above.  |
| 16. Hispanic or Latino or<br>Spanish Ethnicity<br>STS Sequence #: 185     | 1 = Yes<br>2 = No<br>3 = Not Documented | Indicate if the patient is of Hispanic, Latino or Spanish ethnicity as reported by the patient/family.  "Hispanic, Latino or Spanish" refers to a person of Cuban, Mexican, Puerto Rican, South or Central American, or other Spanish culture or origin regardless of race.  CCORP Clarification/Comments: People who identify their origin as Hispanic, Latino or Spanish may be of any race.   |

| Data Element  | Valid Values   | Definition   |
|---|--|--|
| 17. Hospital Discharge Date STS Sequence #: 7006  | Numeric: mmddyyyy  | Indicate the date the patient was discharged from the hospital (acute care facility) If the patient died in the hospital, the hospital discharge date is the date of death.  |
| 18. Status at Hospital Discharge STS Sequence #: 7007   | 3 = Discharged Alive, last<br>known status alive (other<br>than Hospice)<br>4 = Discharged Alive, died<br>after discharge<br>5 = Discharged to Hospice<br>2 = Died in hospital | <ul> <li>Indicate the patient's status at hospital discharge.</li> <li>Intent/Clarification:         <ul> <li>Discharged Alive, last known status alive (other than Hospice) – Includes patients who are discharged alive. Does not include hospice discharge – see choice below for discharged to hospice.</li> <li>Discharged Alive, died after discharge - Includes patients who are discharged alive and expire after discharge.</li> <li>Discharged to Hospice – Includes patients who are discharged to inpatient or outpatient hospice and home hospice.</li> <li>Died in hospital - Includes patients that remain in acute care hospital where the index surgical procedure was performed and die, even if after 30 days following the index surgery. Includes any patient that dies at another acute care hospital after transfer even if after 30 days following the index surgery.</li> </ul> </li> </ul> |
| 19. Patient Transfer to Another Acute Hospital STS Sequence #: 7003                               | 1 = Yes;<br>2 = No   | Indicate if the patient was transferred to another acute care hospital.  |
| 20. Patient Transfer to Acute Care Hospital- Date STS Sequence #:7004                             | Numeric: mmddyyyy  | Indicate the date the patient was transferred to another acute care hospital.  |
| 21. Mortality Date<br>STS Sequence #: 7121  | Numeric: mmddyyyy  | Indicate the date the patient was declared dead.   |
| 22. Mort – Status at 30 Days After Surgery (either discharged or in-hospital) STS Sequence #:7001 | 1 = Alive;<br>2 = Dead<br>3 = Unknown  | Indicate if the patient was alive or dead at 30 days post- surgery (whether in hospital or not).   |

| Data Element  | Valid Values  | Definition   |
|---|---|--|
| 23. Responsible Surgeon<br>Name<br>CCORP-specific<br>variable     | Surgeon Last Name<br>Surgeon First Name<br>Surgeon Middle Initial | Indicate the Surgeon's name.  CCORP Clarification/Comments: Hospitals are encouraged to look up their surgeon names and licensing information DIRECTLY from the California Medical Board. <a href="http://www.mbc.ca.gov/Breeze/License Verification.aspx">http://www.mbc.ca.gov/Breeze/License Verification.aspx</a> and Osteopaths directly from Dept of Consumer Affairs <a href="https://search.dca.ca.gov/">https://search.dca.ca.gov/</a> **See reference on page 55.  |
| 24. Responsible Surgeon CA License Number CCORP-specific variable |   | California physician license number of responsible surgeon assigned by the Medical Board of California of the Department of Consumer affairs.  See page 58 of this training manual for more information criteria.  CCORP Clarification/Comments: Hospitals are encouraged to look up their surgeon names and licensing information DIRECTLY from the California Medical Board. <a href="http://www.mbc.ca.gov/Breeze/License">http://www.mbc.ca.gov/Breeze/License</a> Verification.aspx and Osteopaths directly from Dept of Consumer Affairs <a href="https://search.dca.ca.gov/">https://search.dca.ca.gov/</a> When entering Osteopath license numbers, please include |
| 25. Height (cm)<br>STS Sequence #: 330                            | Usual Range: 122.0 – 213.0<br>Low/High: 20.0 – 251.0              | the leading 20A. Call or email the hotline if you receive errors.  Indicate the height of the patient in centimeters closest to time of OR entry.  CCORP Clarification/Comments: Used to calculate BSA (body surface area), a field for risk calculation.  To convert Inches to centimeters, multiply # of inches by 2.54.  1 inch = 2.54 centimeters.   |
| 26. Weight (kg)<br>STS Sequence #: 335                            | Usual Range: 30.0 – 181.8<br>Low/High: 10.0 – 250.0               | Indicate the weight of the patient in kilograms closest to time of OR entry.  Intent/Clarification: Used to calculate BSA (body surface area) and is a field for risk calculation.  Record in kilograms. 1 Kg = 2.2 pounds. Time frame – capture weight closest to time of OR for index procedure. Use the Anesthesia Record as priority source, followed by the Perfusion record. If weight is not available from the above sources, use the weight recorded in other documents closest to entry to OR for index procedure.   |
| 27. Diabetes<br>STS Sequence #: 360                               | 1 = Yes<br>2 = No<br>3 = Unknown                                  | History of diabetes diagnosed and/or treated by a healthcare provider.  Hemoglobin A1c >=6.5% is indicative of diabetes. Please refer your healthcare providers to the 2017  ADA Standards of Medical Care in Diabetes.  2017 American Diabetes Association Standards of Medicare Care in Diabetes - 2017. Diabetes Care. 40  (Suppl.1):S13. <a href="https://professional.diabetes.org/sites/professional.diabetes.org/files/media/dc_40_s1_final.pdf">https://professional.diabetes.org/sites/professional.diabetes.org/files/media/dc_40_s1_final.pdf</a> .   |

| Data Element                                | Valid Values  | Definition  |
|---|---|---|
|   |   | Intent/Clarification: Indicate if the patient has a history of diabetes mellitus regardless of duration of disease or need for anti-diabetic agents. Code no for patients with steroid induced hyperglycemia and gestational (transient) diabetes if there is no supportive documentation of diabetes such as a HbA1c and/or treatment. Not all patients receiving diabetic medications are considered diabetic. It is important to remember that some medications used to treat diabetes may be used to treat other conditions.  Time frame – capture any occurrence between birth and entry to OR for index procedure.  A HbA1c value > 6.5, collected within 3 months prior to surgery, is acceptable for documentation of diabetes = "yes".  Patients with a history of diabetes who have had a pancreatic transplant are coded as Yes to Diabetes.  Code "Unknown" when there is conflicting information in the medical record and/or with the patient/family and/or patient/family unable to provide history. |
| 28. Diabetes Control<br>STS Sequence #: 365 | 1 = None 2 = Diet only 3 = Oral 4 = Insulin 5 = Other 6 = Other subcutaneous medication 7 = Unknown | Indicate the patient's diabetes control method at home. Choose the most aggressive therapy from the order below  Insulin: insulin treatment (includes any combination with insulin)  Other subcutaneous medications (e.g., GLP-1 agonist)  Oral: treatment with oral agent (includes oral agent with or without diet treatment)  Diet only: Treatment with diet only  None: no treatment for diabetes  Other: other adjunctive treatment, non-oral/insulin/diet  Unknown  2017 American Diabetes Association Standards of Medicare Care in Diabetes - 2017. Diabetes Care. 40 (Suppl.1): S13.  https://professional.diabetes.org/sites/professional.diabetes.org/files/media/dc 40 s1 final.pdf  Intent/Clarification: There must be documentation in chart to code treatment type.  For patients who have had pancreatic transplant, code "other" since the insulin from the new pancreas is not exogenous insulin.  |
| 29. Dialysis<br>STS Sequence #: 375         | 1 = Yes<br>2 = No<br>3 = Unknown  | Indicate whether the patient is currently (prior to surgery) undergoing dialysis on a routine basis.  Intent/Clarification: Includes any form of peritoneal or hemodialysis the patient is currently receiving routinely prior to surgery with the intent to resume post -op. Also, may include Continuous Veno-Venous Hemofiltration (CVVH, CVVH-D), and Continuous Renal Replacement Therapy (CRRT) as dialysis.  |

| Data Element   | Valid Values                     | Definition   |
|--|----------------------------------|--|
|  |                                  | Code "No" for renal dialysis if ultrafiltration is the only documentation found in the record since this is for volume management.  Code "Unknown" when there is conflicting information in the medical record and/or with the patient/family and/or patient/family unable to provide history.   |
| 30. Hypertension<br>STS Sequence #: 380              | 1 = Yes<br>2 = No<br>3 = Unknown | Indicate if the patient has a current diagnosis of hypertension defined by any 1 of the following:  • History of hypertension diagnosed and treated with medication, diet, and/or exercise  • Currently undergoing pharmacological therapy for treatment of hypertension 2017 ACC/AHA/AAPA/ABC/ACPM/AGS/APhA/ASH/ASPC/NMA/PCNA Guideline for the Prevention, Detection, Evaluation, and Management of High Blood Pressure in Adults: A Report of the American College of Cardiology/American Heart Association Task Force on Clinical Practice Guidelines. J Am Coll Cardiol 2018;71:e127-e248.  Intent/Clarification: - Time frame – capture any occurrence between birth and entry to OR for index   |
|  |                                  | procedure. Capturing of HTN as a risk factor must be based on Provider diagnosis of HTN in the medical record.  Code "Unknown" when there is conflicting information in the medical record and/or with the patient/family and/or patient/family unable to provide history.   |
| 31. Endocarditis<br>STS Sequence #: 385              | 1 = Yes<br>2 = No                | Indicate whether the patient has a history of endocarditis. Endocarditis must meet the current CDC definition (See Training manual). Intent/Clarification: Indicate whether the patient has a history of endocarditis documented and diagnosed by a Provider. The below CDC link is provided as a resource. Time frame – capture any occurrence between birth and entry to OR for index procedure. https://www.cdc.gov/nhsn/pdfs/pscmanual/17pscnosinfdef_current.pdf Choose "Yes" for patients with pre-operative endocarditis who begin antibiotics post-op. Code "Yes" for patients who are diagnosed intraoperatively.  Marantic Endocarditis (Nonbacterial Thrombotic Endocarditis) (Lupus) should not be coded as infectious endocarditis. |
| 32. Infectious Endocarditis Type STS Sequence #: 390 | 1 = Treated<br>2 = Active        | Indicate the type of endocarditis the patient has. If the patient is currently being treated for endocarditis, the disease is considered active. If no antibiotic medication (other than prophylactic medication) is being given at the time of surgery and the cultures are negative, then the infection is considered treated.  CCORP Clarification/Comments: If the patient is currently being treated with antimicrobials for endocarditis, the disease is considered active.  |

| Data Element             | Valid Values                             | Definition  |
|--------------------------|--|---|
| 33. Chronic Lung Disease | 1 = No                                   | STS Intent/Clarification: Active - currently being treated; also include patients who were diagnosed in the OR but began treatment postop. Treated - no antibiotic medication at time of surgery (other than prophylactic medication). Indicate whether the patient has chronic lung disease, and the severity level according to the   |
| STS Sequence #: 405      | 2 = Mild                                 | following classification:   |
| ·                        | 3 = Moderate                             | No;   |
|                          | 4 = Severe                               | Mild: FEV1 60% to 75% of predicted, or on chronic inhaled or oral bronchodilator therapy.   |
|                          | 5 = Lung disease<br>documented; severity | Moderate: FEV1 50% to 59% of predicted, or on chronic oral/systemic steroid therapy aimed at lung disease.  |
|                          | unknown                                  | Severe: FEV1 < 50% or Room Air pO2 < 60 or pCO2 > 50.   |
|                          | 6 = Unknown                              | CLD present, severity not documented.   |
|                          |  | Unknown   |
|                          |  | <b>CCORP Clarification/Comments:</b> The diagnosis of chronic lung disease is not based solely on the   |
|                          |  | fact that a person has or currently is smoking or is on home oxygen. Diagnostic testing and/or  |
|                          |  | pharmacological criteria must be met. Chest x-ray findings alone are not included in the data specs for inclusion as chronic lung disease and should not be coded as "Yes"  |
|                          |  | To inclusion as childric lung disease and should not be coded as Tes  |
|                          |  | <b>Time Frame:</b> Do not use values obtained more than 12 months prior to the date of surgery.   |
|                          |  | Intent/Clarification: A history of chronic inhalation reactive disease (asbestosis, mesothelioma, black lung disease or pneumoconiosis) may qualify as chronic lung disease. Radiation induced pneumonitis or radiation fibrosis also qualifies as chronic lung disease. (if above criteria are met) A history of atelectasis is a transient condition and does not qualify. Chronic lung disease can include patients with chronic obstructive pulmonary disease, chronic bronchitis, or emphysema. It can also include a patient who is currently being chronically treated with inhaled or oral pharmacological therapy (e.g., beta-adrenergic agonist, anti-inflammatory agent, leukotriene receptor antagonist, or steroid). |
|                          |  | Spirometry results that have not been interpreted by a pulmonologist may be used to quantify chronic lung disease.  |
|                          |  | Asthma can be considered a chronic lung disease if the patient meets the criteria based on pulmonary function studies, use of inhaled medications or steroids aimed at the lungs.   |

| Data Element                         | Valid Values         | Definition  |
|--------------------------------------|----------------------|---|
|                                      |                      | Sarcoidosis can be considered a chronic lung disease if the patient meets the criteria based on pulmonary function studies, use of inhaled medications or steroids aimed at the lungs. Patients who have had previous lung transplant due to severe CLD no longer have chronic lung disease unless the patient meets the criteria based on pulmonary function studies, use of inhaled medications or steroids aimed at the lungs.  Code "Unknown" when there is conflicting information in the medical record and/or with the patient/family and/or patient/family unable to provide history.  Chronic lung disease, and the severity level is determined by the highlighted above criteria per the data definition for example:  Bedside spirometry results have an FEV1 of 57% but the pulmonologist states the patient has mild chronic lung disease. Code as moderate based on the FEV1 of 57%.  Patient had PFT with FEV1 of 48%. I have not seen any documented diagnosis of chronic lung disease in the chart. Code as severe per FEV1.  Patient had PFT with FEV1 of 78% and a room air ABG with a PCO2 of 36.5 and a PO2 of 56. Code as severe per the room air ABG.  Patient had PFT with FEV1 of 79% and is on Spiriva inhaler and albuterol inhalers at home.  Pulmonary physician cleared patient for surgery documenting the patient had stable COPD and normal PFTs. Code as mild per the use of chronic inhalers at home. |
| 34. Pneumonia<br>STS Sequence #: 465 | 1 = No<br>2 = Recent | Indicate whether patient has a recent (within 30 days) or remote (more than 30 days) history of pneumonia.  |
| 2.2.2.4 # 2.1.00                     | 3 = Remote           | Intent/Clarification: Pneumonia is an infection of one or both lungs caused by bacteria, viruses, fungi,  |
|                                      | 4 = Unknown          | chemicals, or aspiration. It can be community acquired or acquired in a health care setting. Typical  |
|                                      |                      | symptoms associated with pneumonia include cough, chest pain, fever, and difficulty in breathing. Diagnostic tools include x-rays and examination of the sputum. Treatment depends on the cause of pneumonia; bacterial pneumonia is treated with antibiotics.  |

| Data Element                                      | Valid Values                     | Definition   |
|---|----------------------------------|--|
|   |                                  | Code as: Recent- pneumonia diagnosis within 30 days of procedure or Remote - pneumonia diagnosis more than 30 days prior to the procedure. No - meaning no history of pneumonia Unknown - Code "Unknown" when there is conflicting information in the medical record and/or with the patient/family and/or patient/family unable to provide history.  There must be documentation of pneumonia to code "Yes". "Possible pneumonia" with antibiotic treatment should be coded "Unknown". Pneumonitis, inflammation of the lung tissue, without infection is not considered pneumonia and should be coded as "no".   |
| 35. Liver Disease STS Sequence #: 485             | 1 = Yes<br>2 = No<br>3 = Unknown | Indicate whether the patient has a history of hepatitis B, hepatitis C, drug induced hepatitis, auto- immune hepatitis, cirrhosis, portal hypertension, esophageal varices, liver transplant, or congestive hepatopathy. Exclude NASH in the absence of cirrhosis.  Intent/Clarification: LFTs or a MELD score alone cannot be used to code "Yes" to liver disease since other conditions impact these lab values. Liver fibrosis with recurrent ascites, supported by the MELD can be coded as liver disease. Time frame – capture any occurrence between birth and entry to OR for index procedure. The following are not coded as liver disease: Hepatitis A / Hepatitis E Gilberts syndrome Fatty liver Liver Cancer  Hepatic Sarcoidosis should not be coded alone as liver disease. To code liver disease other qualifying disease criteria must be met (cirrhosis, hepatitis, MELD score).  Code "Unknown" when there is conflicting information in the medical record and/or with the patient/family and/or patient/family unable to provide history |
| 36. Immunocompromised Present STS Sequence #: 492 | 1 = Yes<br>2 = No<br>3= Unknown  | Indicate whether immunocompromise is present due to immunosuppressive medication therapy within 30 days preceding the operative procedure or existing medical condition.   |

| Data Element                              | Valid Values  | Definition   |
|---|---|--|
| 37. COVID-19 STS TempCode Sequence#: 7230 | 10 = No<br>11 = Yes, prior to<br>hospitalization for this<br>surgery<br>12 = Yes, in hospital prior to<br>surgery<br>13 = Yes, in hospital after<br>surgery<br>14 = Yes, after discharge<br>within 30 days of surgery | Intent/Clarification: This includes, but is not limited to systemic steroid therapy, anti-rejection medications and chemotherapy. This does not include topical steroid applications, one-time systemic therapy, inhaled steroid therapy or pre-procedure protocol.  Include patients being treated with IVIG, Methotrexate, AntiTNF, Azasan, Imuran, and Hydroxurea.  Patients who have had splenectomy are considered immunocompromised. Examples of conditions causing immunocompromise include Hypogammaglobulinemia, HIV infection, HGB H disease Thalassemia, and patients with systemic lupus taking Plaquenil QD.  Examples of patients who are not considered immunocompromised include:  • Splenic sequestration  • Partial Splenectomy - partial splenectomy may reduce both short and long-term mortality by preserving immune system functioning.  • Patient with IgG4 related sclerosing disease  Code "Unknown" when there is conflicting information in the medical record and/or with the patient/family and/or patient/family unable to provide history. Code unknown if the patient has used immunosuppressive medication therapy, but you do not know if it was within 30 days of surgery.  Did the patient have a laboratory confirmed diagnosis of COVID-19?  No (Harvest Code 10)  Yes, prior to hospitalization for this surgery (Harvest Code 11)  Yes, in hospital prior to surgery (Harvest Code 12)  Yes, in hospital prior to surgery (Harvest Code 13)  Yes, after discharge within 30 days of surgery (Harvest Code 14)  Positive antibody testing is not captured. There are many tests for different types of coronavirus. The one that causes COVID 19 is SARS-CoV-2. Human Coronaviruses types:  229E (alpha coronavirus)  NL63 (alpha coronavirus)  MERS-CoV (the beta coronavirus that causes Middle East Respiratory Syndrome, or MERS)  SARS-CoV-2 (the novel coronavirus that causes coronavirus disease 2019, or COVID19) |

| Data Element  | Valid Values                     | Definition   |
|---|----------------------------------|--|
|   |                                  | Note: During a follow up phone call, a patient says that they tested positive for COVID-19. In this scenario, code Yes, after discharge within 30 days of surgery for patients who self report testing positive for COVID-19 within 30 days of surgery.  Note: For Temporary Code 11 Yes, prior to hospitalization for this surgery. There is no timeframe for Temporary Code 11. Capture any COVID 19 positive test pre-op.  Note: Temporary Code 10 NO applies to any of the above timeframe's pre-op, during hospitalization, and post-op. For example, if the patient tested negative or was not tested pre-op, then code as NO. If the patient is then tested and is negative or not tested during the hospitalization, code NO. If the patient is discharged and is found to be COVID 19 positive within 30 days of surgery, remove code 10 and code Yes to Code 14. |
| 38. Cancer within 5 Years<br>STS Sequence #: 500        | 1 = Yes<br>2 = No<br>3 = Unknown | Indicate whether the patient has a history of cancer diagnosed within 5 years of procedure. Do not capture low grade skin cancers such as basal cell or squamous cell carcinoma.  Intent/Clarification: Capture cancers that have or will require surgical intervention, chemotherapy and or radiation therapy. If the date of diagnosis is not known, then the date of the last treatment may be used to determine the 5-year interval.  Yes - within 5 years  No - patient has never had cancer or has had cancer but not within 5 years  Unknown - Code "Unknown" when there is conflicting information in the medical record and/or with the patient/family and/or patient/family unable to provide history. Code unknown if patient has a history of cancer but you do not know if the cancer occurred within 5 years.  |
| 39. Peripheral Artery<br>Disease<br>STS Sequence #: 505 | 1 = Yes<br>2 = No<br>3 = Unknown | Indicate whether the patient has a history of peripheral arterial disease (includes upper and lower extremity, renal, mesenteric, and abdominal aortic systems). This can include:  1. Claudication, either with exertion or at rest,  2. Amputation for arterial vascular insufficiency,  3. Vascular reconstruction, vascular bypass surgery, or percutaneous intervention to the extremities (excluding dialysis fistulas and vein stripping),  4. Documented abdominal aortic aneurysm with or without repair.  5. Documented subclavian artery stenosis.  Peripheral arterial disease excludes disease in the carotid, cerebrovascular arteries or thoracic aorta.  PVD does not include DVT, pulmonary artery aneurysm, Raynaud's Disease or AVM.  |

| Data Element                                    | Valid Values                     | Definition  |
|---|----------------------------------|---|
|   |                                  | Intent/Clarification: PAD (Peripheral Arterial Disease) is sometimes called PVD (Peripheral Vascular Disease), which can include disease of either peripheral vein or peripheral artery. Code only arterial disease, not venous disease.  Time frame – capture any occurrence between birth and entry to OR for index procedure.  Documentation from a healthcare provider is required to code this field. If documentation is vague or further clarification is needed to abstract correctly then a data manager may use diagnostic results to confirm abstraction. Positive testing, invasive or non-invasive, showing a >50% diameter stenosis in any peripheral artery is a possible indicator of PVD. You must have supporting documentation from the healthcare provider.  If there is conflicting information, then please clarify with patients HCP. For example, a CT Angiogram shows 80% stenosis in the iliac artery but there is no documentation from the patient's healthcare provider to clarify that this is PAD. Clarification and documentation from the Provider is needed to abstract PAD.  Code "Unknown" when there is conflicting information in the medical record and/or with the patient/family and/or patient/family unable to provide history |
| 40. Cerebrovascular Disease STS Sequence #: 525 | 1 = Yes<br>2 = No<br>3 = Unknown | Indicate whether the patient has a current or previous history of any of the following:  A. Stroke: Stroke is an acute episode of focal or global neurological dysfunction caused by brain, spinal cord, or retinal vascular injury as a result of hemorrhage or infarction, where the neurological dysfunction lasts for greater than 24 hours.  B. TIA: is defined as a transient episode of focal neurological dysfunction caused by brain, spinal cord, or retinal ischemia, without acute infarction, where the neurological dysfunction resolves within 24 hours.  C. Noninvasive or invasive arterial imaging test demonstrating >=50% stenosis of any of the major extracranial or intracranial vessels to the brain  D. Vertebral artery and internal carotid and intercranial consistent with atherosclerotic disease with document presence as CVD. External carotid disease is excluded.  E. Previous cervical or cerebral artery revascularization surgery or percutaneous intervention  F. Brain/cerebral aneurysm.  G. Occlusion of vertebral artery, internal carotid artery, and intercranial due to dissection.   |

| Data Element                              | Valid Values                                  | Definition  |
|---|---|---|
|   |   | This does not include chronic (nonvascular) neurological diseases or other acute neurological insults such as metabolic and anoxic ischemic encephalopathy. Subdural hematoma or AVM is not cerebral vascular disease.  Intent/Clarification: Internal carotid and common carotid disease are captured. External carotid disease is not captured. A positive CT scan, even in the patient with no symptoms, should be coded as cerebral vascular disease. A CT scan following surgery with evidence of old infarct or chronic should be coded yes.  Code "Unknown" when there is conflicting information in the medical record and/or with the  |
| 41. Prior CVA                             | 1 = Yes                                       | patient/family and/or patient/family unable to provide history.  Indicate whether the patient has a history of stroke. Stroke is an acute episode of focal or global  |
| STS Sequence #: 530                       | 2 = No<br>3 = Unknown                         | neurological dysfunction caused by brain, spinal cord, or retinal vascular injury as a result of hemorrhage or infarction, where the neurological dysfunction lasts for greater than 24 hours  Intent/Clarification: Include any confirmed neurological deficit of abrupt onset caused by a disturbance in cerebral blood supply that did not resolve within 24 hours of the event. The physical deficit can be in the form of extremity weakness, facial asymmetry, language (speech and/or cognitive thinking) impairment. The intent is to differentiate between neurological events that resolve within 24 hours and those that don't. Time frame — capture any occurrence between birth and entry to OR for index procedure  Code "yes" to prior CVA if the patient has no history of stroke and no symptoms but imaging study results show an infarct (old/chronic or new) or cerebral septic emboli. |
|   |   | Not all subarachnoid hemorrhages will create a stroke. There must be some form of deficit (symptoms lasting > 24 hr) documented in the chart to code SAH as a CVA.  Unknown should be selected if any neurologic dysfunction occurred or was suspected, did not resolve in 24 hours, and could not be confirmed or when there is conflicting information in the medical record and/or with the patient/family and/or patient/family unable to provide history.  |
| 42. Prior CVA When<br>STS Sequence #: 535 | 3 = Recent <= 30 days<br>4 = Remote > 30 days | Indicate when the CVA events occurred. Those events occurring within 30 days prior to the surgical procedure are considered recent, while all others are considered remote.   |

| Data Element   | Valid Values                      | Definition  |
|--|-----------------------------------|---|
| 43. CVD TIA<br>STS Sequence #: 540                   | 1 = Yes<br>2 = No<br>3 = Unknown  | Indicate whether the patient has a history of a Transient Ischemic Attack (TIA). TIA is defined as a transient episode of focal neurological dysfunction caused by brain, spinal cord, or retinal ischemia, without acute infarction, where the neurological dysfunction resolves within 24 hours.  Intent/Clarification: Time frame – capture any occurrence between birth and entry to OR for index |
|  |                                   | procedure.  |
| 44. CVD – Carotid<br>Stenosis<br>STS Sequence #: 545 | 1 = None<br>2 = Right<br>3 = Left | Indicate which carotid artery was determined from any diagnostic test to be >= 50% stenotic.  Intent/Clarification: Code what is found on the study closest to entry into OR for index procedure even   |
|  | 4 = Both<br>5 = Not Documented    | if a prior stent / CEA is in place. Internal carotid and common carotid disease are captured. External carotid disease is not captured.   |
|  |                                   | If the results are reported in a range, such as "40-50%", choose the highest level in the range.  If dissection occluded the artery, then code as 100%. A dissection because the blood flow is null acts like an occlusion.  When a carotid duplex reports stenosis as 0-59% but states "no evidence of hemodynamic significance,"  |
|  |                                   | code No for CVD given the range of 0-59% and documentation of no hemodynamic significance.  |
| 45. CVD Carotid Stenosis                             | 1 = 80-99%                        | Indicate the severity of stenosis reported on the right carotid artery.   |
| – Right  | 2 = 100%                          |   |
| STS Sequence #: 550                                  | 3 = 50-79%                        | Intent/Clarification: Indicate % stenosis:  |
|  | 4 = Not Documented                | 50 - 79% or "moderate"  |
|  |                                   | 80 - 99% or "critical", "severe", or "subtotal". 100% or "total" or "occluded"  |
|  |                                   | Not documented  |
|  |                                   | The accumented  |
|  |                                   | If the results are reported in a range, such as "40-50%", choose the highest level in the range.  |
| 46. CVD Carotid Stenosis                             | 1 = 80-99%                        | Indicate the severity of stenosis reported on the left carotid artery.  |
| – Left   | 2 = 100%                          |   |
| STS Sequence #: 555                                  | 3 = 50-79%                        | Intent/Clarification: Indicate % stenosis:  |
|  | 4 = Not Documented                | 50 - 79% or "moderate"  |
|  |                                   | 80 - 99% or "critical", "severe", or "subtotal".  |
|  |                                   | 100% or "total" or "occluded"   |
|  |                                   | Not documented  |
|  |                                   | If the results are reported in a range, such as "40-50%", choose the highest level in the range.  |

| Data Element  | Valid Values  | Definition   |
|---|---|--|
| 47. CVD Prior Carotid<br>Surgery<br>STS Sequence #: 560 | 1 = Yes<br>2 = No   | Intent/Clarification: Time frame – capture any occurrence between birth and entry to OR for index procedure.  Carotid endarterectomy is a surgical procedure during which a surgeon removes atherosclerotic plaque or other material obstructing the flow of blood from the artery. This procedure eliminates a substance called plaque from the artery and can restore blood flow. Carotid artery stenting is a procedure in which a slender, metal-mesh tube, called a stent, is inserted and expands inside the carotid artery to increase blood flow in areas blocked by plaque.   |
|   | H I D 0.10 12.00  | Also includes internal carotid artery aneurysm coils and a history of carotid angioplasty.   |
| 48. Last Creatinine Level<br>STS Sequence #: 605        | Usual Range: 0.10 – 12.00<br>Low/ High: 0.10 – 30.00          | Indicate the creatinine level closest to the date and time prior surgery but prior to anesthetic management (induction area or operating room). A creatinine level should be collected on all patients, even if they have no prior history of renal disease. A creatinine value is a high predictor of a patient's outcome and is used in the predicted risk models.  Intent/Clarification: Creatinine (Cr) is a chemical waste molecule excreted by the kidneys that is generated from muscle metabolism. If the kidneys become impaired for any reason, the creatinine level in the blood will rise due to poor clearance by the kidneys. Abnormally high levels of creatinine thus warn of possible malfunction or failure of the kidneys. The unit of measurement for Creatinine is mg/dl or mg/100ml or mg%   |
| 49. Total Albumin<br>STS Sequence #: 585                | Usual range: 3.50 - 5.00<br>Low/High: 1.00 - 10.00<br>(mg/dL) | Indicate the total albumin closest to the date and time prior to surgery but prior to anesthetic management (induction area or operating room).  Intent/Clarification: Albumin (alb), produced only in the liver, is the major plasma protein that circulates in the bloodstream. Albumin is essential for maintaining the oncotic pressure in the vascular system. A decrease in oncotic pressure due to a low albumin level allows fluid to leak out from the interstitial spaces into the peritoneal cavity, producing ascites. Albumin is also especially important in the transportation of many substances such as drugs, lipids, hormones, and toxins that are bound to albumin in the bloodstream. A low serum albumin indicates poor liver function. Decreased serum albumin levels are not seen in acute liver failure because it takes several weeks of impaired albumin production before the serum albumin level drops. The most common reason for a low albumin is chronic liver failure caused by cirrhosis. The serum albumin concentration is usually normal in chronic |

| Data Element                               | Valid Values  | Definition   |
|--|---|--|
|  |   | liver disease until cirrhosis and significant liver damage has occurred. The unit of measurement for Albumin is g/dl or g/100 ml or g%.  Timeframe - Capture results up to 6 weeks prior to surgery provided there is no known acute liver disease process.  |
| 50. Total Bilirubin<br>STS Sequence #: 610 | Usual range: 0.20 - 1.30<br>Low/High: 0.10 - 50.00<br>(mg/dL) | Indicate the total Bilirubin closest to the date and time prior to surgery but prior to anesthetic management (induction area or operating room).  |
|  | (9/ 02)   | <b>Intent/Clarification:</b> Bilirubin (Tbili) testing checks for levels of bilirubin, an orange-yellow pigment, in blood. Bilirubin is a natural byproduct that results from the normal breakdown of red blood cells. As a normal process, bilirubin is carried in the blood and passes through the liver. Too much bilirubin may indicate liver damage or disease.   |
|  |   | The unit of measurement for Bilirubin is mg/dl or mg/100 ml or mg%.  |
|  |   | Timeframe - Capture results up to 6 weeks prior to surgery provided there is no known acute liver disease process  |
| 51. INR<br>STS Sequence #: 615             | Usual range 0.90 - 1.30<br>Low/High: 0.50 - 30.00             | Indicate the International Normalized Ratio (INR) closest to the date and time prior to surgery but prior to anesthetic management (induction area or operating room).   |
|  |   | Intent/Clarification: INR is the standard unit used to report the result of a prothrombin (PT) test. An individual whose blood clots normally and who is not on anticoagulation should have an INR of approximately 1. The higher the INR, the longer it takes blood to clot. As the INR increases above a given level, the risk of bleeding and bleeding-related events increases. As the INR decreases below a given level, the risk of clotting events increases. |
| 52. Sodium<br>STS Sequence #: 600          | Usual range 130.0 – 145.0<br>Low/High: 30.0 - 200.00          | Indicate the Sodium level closest to the date and time prior to surgery but prior to anesthetic management (induction area or operating room).   |
| 53. Previous CABG STS Sequence #: 670      | 1 = Yes<br>2 = No   | Indicate whether the patient had a previous Coronary Bypass Graft prior to the current admission.  Intent/Clarification: This applies only to surgical approach to revascularization. Angioplasty or   |
|  |   | other catheter based coronary artery occlusion treatment does not apply.   |
| 54. Previous Valve<br>STS Sequence #: 675  | 1 = Yes<br>2 = No   | Indicate whether the patient had a previous surgical replacement and/or surgical repair of a cardiac valve. This may also include percutaneous valve procedures or transcatheter valve procedures.   |

| Data Element  | Valid Values  | Definition   |
|---|---|--|
|   |   | Intent/Clarification: This may include percutaneous valve procedures such as percutaneous valvotomy or valvuloplasty, as well as surgical or transcatheter valve repair or replacement. Capture all procedures that apply.   |
| 55. Previous PCI<br>STS Sequence #: 775               | 1 = Yes<br>2 = No   | Indicate whether a previous Percutaneous Coronary Intervention (PCI) was performed any time prior to this surgical procedure. PCI is the placement of an angioplasty guide wire, balloon, or other device (e.g. stent, atherectomy, brachytherapy, or thrombectomy catheter) into a native coronary artery or coronary artery bypass graft for the purpose of mechanical coronary revascularization.  Intent/Clarification: An attempted, even if unsuccessful, PCI should be coded as a Previous CV intervention-PCI. This is to harmonize with ACC-NCDR. |
| 56. Previous PCI –<br>Interval<br>STS Sequence #: 800 | 1 = ≤ 6 Hours<br>2 = > 6 Hours  | Indicate the interval of time between the most recent PCI procedure and the current surgical procedure.  Intent/Clarification: The choices are ≤ 6 hours or > 6 hours prior to OR entry. The timing of surgery after PCI may influence outcomes such as renal failure due to contrast given during PCI.  |
| 57. Prior MI<br>STS Sequence #: 885                   | 1 = Yes<br>2 = No<br>3 = Unknown  | Indicate if the patient has had at least one documented previous myocardial infarction at any time prior to this surgery.  Intent/Clarification: Indicate if the patient has a history of MI. Provider documentation should indicate MI. Do not code slight troponin increase and no EKG changes alone as MI without confirmation in the medical record by a physician or physician extender. Do not use phrases such as "cannot rule out", "suggestive", "probable", "cannot exclude", etc. to code MI.   |
| 58. MI - When<br>STS Sequence #: 890                  | 1 = ≤ 6 Hrs<br>2 = > 6 Hrs but < 24 Hrs<br>3 = 1 to 7 Days<br>4 = 8 to 21 Days<br>5 = > 21 Days | Indicate the time period between the last documented myocardial infarction and surgery.  Intent/Clarification: Time of surgery is documented as the hour the patient entered the operating room. Select the time-interval category based on information available on when the MI occurred. MI occurrence is the time of diagnosis and/or when confirmation of the last MI is documented prior to surgery.  |
| 59. Heart Failure<br>STS Sequence #: 911              | 1 = Yes<br>2 = No<br>3 = Unknown  | Indicate whether there is physician documentation or report that the patient has a history of heart failure. Capture either right or left heart failure.   |

| Data Element                                     | Valid Values   | Definition  |
|--|--|---|
|  |  | Intent/Clarification: Heart failure is described as unusual dyspnea on light exertion, recurrent dyspnea occurring in the supine position, fluid retention; or the description of rales, jugular venous distension, pulmonary edema on physical exam, or pulmonary edema on chest x-ray presumed to be cardiac dysfunction. A low ejection fraction alone, without clinical evidence of heart failure does not qualify as heart failure. An elevated BNP without other supporting documentation should not be coded as CHF. Time frame – capture any occurrence between birth and entry to OR for index procedure. NYHA Class documentation alone cannot be used for diagnosis for heart failure, you must have physician documentation that states heart failure. There needs to be documentation in the chart that the patient has been in or was in a state of heart failure.  Do not code heart failure for a diagnosis of Cardiomyopathy. A diagnosis of heart failure must be documented in the medical record to code heart failure. Cardiomyopathy may or may not be associated with a heart failure diagnosis.  Code "Unknown" when there is conflicting information in the medical record and/or with the patient/family and/or patient/family unable to provide history. |
| 60. Heart Failure Timing<br>STS Sequence #: 912  | 1 = Acute<br>2 = Chronic<br>3 =Both                                    | Indicate whether heart failure is acute, chronic or both (acute on chronic).  Acute is new onset/ worsening heart failure within 2 weeks prior to this procedure.  Chronic is greater than 2 weeks prior to this procedure.  Both are worsening heart failure within 2 weeks in a patient with a known history of heart failure.  |
| 61. Classification – NYHA<br>STS Sequence #: 915 | 1 = Class I 2 = Class II 3 = Class III 4 = Class IV 5 = Not Documented | Indicate the patient's worst dyspnea or functional class, coded as the New York Heart Association (NYHA) classification documented by a MD/Provider within the past 2 weeks.  Select the highest level of heart failure within the two weeks leading up to episode of hospitalization or at the time of the procedure. The intent is to capture the highest level of failure. Physician documentation should be in the medical record.  Class I: Patient has cardiac disease but without resulting limitations of ordinary physical activity. Ordinary physical activity (e.g., walking several blocks or climbing stairs) does not cause undue fatigue, palpitation, or dyspnea. Limiting symptoms may occur with marked exertion.   |

| Data Element          | Valid Values                         | Definition  |
|-----------------------|--------------------------------------|---|
|                       |                                      | Class II: Patient has cardiac disease resulting in slight limitation of ordinary physical activity. Patient is comfortable at rest. Ordinary physical activity such as walking more than two blocks or climbing more than one flight of stairs results in limiting symptoms (e.g., fatigue, palpitation, or dyspnea, or anginal pain) |
|                       |                                      | Class III: Patient has cardiac disease resulting in marked limitation of physical activity. Patient is comfortable at rest. Less than ordinary physical activity (e.g., walking one to two level blocks or climbing one flight of stairs) causes fatigue, palpitation, dyspnea, or anginal pain).                                     |
|                       |                                      | <b>Class IV</b> : Patient has dyspnea at rest that increases with any physical activity. Patient has cardiac disease resulting in inability to perform any physical activity without discomfort. Symptoms may be present even at rest. If any physical activity is undertaken, discomfort is increased.                               |
| 62. Cardiogenic Shock | 2 = No                               | Indicate if the patient developed cardiogenic shock. Cardiogenic shock is defined as a sustained (>30   |
| STS Sequence #: 930   | 3 = Yes, at the time of              | min) episode of hypoperfusion evidenced by systolic blood pressure <90 mm Hg and/or, if available,  |
|                       | procedure                            | cardiac index <2.2 L/min per square meter determined to be secondary to cardiac dysfunction and/or  |
|                       | 4 = Yes, not at the time of          | the requirement for parenteral inotropic or vasopressor agents or mechanical support (e.g., IABP,   |
|                       | procedure, but within prior 24 hours | extracorporeal circulation, VADs) to maintain blood pressure and cardiac index above those specified levels.  |
|                       |                                      | Note: Transient episodes of hypotension reversed with IV fluid or atropine do not constitute cardiogenic shock. The hemodynamic compromise (with or without extraordinary supportive therapy) must persist for at least 30 min.  ACCF/AHA 2013  |
|                       |                                      | At the time of the procedure is defined as entry into OR for index procedure and prior to induction of anesthesia. Do not code cardiogenic shock after induction.   |
|                       |                                      | This includes patients with cardiogenic shock who have been stabilized on IABP/inotropes at the time of the procedure.  |
|                       |                                      | • Do not code yes to cardiogenic shock for patients with a low cardiac index who are asymptomatic and do not require mechanical or inotropic support.   |
|                       |                                      | • Hemodynamic issues that could be contributed to anesthesia induction problems should not count in the preoperative status of the patient.   |
|                       |                                      | Elective procedures should not be coded as cardiogenic shock.   |

| Data Element | Valid Values | Definition   |
|--------------|--------------|--|
|              |              | • Do not code yes to cardiogenic shock just because the patient has a LVAD; the patient must meet the blood pressure and/or cardiac index parameters of the definition of cardiogenic shock.   |
|              |              | CCORP Clarification/Comments: "Shock" = Yes if the patient:  |
|              |              | <ol> <li>currently SBP &lt;90 mmHg or cardiac index &lt;2.2 or</li> <li>previously had a SBP &lt; 90 or CI &lt;2.2 but now are on inotropes/ IABP to maintain higher #s.</li> </ol>  |
|              |              | NOTE: sustained (>30 min) episode  |
|              |              | STS: "or requirement for vasopressor agentsto maintain blood pressure and cardiac index"   |
|              |              | Patients left on inotropes/pressors/IABP whose BP/CI has improved so that it is probable BP/CI would be above criteria off therapy should be coded "No." This is more often the case the longer the patient has received these therapies prior to surgery. |
|              |              | 1) CI < 2.2 or unassisted/unaugmented SBP < 90 → shock   |
|              |              | 2) CI>=2.8 or unassisted/unaugmented SBP >=130 → not shock   |
|              |              | <ul> <li>3) CI 2.2-2.39, unassisted/unaugmented SBP 90-99 on <u>any</u> active inotrope/vasopressor/IABP or impella</li> <li>→ shock</li> </ul>  |
|              |              | 4) CI 2.4-2.79, unassisted/unaugmented SBP 100-129 on high dose inotrope/vasopressor/  |
|              |              | impella → shock  |
|              |              | 5) CI 2.4-2.79, unassisted/unaugmented SBP 100-129 on <u>low dose</u> inotrope/ vasopressor/  → not shock  |
|              |              | High Dose Inotropes/Vasopressor dosage High dosage is greater than dosages below:  |
|              |              | a. Dopamine < 5 mcg/kg/min   |
|              |              | b. Dobutamine < 5 mcg/kg/min   |
|              |              | c. Milrinone < 0.375 mcg/kg/min  |
|              |              | d. Norepinephrine (Levophed) < 0.3 mcg/kg/min  |
|              |              | e. Epinephrine < 0.3 mcg/kg/min  |
|              |              | f. Phenylephrine < 0.5 mcg/kg/min g. Vasopressin < 0.03 units per min  |
|              |              | g. Vasopressin < 0.03 units per min  6) VAD, ECMO → shock  |
|              |              | 7) Chart label "shock," inotrope/pressor/IABP, but no CI/BP criteria → not shock   |

| Data Element | Valid Values | Definition   |
|--------------|--------------|--|
|              |              | IABPs are often used to treat coronary ischemia in absence of shock and their use alone does not meet shock criteria (eg, IABP put in for severe left main disease and ACS to stabilize ischemia while waiting for surgery). Some patients have mild cardiogenic shock which does not meet STS criteria even if treated with IABP, inotropes, or pressors. Inotropes may be used or continued to augment diuresis in patients not meeting shock criteria. Note IABPs usually lower systolic BP (assisted SBP < unassisted SBP) therefore assisted SBP should not be used as evidence for shock.  Note: "At the time of the procedure" is defined as prior to induction. This includes patients with CS who have been stabilized on IABP/inotropes at the time of surgery |
|              |              | See following diagram.  140  Diostolia Augmentation  |
|              |              | Diastolic Augmentation  Coronary Perfusion  Unassisted 120 Systole  Mm Hg 100  Balloon Inflation  Unassisted Aortic End-Diastolic Pressure  60   |

| Data Element                          | Valid Values   | Definition   |
|---------------------------------------|--|--|
| 63. Resuscitation STS Sequence #: 935 | 2 = No 3 = Yes, within 1 hour of start of the procedure 4 = Yes, > 1 hour before, but < 24 hours of the start of the procedure | Indicate whether the patient required cardiopulmonary resuscitation before induction of anesthesia. Capture resuscitation timeframe: within 1 hour or 1-24 hours pre-op.  Intent/Clarification: Indicate whether the patient required cardiopulmonary resuscitation within 24 hours of the start of the operative procedure. The start of the procedure begins with the induction of anesthesia. Capture resuscitation timeframe: within 1 hour of surgery or 1-24 hours pre-operatively.  • Resuscitation may include complete circulatory support such as ECMO/other mechanical assist devices (ex. Impella 5.0/LP, LVAD) initiated emergently prior to surgery. Intra-aortic balloon counterpulsation (IABP) by itself does not qualify as complete circulatory support.  • Do not code yes for resuscitation started after induction of anesthesia. The goal is to identify patients who require CPR and/or mechanical circulatory support to maintain life in the 24-hour period preceding surgery.  ECMO: ECMO is to be captured as a status of 'Salvage' and as 'Resuscitation – Yes'. ECMO is a supportive modality and not a procedural type. The risk of the patient on ECMO is accounted for when 'Status = salvage'.  CCORP Clarification/Comments: Impella (other than Impella 5.0/LP) is NOT complete circulatory support and does not qualify as ongoing resuscitation (trumps STS). Impella 5.0/LP is complete circulatory support and qualifies as ongoing resuscitation. |
| 64. Cardiac Arrhythmia                | 1 = Yes  | Indicate whether the patient has a history of a cardiac rhythm disturbance prior to the induction of   |
| STS Sequence #: 945                   | 2 = No   | anesthesia.  |
| 65. Cardiac Arrhythmia –              | 1 = None   | Indicate whether arrhythmia was VTach or VFib.   |
| Vtach/VFib<br>STS Sequence #: 950     | 2 = Remote (> 30 days)<br>3 = Recent (≤ 30 days)   | Intent/Clarification: V-tach rhythm must be sustained/persistent or paroxysmal and require some type of intervention (pharmacological and/or electrical shock) to interrupt and cease the arrhythmia. Do not include short runs of VT.  None  Remote - more than 30 days prior to procedure  Recent - within 30 days of this procedure   |
| 66. Cardiac Arrhythmia –              | 1 = None   | Indicate whether arrhythmia was atrial flutter.  |
| Aflutter                              | 2 = Remote (> 30 days)   |  |
| STS Sequence #: 960                   | 3 = Recent (≤ 30 days)   | Intent/Clarification: Atrial flutter (AFL) is an abnormal heart rhythm that occurs in the atria of the heart. When it first occurs, it is usually associated with a fast heart rate or tachycardia (beats over 100 per minute) which falls into the category of supra-ventricular tachycardias. While this rhythm occurs   |

| Data Element   | Valid Values   | Definition   |
|--|--|--|
|  |  | most often in individuals with cardiovascular disease (e.g. hypertension, coronary artery disease, and cardiomyopathy) and diabetes, it may occur spontaneously in people with otherwise normal hearts. It is typically not a stable rhythm, and frequently degenerates into atrial fibrillation (AF). However, it does rarely persist for months to years. If rhythm is described as fib/flutter, code fibrillation.  None  Remote - more than 30 days prior to procedure  Recent - within 30 days of this procedure  |
| 67. Cardiac Arrhythmia –<br>Third Degree Heart<br>Block<br>STS Sequence #: 970 | 1 = None<br>2 = Remote (> 30 days)<br>3 = Recent (≤ 30 days) | Indicate whether arrhythmia was third degree heart block.  Intent/Clarification: Heart block is applicable only if the patient has or did have 3rd degree heart block (complete heart block). Complete heart block, also referred to as third-degree heart block, or third-degree atrioventricular (AV) block, is a disorder of the cardiac conduction system where there is no conduction through the AV node. Therefore, complete dissociation of the atrial and ventricular activity exists.  |
| 68. Cardiac Arrhythmia –<br>Atrial Fibrillation<br>STS Sequence #: 961         | 1 = None<br>2 = Remote (> 30 days)<br>3 = Recent (<=30 days) | Indicate whether arrhythmia was atrial fibrillation.   |
| 69. Atrial Fibrillation-<br>Type<br>STS Sequence # 971                         | 2 = Paroxysmal<br>4 = Persistent                             | Indicate whether arrhythmia was atrial fibrillation and if so, which type.  Intent/Clarification: If the diagnosis of atrial fibrillation is present code the type: Paroxysmal: Paroxysmal AF is defined as AF that terminates spontaneously or with intervention within seven days of onset. Episodes may recur with variable frequency. Persistent: - Persistent AF is defined as AF that fails to self-terminate within seven days. Episodes often require pharmacologic or electrical cardioversion to restore sinus rhythm. Included in this category are Persistent Afib, Early Persistent Afib, Long-Standing Persistent Afib, Permanent Afib, and Chronic Afib.  Data Source: Heart Rhythm. 2017;S1547 |

| Data Element   | Valid Values                                | Definition   |
|--|---|--|
| 70. Warfarin Use (within<br>5 days)                              | 1 = Yes<br>2 = No                           | Indicate whether the patient received Warfarin (Coumadin) within 5 days preceding surgery.   |
| STS Sequence # 1091  | 3 = Unknown                                 | Intent/Clarification: This is collected to capture the risk of bleeding related to anticoagulation therapy.  |
|  |   | Yes - Capture those who took Coumadin within 5 days preceding surgery.   |
|  |   | <b>No</b> – Patient did not receive Coumadin within 5 days prior to OR entry date and time.  |
|  |   | <b>Unknown</b> – Conflicting information in the medical record and/or with the patient/family or no information is available   |
| 71. Coronary<br>Anatomy/Disease<br>Known<br>STS Sequence #: 1155 | 1 = Yes<br>2 = No                           | Indicate whether coronary artery anatomy and/or disease is documented and available prior to surgery.  Intent/Clarification: Indicate if coronary artery anatomy and/or disease is documented or confirmed by testing prior to surgery. Code Yes for patients who have no coronary artery disease if this has been confirmed by testing.  Sometimes the results are known and verbally communicated to the surgeon, but the Cath Lab Report is not documented in the medical record until after surgery has started; this is particularly true for emergent cases. This can be captured even if dictation was not completed until after the surgery.   |
| 72. Number of Diseased<br>Vessels<br>STS Sequence #: 1170        | 1 = None<br>2 = One<br>3 = Two<br>4 = Three | Indicate the number of diseased major native coronary vessel systems. A vessel that has ever been considered diseased, should always be considered diseased  Intent/Clarification: Indicate if coronary artery anatomy and/or disease is documented or confirmed by testing prior to surgery. Code Yes for patients who have no coronary artery disease if this has been confirmed by testing.  Sometimes the results are known and verbally communicated to the surgeon, but the Cath Lab Report is not documented in the medical record until after surgery has started; this is particularly true for emergent cases. This can be captured even if dictation was not completed until after the surgery. Results dictated following the procedure may be used. |

| Data Element | Valid Values | Definition   |
|--------------|--------------|--|
|              |              | Right coronary artery  Left coronary artery (right oblique)  Proximal circumflex Proximal LAD  Atrial branch 1st septal perforator Obuse marginal  Acute marginal  Distal RCA  Posterior Gescence  (A)  Posterior lateral  Pos |

| Data Element              | Valid Values | Definition  |
|---------------------------|--------------|---|
|                           |              | Coronary artery trifurcation  Ramus  Picture courtesy: Clinical Cardiac CT: Anatomy and Function  |
|                           |              | By Halpern, Ethan J. Halpern Published by Thieme, 2008  |
| 73. Left Main Stenosis >= | 1 = Yes      | Indicate if main stenosis greater or equal to 50% is known.   |
| 50% Known                 | 2 = No       |   |
| STS Sequence #: 1174      | 3 = N/A      | Stenosis at the ostia of the LAD and circumflex is not considered left main disease for the purpose   |
|                           |              | of Society of Thoracic Surgeons (STS). <b>Stenosis needs to be in the left main artery.</b>   |
| 74. Hemo Data EF Done     | 1 = Yes      | Indicate whether the Ejection Fraction was measured prior induction of anesthesia.  |
| STS Sequence #: 1540      | 2 = No       |   |
|                           |              | Intent/Clarification: Some patients may not have had an LV Gram performed during cardiac  |
|                           |              | catheterization due to existing clinical conditions. Ejection fraction (EF) and hemodynamic pressures   |
|                           |              | may be obtained from other sources other than coronary angiogram, such as echo, or MUGA.  |
|                           |              | Because anesthesia can alter the values to be collected, do not collect data from intra-operative   |
|                           |              | transesophageal echo (TEE) after the induction of anesthesia, unless you have no other source to collect the information.   |
|                           |              | Note: If the patient has an echo and a Cath done on the same day and you are not able to determine which study was performed closest to surgery, use the EF from the LHC. |

| Data Element   | Valid Values                                       | Definition   |
|--|--|--|
| 75. Hemo Data EF<br>STS Sequence #: 1545                                     | Usual Range: 5.0 – 90.0<br>Low/ High: 1.0 – 99.0   | Indicate the percentage of the blood emptied from the left ventricle at the end of the contraction).  • Hyperdynamic: >70%  • Normal: 50%–70% (midpoint 60%)  • Mild dysfunction: 40%–49% (midpoint 45%)  • Moderate dysfunction: 30%–39% (midpoint 35%)  • Severe dysfunction: <30%  Note: If no diagnostic report is in the medical record, a value documented in the medical record is acceptable. ACCF/AHA 2013  |
| 76. PA Systolic Pressure   | 1 = Yes  | Indicate whether the PA systolic pressure was measure.   |
| Measured STS Sequence #: 1570  77. PA Systolic Pressure STS Sequence #: 1575 | Usual Range: 15.0 – 40.0<br>Low/High: 10.0 – 150.0 | Intent/Clarification: Elevated pulmonary artery pressures are indicative of pulmonary hypertension, mitral valve disease and other pulmonary/cardiac diseases. Normal systolic pulmonary artery pressure readings are between 15-30 mm of pressure.  If there are no PA pressures recorded or available pre-op from heart cath or echo —one may use PA pressure values from Swan Ganz Catheter inserted for surgery prior to induction of anesthesia.  If PA systolic pressure is not available, it is acceptable to code the peak RV systolic pressure (RVSP).  RVSP and PA systolic pressures will be the same if there is no pulmonary valve disease or outflow obstruction. Note a RVSP cannot be obtained if there is no tricuspid regurgitation present.  Capture PA systolic pressure recorded.  Intent/Clarification: For a PA pressure that is documented as a range value, for example, "30-35 mmHg", capture the highest value in the range |
| 78. Mitral Valve Regurgitation STS Sequence #: 1679                          | 1 = Yes;<br>2 = No                                 | Indicate whether there is evidence of Mitral valve insufficiency/regurgitation prior to surgery.  Intent/Clarification: Mitral regurgitation/insufficiency may be an acute or chronic condition manifesting itself as increased left heart filling pressures which increase the left ventricular stroke volume (amount of blood ejected from the Left Vent. with each heartbeat). Over time, and depending upon the severity, MR can result in pulmonary edema and systemic volume overload. In chronic MR, Left Ventricular Hypertrophy may result. Mitral prolapse and rheumatic fever are the most common cause of MR.  |

| Data Element             | Valid Values             | Definition   |
|--------------------------|--------------------------|--|
| 79. Mitral Regurgitation | 1 = Trivial/Trace;       | Indicate whether there is evidence of Mitral valve insufficiency/regurgitation.  |
| STS Sequence # 1680      | 2 = Mild;                |  |
|                          | 3 = Moderate;            | Intent/Clarification: Indicate the degree of mitral valve insufficiency/regurgitation. Code the highest  |
|                          | 4 = Severe;              | level of valve dysfunction for example mild – moderate will be coded as moderate   |
|                          | 5 = Not documented       |  |
| 80. Incidence            | 1 = First cardiovascular | Indicate if this is the patient's:   |
| STS Sequence #: 1970     | surgery                  |  |
|                          | 2 = First re-op          | -First surgery   |
|                          | cardiovascular surgery   | -First re-op surgery   |
|                          | 3 = Second re-op         | -Second re-op surgery  |
|                          | cardiovascular surgery   | -Third re-op surgery   |
|                          | 4 = Third re-op          | -Fourth or more re-op surgery  |
|                          | cardiovascular surgery   |  |
|                          | 5 = Fourth or more re-op | Intent/Clarification: Incidence will be defined by the number of times of entry into the space for a   |
|                          | cardiovascular surgery   | specific procedure. Please see this resource for an overview of anatomical spaces:   |
|                          |                          | https://en.wikipedia.org/wiki/Pleural_cavity#/media/File:Body_Cavities_Frontal_view_labeled.jpg  |
|                          |                          | A CABG/AVR/ MVR would be in pericardial space. A root/ascending/arch would be the pericardial  |
|                          |                          | space. An open distal arch/descending would be the pleural space. An open thoracoabdominal would   |
|                          |                          | involve the pleural and abdominal space. See examples below:   |
|                          |                          | Previous descending with a current AVR/Root/CAB. This would be first incidence into the pericardial  |
|                          |                          | space.   |
|                          |                          | Previous AVR/Root/hemiarch with a current CAB. This would be first re-op into the pericardial space.   |
|                          |                          | Previous CAB/AVR with a current open descending. This would be first incidence for pleural space   |
|                          |                          | , and the second of the second |
|                          |                          | For the purposes of this field surgery is defined as cardiothoracic surgical procedures performed on the   |
|                          |                          | heart, great vessels or major pericardial procedures, with or without cardiopulmonary bypass (CPB).  |
|                          |                          | Ascending aortic and arch procedures also qualify. Similarly, catheter-based procedures such as TAVR,  |
|                          |                          | TEVAR, mitral-clip, are endovascular procedures and are not classified as prior surgery. Also include lung   |
|                          |                          | procedures utilizing CPB or tracheal procedures utilizing CPB. Reoperation increases risk due to   |
|                          |                          | presence of scar tissue or adhesions.  |
|                          |                          | Great vessels are vessels that are directly attached to the heart - Superior vena cava; Inferior vena cava;  |
|                          |                          | Pulmonary arteries; Pulmonary veins; Aorta.  |

| Data Element | Valid Values | Definition  |
|--------------|--------------|---|
|              |              | The intent of this field is to capture the incidence of the procedure that the patient is about to go through during the current hospitalization, as compared to those procedures prior to this hospitalization. First operative means the patient has never had any surgical procedure on the heart and/or great vessels. Note: previous surgical intervention increases risk for morbidity and mortality and severity of disease process. Choosing N/A does not automatically exclude the patient from analysis. Other Examples of Incidence: |
|              |              | o Patient has a history of a CABG, then later a VAD, then a heart transplant. The patient is now having a CABG on his transplanted heart. Code incidence as third reoperation since the pericardial space has been entered 3 times.   |
|              |              | o Patient underwent a percutaneous aortic valvuloplasty at age 12. He now enters the OR for a surgical AVR. Code incidence as first cardiovascular surgery since the first procedure was percutaneously and this will be the first time the pericardial space is entered.   |
|              |              | o Prior TAVR case that needs a redo-AVR should be coded as first reoperation.   |
|              |              | o Prior TAVR case that needs another cardiac procedure such as a CAB or MVR should be code as incidence as first CV surgery.  |
|              |              | o Prior Mitral Clip Procedure that needs an MVR should be coded as the first operation.   |
|              |              | o Prior TMVR case that needs a redo-MVR should be coded as first reoperation.   |
|              |              | o Prior TMVR case that needs another cardiac procedure such as a CAB or AVR should be code as incidence as first CV surgery.  |
|              |              | CCORP Clarification/Comments:  -CV surgeries INCLUDE: CABG, valve replacement/repair, intracardiac repairs (ASD, VSD), ventricular aneurysmectomy, or surgery on the aortic arch. Use of CPB is not required.  -CV surgeries DO NOT INCLUDE: PCI's and non-cardiac vascular surgeries such as abdominal aortic aneurism repairs or fem-pop bypasses, percutaneous aortic stent grafts, percutaneous valves or pacemaker/ICD implantations.  |

| Data Element                    | Valid Values  | Definition   |
|---------------------------------|---|--|
|                                 |   | The intent of this field is to capture the incidence of the procedure that the patient is about to go through during the current hospitalization, as compared to those procedures prior to this hospitalization.   |
| 81. Status STS Sequence #: 1975 | 1 = Elective 2 = Urgent 3 = Emergent 4 = Emergent Salvage | Indicate the clinical status of the patient prior to entering the operating room.  Elective- The patient's cardiac function has been stable in the days or weeks prior to the operation. The procedure could be deferred without increased risk of compromised cardiac outcome.  Urgent- Procedure required during same hospitalization in order to minimize chance of further clinical deterioration. Examples include but are not limited to: Worsening, sudden chest pain, CHF, acute myocardial infarction (AMI), anatomy, IABP, unstable angina (USA) with intravenous (IV) nitroglycerin (NTG) or rest angina. Any of the conditions that require that the patient remain in the hospital until surgery can take place, but the patient is able to wait for surgery until the next available OR schedule time. Delay in the operation may be necessitated by attempts to improve the patient's condition, availability of a spouse or parent for informed consent, availability of blood products, or the availability of results of essential laboratory procedures or tests. For example, if a patient is brought in for an elective cardiac cath and kept in the hospital for surgery, select urgent.  Emergent- Patients requiring emergency operations will have ongoing, refractory (difficult, complicated, and/or unmanageable) unrelenting cardiac compromise, with or without hemodynamic instability, and not responsive to any form of therapy except cardiac surgery. An emergency operation is one in which there should be no delay in providing operative intervention. Patients requiring emergency operations will have ongoing, refractory (difficult, complicated, and/or unmanageable) cardiac compromise, with or without hemodynamic instability, and not responsive to any form of therapy except cardiac surgery. Hemodynamic picture of shock that is being chemically or mechanically supported. (IV inotrope or IABP to maintain cardiac output [CO]. Requires intubation and ventilation for pulmonary edema. The patient is extending an MI and requires immediate surgery. The patient co |

| Data Element | Valid Values | Definition   |
|--------------|--------------|--|
| Data Element | Valid Values | If a patient presents with a scenario that does not fit into a definite category; it is reasonable to code the reason that most closely matches the patient's presentation.  Emergent Savage- The patient is undergoing CPR en route to the OR prior to anesthesia induction or has ongoing ECMO to maintain life.  CCORP Clarification/Comments: Status refers to the patient's condition immediately before surgery; it should not reflect instability which occurs after the induction of anesthesia or the operative risk but rather how expediently surgery must be performed. Thus, some elective patients may be at higher risk than urgent patients; for example, an elderly patient with an ejection fraction of 20% and COPD operated on electively compared to a young patient with a normal ejection fraction that has ongoing unstable angina.  RULE OF THUMB: Elective – waits at home. Urgent – waits in hospital. Emergent – cannot wait or is not safe to wait. Emergent Salvage – no pulse.  -Elective surgeries are performed on patients whose cardiac function has been stable. They are usually scheduled at least one day prior to surgery, and the clinical picture allows discharge from the hospital with readmission for surgery later.  -Urgent surgeries are performed on patients whose medical condition requires continuous hospitalization prior to CABG. A critical feature that distinguishes urgent from elective patients is that |
|              |              |  |

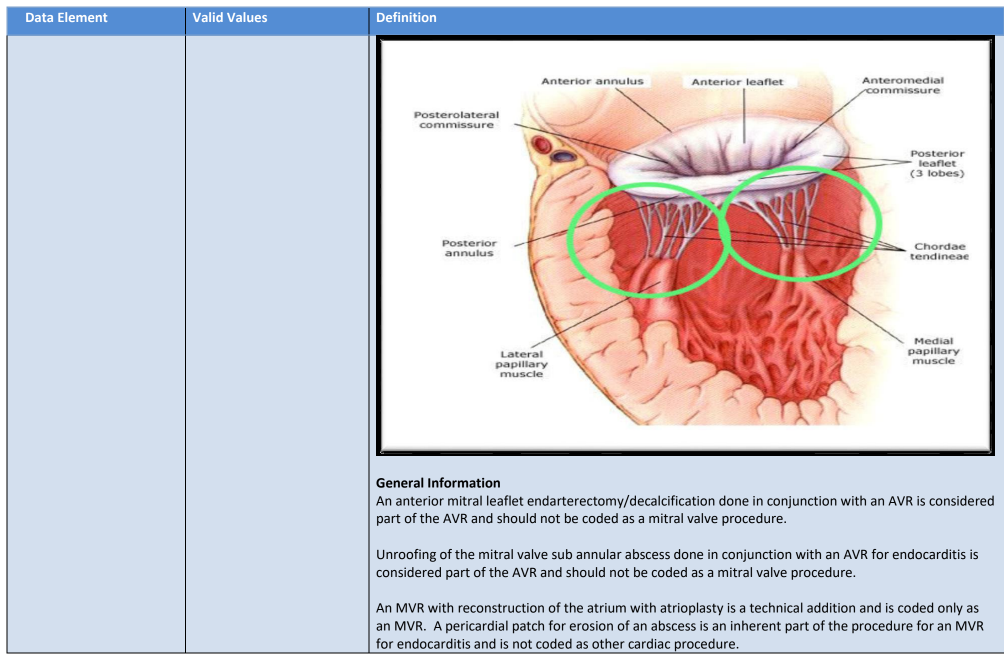
| Data Element          | Valid Values                | Definition   |
|-----------------------|-----------------------------|--|
| 82. Urgent / Emergent | 1 = AMI                     | Choose one reason from the list below that best describes why this operation was considered urgent         |
| /Emergent Salvage     | 2 = Anatomy                 | emergent, or emergent/salvage.   |
| Reason                | 3 = Aortic Aneurysm         |  |
| STS Sequence #: 1990  | 4 = Aortic Dissection       | Intent/Clarification: See list for options. There is no hierarchy - choose the primary reason the          |
|                       | 5 = CHF                     | <b>procedure is urgent or emergent.</b> There may be multiple reasons, choose one that best describes this |
|                       | 6 = Device Failure          | patient's clinical state. The reason is patient specific and needs to be taken from the surgeon's notes.   |
|                       | 7=Diagnostic/Interventional | For example:   |
|                       | Procedure Complication      |  |
|                       | 8 = Endocarditis            | Unstable angina at the time of admission would be coded unstable angina at the time of surgery.            |
|                       | 10 = IABP                   |  |
|                       | 11 = Infected Device        | If a patient has severe aortic and mitral valve stenosis, but also has symptoms such as dyspnea on         |
|                       | 12 = Intracardiac mass or   | exertion (DOE), paroxysmal nocturnal dyspnea (PND), congestion on x-ray or pedal edema that has            |
|                       | thrombus                    | been treated as CHF, code "CHF" as the most appropriate choice.  |
|                       | 13 = Ongoing Ischemia       |  |
|                       | 14 = PCI Incomplete without | Valve dysfunction is defined as a structural failure with that valve. For prosthetic valves – fractured    |
|                       | clinical deterioration      | leaflet, thrombus formation, pannus development which impedes flow through the valve orifice, or           |
|                       | 15 = PCI or attempted PCI   | valvular dehiscence (coming loose or disconnected at the suture line). Native valve dysfunction includes   |
|                       | with Clinical Deterioration | papillary rupture or torn leaflet. Rupture or dissection during cardiac cath; Perforation, tamponade       |
|                       | 16 = Pulmonary Edema        | following cardiac cath-does not include stent closure.   |
|                       | 17 = Pulmonary Embolus      |  |
|                       | 18 = Rest Angina            |  |
|                       | 19 = Shock Circulatory      |  |
|                       | Support                     |  |
|                       | 20 = Shock No Circulatory   |  |
|                       | Support                     |  |
|                       | 21 = Syncope                |  |
|                       | 22 = Transplant             |  |
|                       | 23 = Trauma                 |  |
|                       | 24 = USA                    |  |
|                       | 25 = Valve Dysfunction      |  |
|                       | 26 = Worsening CP           |  |
|                       | 27 = Other                  |  |

| Data Element                                   | Valid Values  | Definition   |
|--|---|--|
| 83. Perfusion Strategy<br>STS Sequence #: 2325 | 28 = Failed Transcatheter Valve Therapy- Acute Annular Disruption 29 = Failed Transcatheter Valve Therapy- Acute Device Malposition 30 = Failed Transcatheter Valve Therapy - Subacute Device Dysfunction 1 = None 2 = Combination 3 = Full 4 = Left Heart Bypass | Indicate the level of CPB or coronary perfusion used during the procedure.  Intent/Clarification:  None: No CPB (cardiopulmonary bypass) or coronary perfusion used during the procedure.  Left Heart Bypass: Left heart bypass is utilized to remove oxygenated blood from the left atrium and return it to the distal descending aorta or femoral artery. This procedure allows repair or replacement of the descending thoracic aorta while regulating blood flow, minimizing surface area contact activation, and reducing heparin requirements.  Combination: With or without CPB and/or with or without coronary perfusion at any time during the procedure (capture conversions from off-pump to on-pump only):  At start of procedure: No CPB/No Coronary Perfusion -> conversion to -> CPB  At start of procedure: No CPB/No Coronary Perfusion -> conversion to -> Coronary perfusion -> conversion to -> CPB/No CPB/No Coronary Perfusion -> conversion to -> CORONARY PERFUSIO |
| 84. CPB Utilization –                          | 1 = Planned   | Full CPB or coronary perfusion was used for the entire procedure  Indicate whether the combination procedure from off-pump to on-pump was a planned or an  |
| Combination Plan STS Sequence #: 2330          | 2 = Unplanned   | unplanned conversion.  |
|  |   | Intent/Clarification: To capture if the operation was intended to be an off-pump case and, for some clinical reason, required cardiopulmonary bypass to complete the operation.  |
|  |   | -Planned: The surgeon intended to treat with any of the combination options described in "CPB utilization"Unplanned: The surgeon did not intend to treat with any of the combination options described in "CPB utilization".   |

| Data Element  | Valid Values  | Definition   |
|---|---|--|
| 85. Internal Mammary<br>Artery Used<br>STS Sequence #: 2626 | 1 = Yes<br>2 = No   | Indicate whether an internal mammary artery conduit was used.  Intent/Clarification: To capture the use of an internal mammary artery (also known as the internal thoracic artery) to construct one or more distal anastomoses: LIMA, RIMA, both or none. IMA may be used as a free or in-situ graft; pedicle, skeletonized.  The patient must leave the OR with an IMA graft in place, in order to code Yes to IMA used. For example, the flow via the IMA graft was poor so the surgeon removed the IMA graft and used a venous graft to the LAD. In this scenario, the IMA was not used.  |
| 86. Reason for No IMA<br>STS Sequence #: 2629               | 2 = Subclavian stenosis 3 = Previous cardiac or thoracic surgery 4 = Previous mediastinal radiation 5 = Emergent or salvage procedure 6 = No (bypassable) LAD disease 7 = Other Not Acceptable STS Provided Exclusion 8 = Other-Acceptable STS Provided Exclusion | Indicate PRIMARY reason Internal Mammary artery was not used as documented in medical record.  Other - acceptable STS provided exclusion – This is an exclusion that has been adjudicated by Surgeon Leadership of STS and deemed to be acceptable. If you have a documented reason for not using the IIMA that does not fall into one of the above approved reasons and it not addressed below, please send in a question to the FAQ Mailbox Ask an Abstraction Question . https://www.sts.org/sts-clinical-question-request-form It is important for sites to keep a copy of the FAQ email documenting the exclusion in the event of an audit.  Examples of Acceptable Reasons:  The IMA was not harvested because the patient had a left upper extremity fistula for hemodialysis and the surgeon was concerned about coronary steal syndrome. Code this as 'Other – acceptable STS provided exclusion'  The patient has a history of severe PVD. The aortogram at the time of the cardiac catheterization showed an occluded distal aorta. The lower extremities perfusion was supplied by the mammary arteries. It was felt that the lower extremities would be in jeopardy if the mammary is used. Code this as an exclusion due to Other - acceptable STS provided exclusion.  A patient had a bilateral mastectomy and immediate reconstruction with flap. During that procedure both the RIMA and LIMA were harvested. 7 months later the patient required a CABG and no IMA could be used as they were not present. Code this as an exclusion due to previous thoracic surgery.  The patient has pectus carinatum and it is documented "We attempted to harvest the left internal mammary artery however due to the patient's severe pectus, it was impossible to visualize the mammary artery. Code this as an exclusion due to Other - acceptable STS provided exclusion.  Examples of Unacceptable Reasons: |

| Data Element                  | Valid Values   | Definition  |
|-------------------------------|--|---|
| 87. Valve                     | 1 = Yes  | I have had multiple cases where the IMA is not used but the physician places an SVG to the LAD. The cardiac cath shows no LAD disease but does have LM disease. The surgeon documents "I elected not to use IMA since the patient does not have any specific LAD disease". No LAD disease is not an acceptable exclusion in this situation. Left main is functionally 2 VD LAD and CX disease. Code OTHER not acceptable STS exclusion as reason for no IMA.  The patient has severe COPD and emphysema with hyperinflated lungs. He had a CABx1 with MVR. The surgeon chooses to use a vein instead of an IMA. This was his documentation in his consult note: "Has single-vessel coronary artery disease and I believe that he should have a vein graft to his LAD as given his large lungs a LIMA would be too hazardous. It simply will not reach." Code OTHER not acceptable STS exclusion as reason for no IMA.  IMA is not used related to poor or low blood flow. Code OTHER not acceptable STS exclusion as reason for no IMA.  Indicate whether a surgical procedure was done on the Aortic, Mitral, Tricuspid or Pulmonic valves |
| STS Sequence #: 2129          | 2 = No   | indicate whether a surgical procedure was done on the Aortic, whitral, incuspid or Full nonic valves  |
| 88. Aortic Valve              | 2 = No   | Indicate whether an aortic valve procedure was performed.   |
| STS Sequence #: 2131          | 3 = Yes, planned<br>4 = Yes, unplanned due to<br>surgical complication<br>5 = Yes, unplanned due to<br>unsuspected disease or<br>anatomy |   |
| 89. Aortic Valve<br>Procedure | 1 = Replacement<br>2 = Repair/Reconstruction   | Indicate the type of procedure that was performed on the aortic valve.  |
| STS Sequence #: 3395          | 3 = Surgical Prosthetic Valve Intervention (not explant of valve)  | Anterior mitral leaflet endarterectomy/decalcification is considered part of the AVR and should not be coded as a mitral valve procedure.  An aortic endarterectomy is considered part of the AVR procedure and should not be coded elsewhere.  Aortoplasty done in conjunction with AVR to reduce the size of the ascending aorta is considered part of the closure and is not coded as an additional procedure.   |
|                               |  | Aortic resection to merely remove excessive aortic tissue prior to aortoplasty is considered part of the closure and is not coded as an additional procedure.   |

| Data Element  | Valid Values   | Definition  |
|---|--|---|
|   |  | Wrapping the dilated portion of the aorta to reinforce it does not constitute an "other or aorta" procedure when done in conjunction with an AVR.  Note: Patient had an AVR for endocarditis. The surgeon also performed unroofing of the mitral valve sub annular abscess. Don't code the unroofing of the mitral valve sub annular abscess. This is part of the AVR for endocarditis. |
| 90. Mitral Valve<br>STS Sequence #: 2133              | 2 = No 3 = Yes, planned 4 = Yes, unplanned due to surgical complication 5 = Yes, unplanned due to unsuspected disease or anatomy | Indicate whether a mitral valve procedure was performed.  |
| 91. Mitral Valve<br>Procedure<br>STS Sequence #: 3500 | 1 = Repair<br>2 = Replacement<br>3 = Surgical Prosthetic Valve<br>Intervention (Not explant of<br>valve)                         | Indicate the type of procedure that was performed on the mitral valve.  |



| Data Element  | Valid Values   | Definition   |
|---|--|--|
| 92. Tricuspid Valve<br>STS Sequence #: 2134               | 2 = No 3 = Yes, planned 4 = Yes, unplanned due to surgical complication 5 = Yes, unplanned due to unsuspected disease or anatomy | Indicate whether a tricuspid valve procedure was performed.  |
| 93. Pulmonic Valve<br>STS Sequence #: 2135                | 2 = No 3 = Yes, planned 4 = Yes, unplanned due to surgical complication 5 = Yes, unplanned due to unsuspected disease or anatomy | Indicate whether a pulmonic valve procedure was performed.   |
| 94. Reoperation for Bleed/ Tamponade STS Sequence #: 6755 | 1 = Yes<br>2 = No  | Indicate whether the patient was re-explored for mediastinal bleeding with or without tamponade either in the ICU or returned to the operating room.  Intent/Clarification: The intent of this field is to capture patients who are re-explored for bleeding / suspected bleeding. Include patients that require surgical re-intervention to investigate or correct bleeding with or without tamponade.  Tamponade occurs when there is compression or restriction placed on the heart within the chest that creates hemodynamic instability or a hypo-perfusion state. Do not include medically (non-operatively) treated excessive post-operative bleeding/tamponade events.  Do not capture reopening of the chest or situations of excessive bleeding that occur prior to the patient leaving the operating room at the time of the primary procedure.  Include patients that return to an OR suite or equivalent OR environment (i.e., ICU setting) as identified by your institution, that require surgical re-intervention to investigate or correct bleeding with or without tamponade. Include only those interventions that pertain to the mediastinum or thoracic cavity. |

| Data Element  | Valid Values  | Definition  |
|---|---|---|
|   |   | Note: Pt returns to OR for exploration of bleed without tamponade. The surgeon documents hematoma evacuation with washout no active bleeding. This is a reop bleed. The patient was re-explored for mediastinal bleeding and a hematoma was evacuated.  |
| 95. Unplanned Coronary<br>Artery Intervention<br>STS Sequence #: 6771           | 1 = Yes<br>2 = No   | Indicate if the patient had an unplanned coronary intervention (PCI) or unplanned surgical intervention on a coronary artery.  Intent/Clarification: Only capture surgical or Cath lab interventions that occur during the hospitalization prior to discharge. Capture an unplanned coronary intervention (PCI) or unplanned surgical intervention on a coronary artery in this field.  Yes No  |
|   |   | Percutaneous coronary intervention (PCI) is the placement of an angioplasty guide wire, balloon, or other device (e.g. stent, atherectomy, brachytherapy, or thrombectomy catheter) into a native coronary artery or coronary artery bypass graft for the purpose of mechanical coronary revascularization. An attempted, even if unsuccessful, PCI should be coded as a PCI.  Note: Patient had CABG, coded in ICU postop and a few days postop had to go to cath lab and near total graft occlusion (RCA) found. The MD attempted to perform a PCI of the RCA and was unable to wire/cross the lesion. Code this as Post-Op-Unplanned Coronary Artery Intervention since a PCI was attempted. An LHC cath done without an attempted intervention would not be captured. |
| 96. Unplanned Coronary<br>Artery Intervention-<br>Vessels<br>STS Sequence# 6772 | 1 = Native Coronary<br>2 = Graft<br>3 = Both  | Indicate the type of vessels that required postoperative reintervention.  |
| 97. Deep Sternal<br>STS Sequence #: 6700  | 2 = No;<br>3= Yes, within 30 days of<br>procedure;<br>4=Yes> 30 after procedure,<br>but during initial<br>hospitalization | Indicate whether a deep sternal wound infection or mediastinitis was diagnosed within 30 days of the OR date or at any time during the initial hospitalization.   |
| 98. Neuro-Stroke Permanent STS Sequence #: 6810                                 | 1 = Yes<br>2 = No   | Indicate whether the patient has a postoperative stroke (i.e., any confirmed neurological deficit of abrupt onset caused by a disturbance in blood supply to the brain) that was confirmed on imaging or did not resolve within 24 hours.   |

| Data Element  | Valid Values      | Definition  |
|---|-------------------|---|
|   |                   | Intent/Clarification: The intent is to capture whether the patient has a postoperative stroke (i.e. any confirmed neurological deficit of abrupt onset caused by a disturbance in blood supply to the brain) that was confirmed on imaging or did not resolve within 24 hours.  |
|   |                   | Stroke occurs when the blood supply to part of the brain is suddenly interrupted or when a blood vessel in the brain bursts, spilling blood into the spaces surrounding brain cells. Brain cells die when they no longer receive oxygen and nutrients from the blood or there is sudden bleeding into or around the brain.  |
|   |                   | The symptoms of a stroke include: Sudden numbness or weakness, especially on one side of the body Sudden confusion or trouble speaking or understanding speech Sudden trouble seeing in one or both eyes Sudden trouble with walking, dizziness, or loss of balance or coordination Sudden severe headache with no known cause  |
|   |                   | There are two forms of stroke: Ischemic - Blockage of a blood vessel supplying the brain. Includes embolic. Hemorrhagic - Bleeding into or around the brain   |
|   |                   | Central events are caused by embolic or hemorrhagic events. Neurological deficits such as confusion, delirium and/or encephalopathic (anoxic or metabolic) events are not to be coded in this field.  |
|   |                   | <b>Note:</b> Patient had an MRI that was positive for a post-op stroke. In discussion with the cardiac team it was felt that this would not be captured as a post-op stroke because the symptoms resolved within 24 hours. In this scenario, code YES to post-op CVA. The intent is to capture whether the patient has a postoperative stroke (i.e. any confirmed neurological deficit of abrupt onset caused by a disturbance in blood supply to the brain) that was confirmed on imaging <b>or</b> did not resolve within 24 hours. |
|   |                   | Reference: https://www.ninds.nih.gov/Disorders/All-Disorders/Stroke-Information-Page  |
| 99. Pulm – Ventilation<br>Prolonged<br>STS Sequence #: 6835 | 1 = Yes<br>2 = No | Indicate whether the patient had prolonged post-operative pulmonary ventilation > 24.0 hours.  The hours of postoperative ventilation time include OR exit until extubation, plus any additional hours following reintubation.  |

| Data Element   | Valid Values      | Definition   |
|--|-------------------|--|
|  |                   | Intent/Clarification: Includes any patient requiring mechanical ventilation > 24 hours postoperatively. To calculate total hours, include initial and additional hours of mechanical ventilation   |
| 100. Renal – Renal<br>Failure<br>STS Sequence #: 6870        | 1 = Yes<br>2 = No | Indicate whether the patient had acute renal failure or worsening renal function resulting in ONE OR BOTH of the following:  |
|  |                   | <ul> <li>A) Increase in serum creatinine level 3.0 x greater than baseline, or serum creatinine level ≥4 mg/dL,</li> <li>Acute rise must be at least 0.5 mg/dl</li> <li>B) A new requirement for dialysis postoperatively.</li> </ul>  |
|  |                   | Intent/Clarification: Baseline creatinine is the creatinine level closest to the date and time prior to surgery but prior to anesthetic management (induction area or operating room). If the patient was on dialysis pre-op, then do not code dialysis as a post-op complication. If pre-op dialysis is equal to "No" and if peak postoperative creatinine level is greater than or equal to 3X last creatinine level pre-op or postoperative creatinine is greater than or equal to 4.0 with a 0.5 mg/dL rise or new postoperative dialysis then, renal failure is equal to "Yes". |
| 101. Renal – Dialysis<br>Requirement<br>STS Sequence #: 6875 | 1 = Yes<br>2 = No | Indicate whether the patient had a new requirement for dialysis postoperatively, which may include hemodialysis, peritoneal dialysis.  |
| 102. Other – A Fib<br>STS Sequence #: 6945                   | 1 = Yes<br>2 = No | Indicate whether the patient experienced atrial fibrillation/flutter (AF) after OR Exit that a. last longer than one hour, or b. lasts less than one hour but requires medical or procedural intervention. Exclude patients who were in AFib at the start of surgery.  Intent/Clarification: Capture event(s) in all patients regardless if they have had a history of recent or remote A-fib, A-fib / flutter, or A-flutter who were not in A-fib, A-fib / flutter, or A-flutter at the start of surgery.   |
|  |                   | Medical or procedural intervention includes cardioversion for A-fib, A-fib / flutter, or A-flutter, AFib ablation, and medication to treat A-fib, A-fib / flutter, or A-flutter.   |

| Data Element  | Valid Values | Definition   |
|---|--------------|--|
|   |              | Use the first rhythm documented on the anesthesia record to determine if the patient is in A-fib, A-fib / flutter, or A-flutter at the time of entry to OR. If there is no documentation on the anesthesia record, then use the rhythm documented closest to OR entry to determine if patient in A-fib, A-fib / flutter, or A-flutter. |
|   |              | Example # 1: A patient is on beta blockers post-op and is titrating each day to give higher doses. The second post-op day the patient has a two-hour run of AFib. During this run of AFib, the beta blocker is increased, or an extra dose of beta blocker is given. This is considered a post-op AFib event.                          |
|   |              | Example # 2: A patient who has no history of AFib is on an AFib protocol preoperatively to prevent post-<br>op AFib; the patient then goes in to atrial fibrillation (AF) postoperatively and the protocol is not<br>adjusted: this should be coded "Yes" as a post op AFib event.   |
| 103. Facility Identification Number CCORP-specific variable |              | The six-digit facility identification number assigned to a hospital by the Office of Statewide Health Planning and Development (OSHPD), as defined in Section 97170.   |

### Type of CABG (\*definitional reference):

Was the surgery an Isolated CABG, CABG + Valve, or Other Non-Isolated CABG?

Valid Values

1=Isolated CABG

3= CABG + Valve

4=Other Non-Isolated CABG

Definition

# Type of CABG

#### **Isolated CABG**

#### **Exclusions from Isolated CABG:**

- Valve repairs or replacements
- Operations on structures adjacent to heart valves (papillary muscle, chordae tendineae, traebeculae carneae cordis, annuloplasty, infundibulectomy)
- Ventriculectomy when diagnosed preoperatively as a rupture, aneurysm or remodeling procedure. But not 1) sites intra-operatively diagnosed, 2) patch applications for site oozing discovered during surgery and 3) prophylactic patch applications to reduce chances of future rupture
- Repair of atrial and ventricular septa, but not closure of patent foramen ovale
- Excision of aneurysm of heart
- · Head and neck, intracranial endarterectomy
- Other open heart surgeries, such as aortic arch repair, pulmonary endarterectomy
- Endarterectomy of aorta
- Thoracic endarterectomy (endarterectomy on an artery outside the heart)
- Carotid endarterectomy
- Heart transplantation
- Repair of certain congenital cardiac anomalies, but not closure of patent foramen ovale (e.g., teratology of fallot, atrial septal defect (ASD), ventricular septal defect (VSD), valvular abnormality)
- Any aortic aneurysm repair (abdominal or thoracic)
- Aorta-subclavian-carotid bypass
- Aorta-renal bypass

- Aorta-iliac-femoral bypass
- Caval-pulmonary artery anastomosis
- Extracranial-intracranial (EC-IC) vascular bypass
- Coronary artery fistula
- Resection of a lobe or segment of the lung (e.g., lobectomy or segmental resection of lung). But not simple biopsy of lung nodule in which surrounding lung is not resected, biopsy of a thoracic lymph node or excision or stapling of an emphysematous bleb.
- Pleural decortication
- Mastectomy for breast cancer (not simple breast biopsy)
- Amputation of any extremity (e.g., foot or toe)
- Resection of LV aneurysm
- Planned Ventricular Assist Device (VAD) for long term treatment.
- Septal myectomy with hypertrophic obstructive cardiomyopathy
- Full open mazes
- Repair of aortic dissection

#### CABG + Valve

CABG + Valve includes all CABG cases with aortic valve replacement (AVR), mitral valve replacement (MVR), mitral valve repair (MVRepair) and AVR +MVR/MVRepair

#### Exclusions from CABG + Valve:

- Aortic Valve repair
- Aortic Valve root replacement with valved conduit (Bentall)
- Pulmonic Valve Procedure
- Tricuspid Valve Procedure
- Ventriculectomy when diagnosed preoperatively as a rupture, aneurysm or remodeling procedure. But not 1) sites intra-operatively diagnosed, 2) patch
  applications for site oozing discovered during surgery and 3) prophylactic patch applications to reduce chances of future rupture
- Repair of atrial and ventricular septa, but not closure of patent foramen ovale
- Excision of aneurysm of heart
- Head and neck, intracranial endarterectomy
- Other open heart surgeries, such as aortic arch repair, pulmonary endarterectomy
- Endarterectomy of aorta
- Thoracic endarterectomy (endarterectomy on an artery outside the heart)
- Carotid endarterectomy
- Heart transplantation

- Repair of congenital cardiac anomalies, such as tetralogy of fallot, atrial septal defect (ASD), ventricular septal defect or other complex anomaly
- Any aortic aneurysm repair (abdominal or thoracic)
- Repair of aortic dissection
- Aorta-subclavian-carotid bypass
- Aorta-renal bypass
- Aorta-iliac-femoral bypass
- Caval-pulmonary artery anastomosis
- Extracranial-intracranial (EC-IC) vascular bypass
- Coronary artery fistula
- Resection of a lobe or segment of the lung (e.g., lobectomy or segmental resection of lung). But not simple biopsy of lung nodule in which surrounding lung is not resected, biopsy of a thoracic lymph node or excision or stapling of an emphysematous bleb.
- Pleural decortication
- Mastectomy for breast cancer (not simple breast biopsy)
- Amputation of any extremity (e.g., foot or toe)
- Resection of LV aneurysm
- Planned Ventricular Assist Device (VAD) for long term treatment.
- Infundibulectomy
- Septal myectomy with hypertrophic obstructive cardiomyopathy
- Full Open MAZE for Aortic Valve cases only (epicardial MAZE procedures are coded as CABG + Valve and Full Open MAZE procedures for Mitral Valve are also coded as CABG + Valve.).

### Other Non-Isolated

All other non-isolated CABGs

Must include a CABG (not isolated Valves)

# Responsible Surgeon Name (\*\*definitional reference):

"Responsible surgeon" means the principle surgeon who performs a coronary artery bypass procedure.

The first and last name collected should exactly match the name assigned to the license number issued by the California Medical Board.

The middle initial collected should match the first letter of the middle name assigned to the license number issued by the California Medical Board. Example: if a surgeon's middle name is Harry, the middle initial should be reported as 'H'. NOTE: do not include period (.).

If a trainee performs this procedure, then the responsible surgeon is the physician responsible for supervising this procedure performed by the trainee. In situations in which a responsible surgeon cannot otherwise be determined, the responsible surgeon is the surgeon who bills for the coronary artery bypass procedure.