Overview Statement

The purpose of these clinical guidelines is to assist healthcare professionals in selecting the medical service that may be appropriate and supported by evidence to improve patient outcomes. These clinical guidelines neither preempt the clinical judgment of trained professionals nor advise anyone on how to practice medicine. The healthcare professionals are responsible for all clinical decisions based on their assessment. These clinical guidelines do not provide authorization, certification, explanation of benefits, or guarantee of payment, nor do they substitute for, or constitute, medical advice.

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# Table of Contents

1. Overview Statement  
2. Table of Contents  
3. Cardiac Ablation  
4. Cardiac Catheterization  
5. Cardiac Defibrillation Device  
6. Sub-Cutaneous Implantable Cardiac Defibrillator (SICD)  
7. Automatic Implantable Cardiac Defibrillator – Insertion (ICD)  
8. Automatic Implantable Cardiac Defibrillator – Removal or Replacement (ICD)  
9. Cardiac Electrophysiology Studies  
10. Cardiac Pacemaker Device  
11. Cardiac Positron Emission Test  
12. Cardiac Resynchronization Therapy  
13. Cardiac Resynchronization Therapy: Defibrillator (CRT-D)  
14. Cardiac Resynchronization Therapy: Pacemaker (CRT-P)  
15. Cardiac Single Photon Emission Computerized Tomography  
16. Coronary Computed Tomography Angiography
<table>
<thead>
<tr>
<th>Service</th>
<th>Page</th>
</tr>
</thead>
<tbody>
<tr>
<td>Echocardiogram</td>
<td>73</td>
</tr>
<tr>
<td>Transthoracic Echocardiogram (TTE)</td>
<td>73</td>
</tr>
<tr>
<td>Tranesophageal Echocardiogram (TEE)</td>
<td>92</td>
</tr>
<tr>
<td>Implantable Loop Recorder</td>
<td>101</td>
</tr>
<tr>
<td>Leadless Intracardiac Pacemaker</td>
<td>113</td>
</tr>
<tr>
<td>Left Atrial Appendage Closure</td>
<td>115</td>
</tr>
<tr>
<td>MRA Heart</td>
<td>117</td>
</tr>
<tr>
<td>MRI Heart</td>
<td>119</td>
</tr>
<tr>
<td>Percutaneous Coronary Interventions</td>
<td>121</td>
</tr>
<tr>
<td>Percutaneous Ventricular Assistive Device</td>
<td>128</td>
</tr>
<tr>
<td>Trans catheter Aortic Valve Replacement</td>
<td>131</td>
</tr>
<tr>
<td>Trans catheter Mitral Valve Replacement</td>
<td>136</td>
</tr>
<tr>
<td>Wearable Cardiac Defibrillator</td>
<td>139</td>
</tr>
</tbody>
</table>
Cardiac Ablation

Utilization of a cardiac ablation may be medically appropriate and supported by evidence to improve patient outcomes for the following indications.

- Ablation for atrial fibrillation (a-fib) may be reasonable and appropriate when the patient's medical record demonstrates EITHER of the following: (2, 6)
  o Patient has atrial fibrillation that terminates spontaneously within seven (7) days of onset or within 48 hours or less with electrical or pharmacological cardioversion; (2,6)
  o Patient has continuous atrial fibrillation that is sustained for greater than seven (7) days or does not convert after greater than 48 hours following electrical or pharmacological cardioversion; and EITHER of the following: (1,8)
    ▪ Patient had poor response or was intolerant of at least one antiarrhythmic medication; (2,7)
    ▪ Patient's lifestyle would be severely limited by atrial fibrillation or the patient does not prefer long-term therapy. (1,6-7)

- Ablation for supraventricular tachycardia (SVT) may be reasonable and appropriate when the patient's medical record demonstrates EITHER of the following:
  o Patient is diagnosed with atrioventricular nodal reentrant tachycardia (AVNRT); and ANY of the following:
    ▪ Patient had poor response or was intolerant of at least one antiarrhythmic medication; (2)
    ▪ Patient's lifestyle would be severely limited by atrial fibrillation or the patient does not prefer long-term therapy; (7)
    ▪ Prior EPS illustrated rapid ventricular rate of concomitant arrhythmia;
    ▪ Patient is intolerant of drug therapy with recurrent break-through episodes that are life altering. (2)
  o Patient is diagnosed with Wolf-Parkinson-White (WPW) with tachycardia or other atrioventricular reentrant tachycardia, there is rapid ventricular response via accessory pathway noted; and EITHER of the following: (3)
- Patient had poor response or was intolerant of at least one antiarrhythmic medication; (3,4)
- Patient is intolerant of drug therapy with recurrent break-through episodes that are life altering.

- Ablation for ventricular tachycardia (VT) may be reasonable and appropriate when the patient’s medical record demonstrates ANY of the following:
  - Patient has a history of sustained monomorphic VT; and ANY of the following:
    - Patient is symptomatic with drug-resistant tachycardia; (5)
    - Patient is symptomatic and intolerant of drug therapy; (9)
    - Patient is symptomatic and does not desire long-term drug therapy;
    - Patient has an implantable cardioverter defibrillator (ICD) and is receiving multiple defibrillations not manageable by reprogramming the device or additive drug therapy.
  - Patient has bundle branch reentrant tachycardia; (5)
  - Patient has non-sustained monomorphic VT, which is resistant to drug therapy.
  - Patient is drug intolerant. (5)
  - Patient does not desire long-term drug therapy.
REFERENCES:


Cardiac Catheterization

Utilization of a cardiac catheterization may be medically appropriate and supported by evidence to improve patient outcomes for the following indications.

Left Heart Catheterization

Left heart catheterization for acute coronary syndrome, ST-segment elevated myocardial infarction (STEMI) or non-ST segment elevated myocardial infarction (Non-STEMI) may be reasonable and appropriate when the patient's medical record demonstrates EITHER of the following:

- Documented Non-STEMI within the past three (3) months; and ANY of the following:
  - Previous Coronary Computerized Tomography Angiography (CCTA) demonstrates moderate or severe proximal obstructive vessel disease;
  - Patient is exhibiting persistent cardiac symptoms despite anti-anginal therapy;
  - Previous physiologic or pharmacologic noninvasive imaging (i.e. Cardiac PET, SPECT or Stress Echocardiogram) showing definite reversible ischemia or high-risk treadmill study (i.e. > 2mm ST depression, Non-sustained VT, hypotension or failure to increase BP with exercise).
- Documented STEMI in the past twenty-four (24) hours.

Left heart catheterization to evaluate worsening or recurrent chest pain, ischemic equivalent symptoms or stable patient post revascularization (e.g. PCI or CABG) may be reasonable and appropriate when the patient's medical record demonstrates ANY of the following:

- Patient is exhibiting worsening or limiting chest pain/ischemic equivalent despite anti-anginal therapy; and ANY of the following:
  - This request is for a diagnostic left heart catheterization only, no PCI is currently anticipated;
- Previous angiogram is suggestive of 50-79% stenosis and the ordering physician will perform a FFR for blockage less than or equal to 0.8% prior to PCI;
- Previous physiologic or pharmacologic non-invasive imaging (i.e. Cardiac PET, SPECT or Stress Echocardiogram) or high-risk treadmill study demonstrates greater than or equal to 10% reversible myocardial ischemia with possible PCI. (3)
- Previous non-invasive imaging (i.e. Cardiac PET, SPECT or Stress Echocardiogram) resulted in equivocal, artefactual or discordant findings making interpretation uncertain or unclear and this request is for a diagnostic left heart catheterization with no planned PCI. (3)
  - Patient’s symptoms are stable or controlled with medical management; and
  - EITHER of the following:
    1. Previous physiologic or pharmacologic non-invasive imaging (i.e. Cardiac PET, SPECT or Stress Echocardiogram) or high risk treadmill study demonstrates greater than or equal to 10% reversible myocardial ischemia with or without possible PCI; (3)
    - Patient has a known incomplete revascularization after previous PCI and staged additional PCI is planned.
  - Patient is exhibiting unexplained post procedural symptoms which could possibly represent continuing myocardial ischemia; and the following:
    - Previous physiologic or pharmacologic non-invasive imaging (i.e. Cardiac PET, SPECT or Stress Echocardiogram) or high-risk treadmill study demonstrates greater than or equal to 10% reversible myocardial ischemia with or without possible PCI. (3)
  - Patient has had a return of the symptoms for which the previous revascularization was performed and EITHER of the following: (7)
    - Diagnostic Cath with no planned PCI.
    - Previous angiogram is suggestive of 50-79% stenosis and the ordering physician will perform a FFR for blockage less than or equal to 0.8% prior to PCI with possible PCI.
  - PCI was performed within the past three (3) months or patient has had a previous coronary artery bypass graft (CABG) and the following:
    - Previous physiologic or pharmacologic non-invasive imaging (i.e. Cardiac PET, SPECT or Stress Echocardiogram) or high-risk treadmill study
demonstrates greater than or equal to 10% reversible myocardial ischemia. (3)

Left heart catheterization related to cardiac transplant may be reasonable and appropriate when the patient's medical record demonstrates that the patient is a candidate for a heart transplant or has had a heart transplant and a surveillance study is deemed necessary per the post-transplant protocol of the treating cardiologist.

Left heart catheterization requested with or without planned PCI for follow-up evaluation of an abnormal CCTA, suggesting coronary artery disease (CAD) may be reasonable and appropriate when the patient's medical record demonstrates the following: (13)

- Chest pain or ischemic equivalent; and EITHER of the following:
  - CCTA demonstrates a lesion obstructing 50% or more of at least one (1) proximal coronary artery segment.
  - CCTA is non-diagnostic, equivocal or demonstrates lesions of unclear severity.

Left heart catheterization requested for evaluation of known or suspected congenital heart disease is reasonable.

Left heart catheterization for coronary assessment secondary to aortic dissection, aneurysm or congestive heart failure (CHF) may be reasonable and appropriate when the patient's medical record demonstrates ANY of the following: (10)

- Diagnosis of thoracic aortic dissection;
- Diagnosis of ascending aortic aneurysm;
- Evaluation of signs and symptoms of congestive heart failure persistent or worsening despite pharmacological therapy with confirmed diastolic dysfunction via TTE or previous cardiac catheterization; and EITHER of the following:
  - PCI is planned and previous angiogram is suggestive of 50-79% stenosis; and EITHER of the following:
    - Previous physiologic or pharmacologic non-invasive imaging (i.e. Cardiac PET, SPECT or Stress Echocardiogram) or high-risk treadmill study demonstrates greater than or equal to 10% reversible myocardial ischemia.
• CCTA demonstrates a lesion obstructing 50% or more of at least one (1) proximal coronary artery segment.
  - PCI is not planned; and EITHER of the following:
    - Previous physiologic or pharmacologic non-invasive imaging (i.e. Cardiac PET, SPECT or Stress Echocardiogram) or high-risk treadmill study demonstrates greater than or equal to 10% reversible myocardial ischemia.  (3)
    - CCTA demonstrates a lesion obstructing 50% or more of at least one (1) proximal coronary artery segment.

Left heart catheterization for valvular heart disease may be reasonable and appropriate when the patient's medical record demonstrates EITHER of the following:  (8)
  - Assessing hemodynamics or chamber size when non-invasive testing is incomplete or discordant;
  - Pre-operative planning study for cardiac valve surgery in patient's age 50 or older.  (8)

Left heart catheterization for evaluation of suspected coronary artery disease in a symptomatic patient exhibiting typical angina may be reasonable and appropriate when the patient's medical record demonstrates EITHER of the following:  (12)
  - Male patient 40 years of age of older; and EITHER of the following:
    - PCI is planned and previous angiogram is suggestive of 50-79% stenosis; and ANY of the following:
      - Previous physiologic or pharmacologic non-invasive imaging (i.e. Cardiac PET, SPECT or Stress Echocardiogram) or high-risk treadmill study demonstrates 5-10% reversible myocardial ischemia.  (3)
      - Previous physiologic or pharmacologic non-invasive imaging (i.e. Cardiac PET, SPECT or Stress Echocardiogram) or high-risk treadmill study demonstrates greater than or equal to 10% reversible myocardial ischemia.  (3)
      - Non-invasive testing (i.e. Cardiac PET, SPECT or Stress Echocardiogram) is non-diagnostic or equivocal making interpretation of the test uncertain or unclear.  (3)
    - PCI is not planned; and ANY of the following:
• Previous physiologic or pharmacologic non-invasive imaging (i.e. Cardiac PET, SPECT or Stress Echocardiogram) or high-risk treadmill study demonstrates 5-10% reversible myocardial ischemia. (3)
• Previous physiologic or pharmacologic non-invasive imaging (i.e. Cardiac PET, SPECT or Stress Echocardiogram) or high-risk treadmill study demonstrates greater than or equal to 10% reversible myocardial ischemia. (3)
• Non-invasive testing (i.e. Cardiac PET, SPECT or Stress Echocardiogram) is non-diagnostic or equivocal making interpretation of the test uncertain or unclear. (3)

- Female patient 60 years of age or older; and EITHER of the following:
  - PCI is planned and previous angiogram is suggestive of 50-79% stenosis; and ANY of the following:
    - Previous physiologic or pharmacologic non-invasive imaging (i.e. Cardiac PET, SPECT or Stress Echocardiogram) or high risk treadmill study demonstrates 5-10% reversible myocardial ischemia. (3)
    - Previous physiologic or pharmacologic non-invasive imaging (i.e. Cardiac PET, SPECT or Stress Echocardiogram) or high-risk treadmill study demonstrates greater than or equal to 10% reversible myocardial ischemia. (3)
    - Non-invasive testing (i.e. Cardiac PET, SPECT or Stress Echocardiogram) is non-diagnostic or equivocal making interpretation of the test uncertain or unclear. (3)
  - PCI is not planned; and ANY of the following:
    - Previous physiologic or pharmacologic non-invasive imaging (i.e. Cardiac PET, SPECT or Stress Echocardiogram) or high-risk treadmill study demonstrates 5-10% reversible myocardial ischemia. (3)
    - Previous physiologic or pharmacologic non-invasive imaging (i.e. Cardiac PET, SPECT or Stress Echocardiogram) or high-risk treadmill study demonstrates greater than or equal to 10% reversible myocardial ischemia. (3)
• Non-invasive testing (i.e. Cardiac PET, SPECT or Stress Echocardiogram) is non-diagnostic or equivocal making interpretation of the test uncertain or unclear. (3)

Left heart catheterization with no plan to perform a PCI for evaluation of left ventricular function may be reasonable and appropriate when the patient's medical record demonstrates left ventricular ejection fraction (LVEF) of 45% or less and EITHER of the following:

- PCI is planned and previous angiogram is suggestive of 50-79% stenosis; Patient is under consideration for coronary bypass; and ANY of the following:
  - New or worsening angina or anginal equivalent (i.e. dyspnea, diaphoresis, arm or jaw pain, profuse vomiting in a diabetic patient) and the previous heart catheterization was performed greater than one (1) year ago and demonstrated ischemic coronary disease;
  - Recent non-invasive (i.e. cardiac PET/SPECT, Stress Echocardiogram, MUGA or Diagnostic Echocardiogram) testing shows a new decline in LVEF of greater than 10%;
- PCI is not planned; and ANY of the following:
  - New or worsening angina or anginal equivalent (i.e. dyspnea, diaphoresis, arm or jaw pain, profuse vomiting in a diabetic patient) and the previous heart catheterization was performed greater than one (1) year ago and demonstrated ischemic coronary disease;
  - Recent non-invasive (i.e. cardiac PET/SPECT, Stress Echocardiogram, MUGA or Diagnostic Echocardiogram) testing shows a new decline in LVEF of greater than 10%;
  - Patient is under consideration for coronary bypass.

Left heart catheterization for clearance prior to a non-cardiac surgical procedure may be reasonable and appropriate when the patient's medical record demonstrates ANY of the following:

- PCI is planned and previous angiogram is suggestive of 50-79% stenosis; and ANY of the following:
  - Male 40 years of age or older exhibiting typical angina or anginal equivalent; and ANY of the following:
    - BMI of greater than thirty-five (35);
- Coronary calcium score of 500 or greater;
- History of atrial fibrillation.

- Female 60 years of age or older exhibiting typical angina or anginal equivalent; and **ANY** of the following:
  - BMI of greater than 35;
  - Coronary calcium score of 500 or greater;
  - History of atrial fibrillation.

- Patient with a history of congestive heart failure; and **ANY** of the following: \(^{(11)}\)
  - BMI of greater than 35;
    - Coronary calcium score of 500 or greater;
    - History of atrial fibrillation.

- **PCI is not planned; and ANY** of the following:
  - Male 40 years of age or older exhibiting typical angina or anginal equivalent; and **ANY** of the following:
    - BMI of greater than thirty-five (35);
    - Coronary calcium score of 500 or greater;
    - History of atrial fibrillation.

- Female 60 years of age or older exhibiting typical angina or anginal equivalent; and **ANY** of the following:
  - BMI of greater than 35;
  - Coronary calcium score of 500 or greater;
  - History of atrial fibrillation.

- **Patient with a history of congestive heart failure; and ANY** of the following: \(^{(10)}\)
  - BMI of greater than 35;
  - Coronary calcium score of 500 or greater;
  - History of atrial fibrillation.

- Previous physiologic or pharmacologic non-invasive imaging (i.e. Cardiac PET, SPECT or Stress Echocardiogram) or high risk treadmill study demonstrates greater than or equal to 10% reversible myocardial ischemia \(^{(3)}\)

- CCTA demonstrates a lesion obstructing 50% or more of at least one proximal coronary artery segment
Left and Right Heart Catheterization

Left and Right heart catheterization related to cardiac transplantation may be reasonable and appropriate when the patient's medical record demonstrates ANY of the following:

- Pre-cardiac transplantation evaluation; (5)
- Post cardiac transplantation for routine annual evaluation; (5)
- Post cardiac transplantation for evaluation of possible organ rejection.

Left and Right heart catheterization for evaluation of chest pain when dyspnea is a major associated symptom may be reasonable and appropriate when the patient's medical record demonstrates clinical signs and symptoms suggesting pulmonary hypertension and evaluation with a non-invasive study was equivocal, inadequate or discrepant. (12)

Left and Right heart catheterization for evaluation of chronic heart failure (CHF) may be reasonable and appropriate when the patient's medical record demonstrates the following: (10, 11)

- Clinical signs and symptoms suggesting pulmonary hypertension and evaluation with a non-invasive study was equivocal, inadequate or discrepant; and EITHER of the following:
  - Worsening heart failure with suspected ischemic etiology despite medical management;
  - Diagnosis of CHF with new onset chest pain or chest pain that in not responding to medical management. (11)

Left and Right heart catheterization for new onset heart failure may be reasonable and appropriate when the patient's medical record demonstrates EITHER of the following: (10)

- Clinical signs and symptoms suggesting pulmonary hypertension and evaluation with a non-invasive study was equivocal, inadequate or discrepant; and EITHER of the following:
  - Multiple risk factors for coronary artery disease (CAD) with no previous evaluation;
  - Chest pain or ischemic equivalent is present.
Pulmonary pressure of greater than 25mmHg at rest or 30mmHg during physical activity; and EITHER of the following:
- Multiple risk factors for coronary artery disease (CAD) with no previous evaluation;
- Chest pain or ischemic equivalent is present.

Left and Right heart catheterization for evaluation of pulmonary hypertension may be reasonable and appropriate when the patient's medical record demonstrates EITHER of the following:
- Chest pain or ischemic equivalent is present; and ANY of the following: (12)
  - Request is for an evaluation of new pulmonary hypertension diagnosis;
  - Patient has an established diagnosis of pulmonary hypertension with unexplained worsening symptoms or to titrate therapy;
  - Clinical signs and symptoms suggesting pulmonary hypertension and evaluation with a non-invasive study was equivocal, inadequate or discrepant.
- Non-invasive testing suggestive of ischemia; and EITHER of the following: (12)
  - Request is for an evaluation of new pulmonary hypertension diagnosis;
  - Clinical signs and symptoms suggesting pulmonary hypertension and evaluation with a non-invasive study was equivocal, inadequate or discrepant. (3)

Left and Right heart catheterization for evaluation of valvular heart disease may be reasonable and appropriate when the patient's medical record demonstrates ANY of the following: (6)
- Patient is at high risk for death;
- Pre-operative planning for valvular surgery; and ANY of the following: (6)
  - Non-invasive imaging is discordant, discrepant or inadequate for determining the severity of valvular disease;
  - Non-invasive imaging is discordant, discrepant or inadequate for determining pulmonary hypertension;
  - Non-invasive imaging is discordant, discrepant or inadequate for evaluation of ischemia.
  - Chest pain or ischemic equivalent.
  - To confirm valvular disease is severe. (6)
Features of ischemia are present on non-invasive testing but the non-invasive imaging is discordant, discrepant or inadequate for determining the severity of valvular disease; (6)

Chest pain or ischemic equivalent is present but non-invasive imaging is discordant, discrepant or inadequate for determining the severity of valvular disease; (12)

### Right Heart Catheterization

Right heart catheterization for post cardiac transplant may be reasonable and appropriate when the patient's medical record demonstrates **ANY** of the following:
- Routine annual evaluation; (15)
- Evaluation of possible organ rejection;
- Evaluation of worsening congestive heart failure or unknown etiology.

Right heart catheterization for evaluation of cardiomyopathy/ congestive heart failure (CHF) may be reasonable and appropriate when the patient's medical record demonstrates **EITHER** of the following; (9, 10)
- Cardiomyopathy with clinical deterioration to titrate inotrope or vasodilator therapy;
- CHF with clinical deterioration of unclear etiology. (10)

Right heart catheterization for evaluation of congenital heart disease may be reasonable and appropriate when the patient's medical record demonstrates **BOTH** of the following:
- Diagnosis of congenital heart disease; (14)
- Non-invasive imaging is discordant, discrepant or equivocal.

Right heart catheterization for pulmonary hypertension may be reasonable and appropriate when the patient's medical record demonstrates **ANY** of the following:
- Request is for an evaluation of new pulmonary hypertension diagnosis; (4)
- Patient has an established diagnosis of pulmonary hypertension with unexplained worsening symptoms or to titrate therapy; (13)
- Clinical signs and symptoms suggesting pulmonary hypertension and evaluation with a non-invasive study was equivocal, inadequate or discrepant. (13)

Right heart catheterization for evaluation of valvular heart disease may be reasonable and appropriate when the patient's medical record demonstrates that non-invasive testing was discrepant, equivocal or inadequate to determine the severity of valvular stenosis or regurgitation. (2, 8)
REFERENCES:


Cardiac Defibrillation Device

Utilization of an implantable cardiac defibrillation device may be medically appropriate and supported by evidence to improve patient outcomes for the following indications.

**Sub-Cutaneous Implantable Cardiac Defibrillator (SICD)**

SICD utilization for the primary prevention of Sudden Cardiac Death may be reasonable and appropriate when the patient’s medical record demonstrates that the patient does not have symptomatic bradycardia, incessant ventricular tachycardia, spontaneous, frequently occurring ventricular tachycardia, which has been reliably, terminated with anti-tachycardia pacing or a unipolar pacemaker **ANY** of the following:

- Patient is awaiting a cardiac transplant in an outpatient setting; (2,8)
- Patient who is diagnosed with Cardiac Sarcoidosis or Chagas Disease and has an ejection fraction (EF) of less than or equal to 40%; (3,8)
- Patient is diagnosed with hypertrophic cardiomyopathy and **ANY** of the following: (1, 2, 5)
  - Family history of sudden cardiac death; (3)
  - Non- sustained ventricular tachycardia; (1, 2, 4, 5)
  - Massive Left ventricular hypertrophy (wall thickness of greater than 30mm); (3)
  - Syncope (2)
  - Abnormal blood pressure in response to exercise (8)
- Patient who is diagnosed with ischemic cardiomyopathy and **ANY** of the following: (2, 3)
- NYHA Class 2 or 3, EF less than 35% and it has been more than forty (40) days since a myocardial infarction; (2)
- NYHA Class 1, EF of less than 30% and it has been more than forty (40) days since a myocardial infarction; (2)
- Non-sustained ventricular tachycardia with electrophysiology study (EPS) demonstrating inducible ventricular tachycardia or ventricular fibrillation and an EF of less than 40%; (4)
- NYHA Class 4
  - Primary prevention in a patient diagnosed with long QT syndrome; (3)
  - Patient is diagnosed with Non-Ischemic Cardiomyopathy, EF of less than or equal to 35% and NYHA Class 2 or 3; (2)
  - Patient is diagnosed with right ventricular dysplasia and **ANY of the following:** (2)
    - Patient is less than fifty (50) years of age;
    - Patient is experiencing syncope; (2)
    - Patient is having an increase in QRS dispersion; (8)
    - Patient has ventricular involvement; (2)
    - ARVC2 or ARVC5 positive; (3)
    - Patient has a previous history of cardiac arrest or of ventricular tachycardia with hemodynamic compromise (4)

SICD utilization for the secondary prevention of Sudden Cardiac Death may be reasonable and appropriate when the patient’s medical record demonstrates that the patient does not have symptomatic bradycardia, incessant ventricular tachycardia, spontaneous, frequently occurring ventricular tachycardia, which has been reliably, terminated with anti-tachycardia pacing or a unipolar pacemaker **ANY of the following:**

- Patient is diagnosed with Bragada Syndrome and **EITHER of the following:** (5)
• Patient is experiencing syncope; (2, 5)
• Documentation of ventricular tachycardia is present (3)
  o Patient is a cardiac arrest survivor secondary to Ventricular tachycardia or ventricular fibrillation and no reversible causes for the arrhythmia have been found; (2,4,6)
  o Patient has documented spontaneous sustained ventricular tachycardia and no reversible causes for the arrhythmia have been found; (1, 2, 6, 9)
  o Patient is exhibiting syncope of unknown etiology, and EITHER of the following (2,9)
    ▪ Patient has had a comprehensive work-up to determine cause of syncope without a clear etiology having been identified, EF of less than 50%; (2,8)
    ▪ EPS demonstrates sustained ventricular tachycardia or ventricular fibrillation (3,6)

Automatic Implantable Cardiac Defibrillator – Insertion (ICD)

Single Chamber ICD utilization for the primary prevention of Sudden Cardiac Death may be reasonable and appropriate when the patient's medical record demonstrates ANY of the following:

  o Patient is awaiting a cardiac transplant in an outpatient setting; (11)
  o Patient who is diagnosed with Cardiac Sarcoidosis or Chagas Disease and has an ejection fraction (EF) of less than or equal to 40%; (10, 14)
  o Patient is diagnosed with hypertrophic cardiomyopathy and ANY of the following:
    ▪ Family history of sudden cardiac death; (12)
    ▪ Non-sustained ventricular tachycardia; (12)
- Massive Left ventricular hypertrophy (wall thickness of greater than 30mm); (12)
- Syncope; (12)
- Abnormal blood pressure in response to exercise (12)
  - Patient who is diagnosed with ischemic cardiomyopathy and **ANY** of the following:
    - NYHA Class 2 or 3, EF less than or equal to 35% and it has been more than forty (40) days since a myocardial infarction; (13, 14)
    - NYHA Class 1, EF of less than or equal to 30% and it has been more than forty (40) days since a myocardial infarction; (13, 14)
    - Non-sustained ventricular tachycardia with electrophysiology study (EPS) demonstrating inducible ventricular tachycardia or ventricular fibrillation and an EF of less than or equal to 40%; (14)
  - Primary prevention in a patient diagnosed with long QT syndrome and experiencing syncope or ventricular tachycardia on beta-blocker. (14)
  - Patient is diagnosed with Non-Ischemic Cardiomyopathy, EF of less than or equal to 35% and NYHA Class 2 or 3; (14)
  - Patient is diagnosed with right ventricular dysplasia and **ANY** of the following:
    - Patient is less than fifty (50) years of age; (17)
    - Patient is experiencing syncope; (14)
    - Patient is having an increase in QRS dispersion; (17)
    - Patient has left ventricular involvement; (15)
    - ARVC2 or ARVC5 positive; (17)
    - Patient has a previous history of cardiac arrest or of ventricular tachycardia with hemodynamic compromise (14, 16)

Single Chamber ICD utilization for the secondary prevention of Sudden Cardiac Death may be reasonable and appropriate when the patient's medical record demonstrates **ANY** of the following:
- Patient is diagnosed with Bragada Syndrome and has EITHER syncope or documented ventricular tachycardia; (14)
- Cardiac arrest survivor from ventricular tachycardia or ventricular fibrillation with no reversible cause identified; (18)
- Spontaneous sustained ventricular tachycardia with no reversible cause identified; (14)
- Syncope of unknown etiology and EITHER of the following:
  - Electrophysiology study shows ventricular fibrillation of sustained ventricular tachycardia (14)
  - Comprehensive work up completed with no clear etiology, left ventricular dysfunction with an ejection fraction of <50%, and non-ischemic etiology (18)

Dual Chamber ICD utilization for the primary prevention of Sudden Cardiac Death may be reasonable and appropriate when the patient’s medical record demonstrates ANY of the following:

- Patient is awaiting cardiac transplant and has a non-ischemic etiology with ANY of the following:
  - Documented sinus node disease with bradycardia that may need pacing; (17)
  - 2nd or 3rd degree AV block; (17)
  - Indications for a dual chamber pacemaker; (17)
  - Anti-tachycardia pacing will be an important part of treatment. (10, 17)
- Patient who is diagnosed with Cardiac Sarcoidosis or Chagas Disease and has an ejection fraction (EF) of less than or equal to 40% and ANY of the following:
  - Documented sinus node disease with bradycardia that may need pacing;
  - 2nd or 3rd degree AV block;
  - Indications for a dual chamber pacemaker;
- Anti-tachycardia pacing will be an important part of treatment (10)
  - Patient is diagnosed with hypertrophic cardiomyopathy and ANY of the following:
    - Family history of sudden cardiac death and EITHER
    - Documented sinus node disease with bradycardia that may need pacing; (19)
    - 2nd or 3rd degree AV block; (20)
    - Indications for dual chamber pacing; (19)
    - Anti-tachycardia pacing will be an important part of treatment. (19)
  - Patient is diagnosed with syncope, non-sustained ventricular tachycardia OR is in the outpatient setting, and EITHER:
    - Documented sinus node disease with bradycardia that may need pacing; (20)
    - 2nd or 3rd degree AV block; (20)
    - Indications for dual chamber pacing; (20)
    - Anti-tachycardia pacing will be an important part of treatment. (20)
  - Patient is diagnosed with ischemic cardiomyopathy and ANY of the following:
    - Ejection fraction is less than or equal to 30%, NYHA class 1 and less than 40 days post myocardial infarction and EITHER: (20)
    - Documented sinus node disease with bradycardia that may need pacing; (20)
    - 2nd or 3rd degree AV block; (20)
    - Indications for dual chamber pacing; (20)
    - Anti-tachycardia pacing will be an important part of treatment. (20)
  - Ejection fraction is less than or equal to 35%, NYHA class 2 or 3 and less than 40 days post myocardial infarction and EITHER: (20)
    - Documented sinus node disease with bradycardia that may need pacing; (20)
    - 2nd or 3rd degree AV block; (20)
    - Indications for dual chamber pacing; (20)
- Anti-tachycardia pacing will be an important part of treatment. (20)
  - Ejection fraction is less than or equal to 40%, with non-sustained ventricular tachycardia, and inducible ventricular tachycardia/fibrillation on electrophysiology study, and EITHER: (20)
    - Documented sinus node disease with bradycardia that may need pacing; (20)
    - 2nd or 3rd degree AV block; (20)
    - Indications for dual chamber pacing; (20)
    - Anti-tachycardia pacing will be an important part of treatment. (20)
  - Patient is diagnosed with Long QT Syndrome with syncope or ventricular tachycardia on beta blockers with ANY of the following:
    - Documented sinus node disease with bradycardia that may need pacing; (18)
    - 2nd or 3rd degree AV block; (20)
    - Indications for dual chamber pacing; (18)
    - Anti-tachycardia pacing will be an important part of treatment. (18)
  - Patient is diagnosed with Non-Ischemic Cardiomyopathy, with an ejection fraction of less than or equal to 35%, and an NYHA class 2 or 3, and ANY of the following:
    - Documented sinus node disease with bradycardia that may need pacing; (20)
    - 2nd or 3rd degree AV block; (20)
    - Indications for dual chamber pacing; (20)
    - Anti-tachycardia pacing will be an important part of treatment. (20)
  - Patient is diagnosed with Right Ventricular Dysplasia and ANY of the following: Is less than 50 years old, has a previous history of cardiac arrest or ventricular tachycardia with hemodynamic compromise, has left ventricular involvement, ARVC2 or ARVC5 positive or increased QRS dispersion plus EITHER:
• Documented sinus node disease with bradycardia that may need pacing; (16)
• 2nd or 3rd degree AV block; (16)
• Indications for dual chamber pacing; (16)
• Anti-tachycardia pacing will be an important part of treatment. (16)

Dual Chamber ICD utilization for the secondary prevention of Sudden Cardiac Death may be reasonable and appropriate when the patient's medical record demonstrates ANY of the following:

- Patient is diagnosed with Brugada Syndrome with documented ventricular tachycardia or syncope/ventricular tachycardia while on beta blocker and ANY of the following:
  • Documented sinus node disease with bradycardia that may need pacing; (20)
  • 2nd or 3rd degree AV block; (20)
  • Indications for dual chamber pacing; (20)
  • Anti-tachycardia pacing will be an important part of treatment. (20)
- Patient is a cardiac arrest survivor from ventricular tachycardia/fibrillation with no reversible cause identified and ANY of the following:
  • Documented sinus node disease with bradycardia that may need pacing; (20)
  • 2nd or 3rd degree AV block; (20)
  • Indications for dual chamber pacing; (20)
  • Anti-tachycardia pacing will be an important part of treatment. (20)
- Patient has sustained spontaneous ventricular tachycardia and ANY of the following:
  • Documented sinus node disease with bradycardia that may need pacing; (20)
  • 2nd or 3rd degree AV block; (20)
  • Indications for dual chamber pacing; (20)
Anti-tachycardia pacing will be an important part of treatment. (20)

Automatic Implantable Cardiac Defibrillator – Removal or Replacement (ICD)

The removal of an Implantable Cardiac Defibrillator may be reasonable and appropriate when the patient's medical record demonstrates any of the following:

- The device was implanted more than a year ago, and replacement is needed earlier than replacement interval (less than 6 years) and ANY of the following:
  - Leakage of other cardiac devices/implants; (24)
  - Device related pain; (22)
  - Lead fracture; (24)
  - Implantation related of chronic infection; (21)
  - Erosion of device through skin (21)
- This is an upgrade of a single chamber to a dual chamber defibrillator (22)
- Device is at the end of its life (implant 6 or more years ago) (24)
- Device was implanted 6 months to a year ago (24)
- Initial encounter (24)
- Subsequent encounter (24)

The lead replacement or add on of an Implantable Cardiac Defibrillator may be reasonable and appropriate when the device is at the end of its life and replacement will be with the same device or lead. (23)

The replacement of an Implantable Cardiac Defibrillator due to device error or battery may be reasonable and appropriate when the medical record demonstrates ANY of the following:
The device is at the end of its life and replacement will be with the same device battery. (23)

The lead replacement or add on of an Implantable Cardiac Defibrillator due to patient complication may be reasonable and appropriate when the device was implanted more than a year ago and ANY of the following:

- Leakage of other cardiac devices/implants;
- Device related pain; (22)
- Lead fracture;
- Implantation related of chronic infection; (21)
- Erosion of device through skin (21)

The lead replacement or add on of an Implantable Cardiac Defibrillator due to patient complication may be reasonable and appropriate in the instance of excessive external manipulation. (22)

Repositioning of an Implantable Cardiac Defibrillator may be reasonable and appropriate when the medical record demonstrates EITHER of the following:

- Excessive external manipulation; (22)
- Device was implanted more than and year ago and ANY of the following:
  - Leakage of other cardiac devices/implants;
  - Device related pain;
  - Lead fracture;
  - Implantation related of chronic infection;
  - Erosion of device through skin
REFERENCES:


18. HRS/ACC/AHA Expert Consensus Statement on the Use of Implantable Cardioverter-Defibrillator Therapy in Patients Who Are Not Included or Not Well Represented in Clinical Trials. 2014 Heart Rhythm Society; American College of Cardiology Foundation; and American Heart Association, Inc.


23. Damir Erkapic, Johannes Sperzel, Sascha Stiller, Ulf Meltendorf, Johann Merani, Karl Wegscheider, Burkhard Hügl, for the INSURE Investigators; Long-term benefit of implantable cardioverter/defibrillator therapy after elective device replacement: results of the INCidence free SURvival after ICD RePlacement (INSURE) trial—a prospective multicentre study, European Heart Journal, Volume 34, Issue 2, 7 January 2013, Pages 130-137, https://doi.org/10.1093/eurheartj/ehs177

Cardiac Electrophysiology Studies

Utilization of a cardiac electrophysiology studies (EPS) may be medically appropriate and supported by evidence to improve patient outcomes for the following indications.

Evaluation of Atrioventricular (AV) block may be reasonable and appropriate when the patient's medical record demonstrates that catheter ablation is planned as well as ANY of the following:
- Suspected site of AV block is within the bundle of His (Intra-Hisian); and ANY of the following:
  - Syncope is present; (32)
  - Near syncopal episodes are reported; (32)
  - Transient lightheadedness is reported; (32)
  - Severe fatigue is present; (32)
  - Palpitations are reported. (32)
- Suspected site of AV block is distal or below the bundle of His (Infra-Hisian); and ANY of the following:
  - Syncope is present; (35)
  - Near syncopal episodes are reported; (35)
  - Transient lightheadedness is reported; (35)
  - Severe fatigue is present; (35)
  - Palpitations are reported. (35)
- Complete AV Block is present; and EITHER of the following:
  - Recurrent symptoms despite therapy and pacemaker placement, study to aid in identification of other arrhythmias as the cause of symptoms.
  - Study to aid in understanding the block or conduction delay with respect to site, mechanism or response to therapy in order to assess the patient's prognosis. (35)
- Second degree AV block; and EITHER of the following:
  - Recurrent symptoms despite therapy and pacemaker placement, study to aid in identification of other arrhythmias as the cause of symptoms; (35)
  - Study to aid in understanding the block or conduction delay with respect to site, mechanism or response to therapy in order to assess the patient's prognosis. (35)
Pseudo AV block; (35)

Prior EKG illustrating a bundle branch block and this study will aid in understanding the block or conduction delay with respect to site, mechanism or response to therapy in order to assess the patient’s prognosis; (32)

Other intraventricular conduction delay is present and this study will aid in understanding the block or conduction delay with respect to site, mechanism or response to therapy in order to assess the patient’s prognosis. (35)

Evaluation of atrial fibrillation or atrial flutter may be reasonable and appropriate when the patient’s medical record demonstrates that catheter ablation is planned as well as **EITHER** of the following:

- Patient has atrial fibrillation or atrial flutter that did not terminate spontaneously within seven (7) days of onset or within forty-eight (48) hours or less of electrical or pharmacological cardioversion as well as having had poor response or was intolerant of at least one type of antiarrhythmic medication. (24)
- Patient has atrial fibrillation or atrial flutter that terminated spontaneously within seven (7) days of onset or within forty-eight (48) hours or less of electrical or pharmacological cardioversion; and **ANY** of the following:
  - Patient is compliant with drug therapy but concern exists for proarrhythmia or adverse effects on the sinus node or atrioventricular conduction; (24)
  - Patient has had a poor response or is intolerant of at least one type of antiarrhythmic medication, drug affects may reveal distinct QT abnormality; and **ANY** of the following:
    - EPS is needed to identify proarrhythmic effect of a drug while patient is experiencing ventricular tachycardia or is at risk for cardiac arrest and prior EKG illustrates a prolonged QT interval; (24)
    - Patient is experiencing near syncopal episodes; and **EITHER** of the following:
      - EKG illustrates equivocal abnormality of QT interval duration; (24)
- EKG illustrates equivocal abnormality of TU wave configuration. (24)
- Patient is experiencing transient lightheadedness; and EITHER of the following:
  - EKG illustrates equivocal abnormality of QT interval duration; (24)
  - EKG illustrates equivocal abnormality of TU wave configuration. (24)
- Patient is experiencing severe fatigue; and EITHER of the following:
  - EKG illustrates equivocal abnormality of QT interval duration; (24)
  - EKG illustrates equivocal abnormality of TU wave configuration. (24)
  - Patient is experiencing syncopal episodes; and EITHER of the following:
    - EKG illustrates equivocal abnormality of QT interval duration; (24)
    - EKG illustrates equivocal abnormality of TU wave configuration.

Evaluation of unexplained syncope or palpitations thought to be of cardiac origin may be reasonable and appropriate when the patient's medical record demonstrates that catheter ablation is planned as well as ANY of the following:
- Palpitations present alone or in the presence of an examination revealing discordance between symptoms and EKG findings which are suggestive of an arrhythmia with or without a structurally abnormal heart; (25,31)
- Palpitations preceding a syncopal episode; (25,31)
- Recurrent syncopal episodes with a patient who has a structurally normal heart and has had a negative head-up tilt test. (31)

Evaluation of supraventricular tachycardia (SVT) may be reasonable and appropriate when the patient's medical record demonstrates that catheter ablation is planned as well as ANY of the following:
- EPS is needed to identify proarrhythmic effects of a drug while patient is experiencing SVT or is at risk of cardiac arrest; (25)
o Patient is diagnosed with Wolff-Parkinson-White syndrome (WPW) or other atrioventricular reentrant/reciprocating tachycardia (AVRT); and **ANY** of the following: (25)
  - EPS is needed to evaluate the patient for ablation of an accessory pathway; (28, 25)
  - EPS is needed to understand the properties of an accessory pathway and normal conduction system for determination of appropriate therapy. (25)
  - Prior EKG illustrates a narrow QRS complex and patient is hemodynamically stable; (25)
  - Prior EKG illustrates a wide QRS complex, greater than 120ms with a bundle branch block. (25)

o Patient is diagnosed with atrioventricular nodal reentrant tachycardia (AVNRT); and **EITHER** of the following:
  - Prior EKG illustrates a narrow QRS complex and patient is hemodynamically stable; (25)
  - Prior EKG illustrates a wide QRS complex, greater than 120ms with a bundle branch block. (25)

o Clinical evaluation indicates that drug effects may reveal a distinct QT abnormality; and **ANY** of the following:
  - Patient is experiencing syncopal episodes; and **EITHER** of the following:
    - EKG illustrates equivocal abnormality of QT interval duration; (25)
    - EKG illustrates equivocal abnormality of TU wave configuration. (25)
  - Patient is experiencing near syncopal episodes; and **EITHER** of the following:
    - EKG illustrates equivocal abnormality of QT interval duration; (25)
    - EKG illustrates equivocal abnormality of TU wave configuration. (25)
  - Patient is experiencing transient lightheadedness; and **EITHER** of the following:
    - EKG illustrates equivocal abnormality of QT interval duration; (25)
    - EKG illustrates equivocal abnormality of TU wave configuration. (25)
  - Patient is experiencing severe fatigue; and **EITHER** of the following:
    - EKG illustrates equivocal abnormality of QT interval duration; (25)
• EKG illustrates equivocal abnormality of TU wave configuration. (25)

Evaluation of ventricular tachycardia (VT) may be reasonable and appropriate when the patient's medical record demonstrates ANY of the following:

- EPS is needed to assess candidacy for catheter ablation; and ANY of the following:
  - Prior ambulatory EKG illustrates VT; (29)
  - Premature ventricular complexes (PVC) are present with severe symptoms; (29)
  - Ventricular couplets are present with severe symptoms; (34)
  - Spontaneous sustained VT has occurred. (29)
- Patient has dilated cardiomyopathy. (29)
- LVEF is less than 35%; and ANY of the following:
  - Patient is status post myocardial infarction (MI) with non-sustained VT on ambulatory EKG; (33)
  - EPS is needed to guide therapy in a patient with inducible VT; (30)
  - EPS is needed to further evaluate risk of VT. (30)

Evaluation of sinus node function may be reasonable and appropriate when the patient's medical record demonstrates that catheter ablation is planned as well as ANY of the following:

- Clinical examination reveals discordance of symptoms with EKG findings suggestive of arrhythmia; and ANY of the following:
  - Patient is experiencing syncopal episodes; (35)
  - Patient is experiencing near syncopal episodes; (35)
  - Patient is experiencing transient lightheadedness; (35)
  - Patient is experiencing severe fatigue; (35)
  - Patient is experiencing palpitations. (35)
- EPS study may aid in selecting the most appropriate pacing modality;
- EPS may aid in selecting the best therapeutic option; and ANY of the following:
  - Suspected cause of symptoms is intrinsic disease within the sinus node; (35)
  - Suspected cause of symptoms is an abnormality in the autonomic nervous system; (35)
  - Suspected cause of symptoms is effects of drugs. (35)
- EPS study may aid in identifying other arrhythmias as the cause of symptoms when sinus node bradycardia is present. (35)
REFERENCES:

6. AHA/ACCF Scientific Statement on the Evaluation of Syncope From the American Heart Association Councils on Clinical Cardiology, Cardiovascular Nursing, Cardiovascular Disease in the Young, and Stroke, and the Quality of Care and Outcomes Research Interdisciplinary Working Group; and the American College of Cardiology Foundation In Collaboration With the Heart Rhythm Society. American Heart Association Science Advisory and Coordinating Committee on August 3, 2005
7. LLOYD A. RUNSER, MD, MPH; ROBERT L. GAUER, MD; and ALEX HOUSER, DO, Syncope: Evaluation and Differential Diagnosis. 2017 American Academy of Family Physician.
Cardiac Pacemaker Device

Utilization of a cardiac pacemaker may be medically appropriate and supported by evidence to improve patient outcomes for the following indications.

Removal of a cardiac pacemaker may be reasonable and appropriate when the patient's medical record demonstrates EITHER of the following:
  - Device is at the end of its lifespan, implantation of device was greater than ten (10) years ago;
  - Device replacement is required, implantation greater than one (1) year ago but less than ten (10) years ago; and ANY of the following:
    - Device migration has occurred, excessive external device manipulation;
    - Complication, implantation related or chronic infection;
    - Complication, erosion of device through the skin;
    - Complication, device related pain;
    - Complication, leakage of other cardiac device or implant.

Lead replacement may be reasonable and appropriate when the replacement is for the same device.

Initial placement of a dual chamber pacemaker for 2nd degree heart block may be reasonable and appropriate when the patient's medical record demonstrates ANY of the following:
  - Diagnosis of symptomatic bradycardia; (45)
  - Patient develops 2nd degree block during exercise in the absence of ischemia; (45)
  - Patient has 2nd degree infra-hisian block illustrated on electrophysiology study (EPS); (45)
  - Patient has symptoms of pacemaker syndrome; (45,46)
  - Patient has Type 2, 2nd degree bock with narrow QRST. (45)
Initial placement of a dual chamber pacemaker for 3rd degree heart block may be reasonable and appropriate when the patient's medical record demonstrates ANY of the following:

- Diagnosis of symptomatic bradycardia; (45)
- Patient requires drugs which induce symptomatic bradycardia; (45)
- There are documented pauses in the patient's cardiac rhythm of greater than or equal to three (3) seconds; (45)
- Patient has a documented heart rate of less than forty (40) while awake; (45)
- Patient is undergoing an AV node ablation that will result in a complete AV block; (45)
- Patient has a post-operative AV block that is not expected to resolve;
- Patient is asymptomatic with left ventricular dysfunction; (45)
- Patient develops complete AV block during exercise in the absence of ischemia; (45)
- Patient has a neuromuscular disease associated with AV block. (45)

Initial placement of a dual chamber pacemaker for AV block due to other causes may be reasonable and appropriate when the patient's medical record demonstrates ANY of the following:

- Patient underwent a cardiac transplant and has persistent, inappropriate symptomatic bradycardia; (45)
- Patient has symptomatic bradycardia or pauses in cardiac rhythm of greater than or equal to three (3) seconds with sleep apnea syndrome; (45)
- Patient has cardiac sarcoidosis with conduction system abnormality; (45,46)
- Patient has hypertrophic cardiomyopathy with sinus node disease or AV block and is medically refractory with left ventricular outflow tract obstruction; (45)
- Patient has a neuromuscular disease and conduction system abnormality; (45)
- Patient has congenital heart disease and a conduction system abnormality. (45)

Initial placement of a dual chamber device for AV block with hypersensitive carotid and/or neurocardiogenic syncope may be reasonable and appropriate when the patient's medical record demonstrates EITHER of the following:
o Patient has recurrent syncopal episodes caused by spontaneously occurring carotid sinus stimulation with a greater than or equal to three (3) second pause; (45)
o Patient has syncope without clear provocation of carotid stimulation but known carotid sinus sensitivity. (45)

Initial placement of a dual chamber pacemaker for AV block with pace termination of tachycardia may be reasonable and appropriate when the patient’s medical record demonstrates that the patient has symptomatic, recurrent SVT that is reproducibly terminated by pacing where drugs or ablation have failed. (45)

Initial placement of a dual chamber pacemaker for AV block with pacing to prevent tachycardia may be reasonable and appropriate when the patient’s medical record demonstrates EITHER of the following:
  o Patient has sustained pause dependent VT; (45)
  o Patient is high risk with long QT syndrome. (45)

Initial placement of a dual chamber pacemaker for Bi and Tri fascicular AV block may be reasonable and appropriate when the patient’s medical record demonstrates ANY of the following:
  o Patient has advanced 2nd degree or intermittent 3rd degree block; (45)
  o Patient has Type 2, 2nd degree block; (45)
  o Patient has alternating bundle branch block; (45)
  o Patient is experiencing syncopal episodes and all other causes have been ruled out; (45)
  o EPS illustrates incidental long His ventricle interval of greater than 100msec; (45)
  o EPS illustrates incidental infra-hisian block. (45)

Initial placement of dual chamber pacemaker post myocardial infarction (MI) for AV block may be reasonable and appropriate when the patient’s medical record demonstrates ANY of the following:
  o Patient has a persistent 2nd degree AV block with alternating bundle branch block or 3rd degree block; (45)
  o Patient has transient advanced 2nd or 3rd degree block; (45)
o Patient has persistent and symptomatic second or 3rd degree block. (45)

Initial placement of a dual chamber pacemaker for sinus node disease may be reasonable and appropriate when the patient's medical record demonstrates ANY of the following:

o Patient has documented symptomatic bradycardia or cardiac rhythm pauses greater than or equal to three (3) seconds; (45)

o Patient has chronotropic incompetence; (45)

o Patient has symptomatic bradycardia caused by drug therapy which is required, Tachy Brady Syndrome; (45)

o Patient has documented sinus node disease with syncope of unknown origin; (45)

o Patient has a heart rate of less than forty (40) while aware with symptoms that are not clearly linked. (45)

Initial placement of a single chamber pacemaker for 2nd or 3rd degree AV block may be reasonable and appropriate when the patient's medical record demonstrates ANY of the following:

o Patient has documented symptomatic bradycardia or cardiac rhythm pauses greater than or equal to three (3) seconds; (45)

o Patient has chronotropic incompetence; (45)

o Patient has symptomatic bradycardia caused by drug therapy which is required, Tachy Brady Syndrome; (45)

o Patient has documented sinus node disease with syncope of unknown origin; (45)

o Patient has a heart rate of less than forty (40) while aware with symptoms that are not clearly linked. (45)

Initial placement of a single chamber pacemaker for AV block due to other causes may be reasonable and appropriate when the patient's medical record demonstrates ANY of the following:

o Patient has persistent inappropriate or symptomatic bradycardia post cardiac transplant; (45)

o Patient has sleep apnea syndrome with symptomatic bradycardia or pauses in cardiac rhythm or greater than or equal to three (3) seconds; (45)

o Patient has cardiac sarcoidosis with conduction system abnormalities; (45,46)
Patient has hypertrophic cardiomyopathy with sinus node disease or AV block and is symptomatic with left ventricular outflow tract obstruction that is refractory to medical management; (45)

Patient has a neuromuscular disease with conduction system abnormalities; (45,46)

Patient has congenital heart disease with conduction system abnormalities. (45)

Initial placement of a single chamber pacemaker for AV block with hypersensitive carotid and/or neurocardiogenic syncope may be reasonable and appropriate when the patient's medical record demonstrates EITHER of the following:

- Patient is experiencing recurrent syncope caused by spontaneously occurring carotid sinus stimulation with a pause in cardiac rhythm of greater than three (3) seconds; (45)
- Patient is experiencing syncopal episodes without clear provocation of carotid stimulation but with known carotid sinus sensitivity. (45)

Initial placement of a single chamber pacemaker for AV block with pace termination of tachycardia may be reasonable and appropriate when the patient's medical record demonstrates symptomatic recurrent SVT that has failed drug or ablation therapy but is reproducibly terminated by pacing.

Initial placement of a single chamber pacemaker for Bi and Tri fascicular AV block may be reasonable and appropriate when the patient's medical record demonstrates ANY of the following:

- Patient has advanced 2nd degree or intermittent 3rd degree block; (45)
- Patient has Type 2, 2nd degree block; (45,46)
- Patient has alternating bundle branch block; (45)
- Patient is experiencing syncopal episodes and all other causes have been ruled out; (45)
- EPS illustrates incidental long His ventricle interval of greater than 100msec; (45)
- EPS illustrates incidental infra-hisian block. (45)

Initial placement of single chamber pacemaker post myocardial infarction (MI) for AV block may be reasonable and appropriate when the patient's medical record demonstrates ANY of the following:
- Patient has a persistent 2nd degree AV block with alternating bundle branch block or 3rd degree block; (45)
- Patient has transient advanced 2nd or 3rd degree block; (45)
- Patient has persistent and symptomatic 2nd or 3rd degree block. (45)

Initial placement of a single chamber pacemaker for sinus node disease may be reasonable and appropriate when the patient’s medical record demonstrates **ANY** of the following:
- Patient has documented symptomatic bradycardia or cardiac rhythm pauses greater than or equal to three (3) seconds; (45)
- Patient has chronotropic incompetence; (45)
- Patient has symptomatic bradycardia caused by drug therapy which is required, Tachy Brady Syndrome; (45)
- Patient has documented sinus node disease with syncope of unknown origin; (45)
- Patient has a heart rate of less than forty (40) while aware with symptoms that are not clearly linked. (45)

Replacement of a pacemaker due to device battery error messaging may be reasonable and appropriate when the patient’s medical record demonstrates the current device will be replaced with the same type of device and the device is at the end of its lifespan.

Replacement of a pacemaker due to complication may be reasonable and appropriate when the patient’s medical record demonstrates **EITHER** of the following:
- Implantation of device was more than a year ago; and **ANY** of the following:
  - Complication, lead fracture;
  - Complication, implantation related chronic infection;
  - Complication, erosion of device through the skin;
  - Complication, device related pain.
- Device migration has occurred, excessive external device manipulation.
REFERENCES:


Cardiac Positron Emission Test

PET scan of the heart may be medically appropriate and supported by evidence to improve patient outcomes for the following indications.

PET scan of the heart may be reasonable and appropriate for a patient who has Coronary Artery Disease (CAD), symptoms are present but stable, and the requested study is for follow-up when the patient's medical record demonstrates BOTH of the following:

- Over 2 years have passed since last myocardial perfusion scan;
- Body Mass Index (BMI) is greater than 35.

PET scan of the heart may be reasonable and appropriate for a patient who requires myocardial viability assessment and the patient's medical record demonstrates EITHER of the following:

- Abnormal myocardial perfusion scan (indeterminate for scar vs. hibernating myocardium) with compromised left ventricular function; (57)
- Documented coronary artery stenosis illustrated on previous cardiac catheterization; and BOTH of the following:
  - Documented compromised left ventricular function (finding indeterminate for scar vs. hibernating myocardium); (58)
  - Anatomically bypassable vessels in area(s) of myocardial dysfunction. (58)

PET scan of the heart may be reasonable and appropriate for an asymptomatic patient who requires a Stress-Rest myocardial perfusion scan for detection of CAD when the patient's medical record demonstrates the following:

- BMI is greater than 35 with no myocardial perfusion scan in the last 2 years which illustrated a normal result; and EITHER of the following:
  - Prior history of CAD;
At least **TWO** of the following:

- Male over 50 years of age;
- Diabetes Mellitus;
- Hypertension;
- Hypercholesterolemia;
- Family history of CAD;
- History of tobacco use;
- Recent ECG, which was uninterpretable or illustrates significant changes.

PET scan of the heart may be reasonable and appropriate for an asymptomatic patient who requires a Stress-Rest myocardial perfusion scan for detection of CAD in the presence of a new diagnosis of ventricular tachycardia when the patient’s medical record demonstrates the following:

- BMI is greater than 35; and **EITHER** of the following:
  - Prior history of CAD;
  - At least **TWO** of the following:
    - Male over 50 years of age;
    - Diabetes Mellitus;
    - Hypertension;
    - Hypercholesterolemia;
    - Family history of CAD;
    - History of tobacco use;
    - Recent ECG, which was uninterpretable or illustrates significant changes.

PET scan of the heart may be reasonable and appropriate for an asymptomatic patient who requires a Stress-Rest myocardial perfusion scan for detection of CAD
in the presence of a new diagnosis of atrial fibrillation when the patient’s medical record demonstrates the following:

- BMI is greater than 35; and **EITHER** of the following:
  - Prior history of CAD;
  - At least **TWO** of the following:
    - Male over 50 years of age;
    - Diabetes Mellitus;
    - Hypertension;
    - Hypercholesterolemia;
    - Family history of CAD;
    - History of tobacco use;
    - Recent ECG, which was uninterpretable or illustrates significant changes.

PET scan of the heart may be reasonable and appropriate for a Stress-Rest myocardial perfusion scan for detection of CAD in the presence of a new onset heart failure of systolic dysfunction when the patient’s medical record demonstrates the **ALL** of the following:

- BMI is greater than 35;
- No cardiac catheterization is planned;
- No prior CAD evaluation;
- **ANY** of the following:
  - Male over 50 years of age;
  - Diabetes Mellitus;
  - Hypertension;
  - Hypercholesterolemia;
  - Family history of CAD;
  - History of tobacco use;
Recent ECG, which was uninterpretable or illustrates significant changes.

PET scan of the heart may be reasonable and appropriate for a Stress-Rest myocardial perfusion scan for the evaluation of chest pain when the patient's medical record demonstrates BOTH of the following:

- BMI is greater than 35 with no myocardial perfusion scan performed in the last 2 years which illustrated a normal result;
- ANY of the following:
  - Male over 50 years of age;
  - Diabetes Mellitus;
  - Hypertension;
  - Hypercholesterolemia;
  - Family history of CAD;
  - History of tobacco use;
  - Recent ECG, which was uninterpretable or illustrates significant changes.

PET scan of the heart may be reasonable and appropriate for a Stress-Rest myocardial perfusion scan for the evaluation of a patient with known CAD and worsening symptoms when the patient's medical record demonstrates the following:

- BMI is greater than 35; and EITHER of the following:
  - Revascularization procedure was performed since last myocardial scan;
  - Over 1 year has passed since last myocardial perfusion scan.

PET scan of the heart may be reasonable and appropriate for a Stress-Rest myocardial perfusion scan for the evaluation of a patient with an intermediate Duke Treadmill score when the patient's medical record demonstrates the following:

- BMI is greater than 35; and EITHER of the following:
  - Unable to exercise;
  - ANY of the following:
    - Male over 50 years of age;
• Diabetes Mellitus;
• Hypertension;
• Hypercholesterolemia;
• Family history of CAD;
• History of tobacco use;
• Recent ECG, which was uninterpretable or illustrates significant changes.

PET scan of the heart may be reasonable and appropriate for a Stress-Rest myocardial perfusion scan for evaluation prior to surgery when the patient's medical record demonstrates **ALL** of the following:
  - BMI is greater than 35;
  - Over one year has passed since last cardiac test or intervention;
  - Intermediate or high risk for surgery;
  - Unable to perform adequate exercise.

PET scan of the heart may be reasonable and appropriate for a Stress-Rest myocardial perfusion scan as part of the patient's risk assessment after acute coronary syndrome when the patient's medical record demonstrates **BOTH** of the following:
  - BMI is greater than 35;
  - No cardiac catheterization is planned.

PET scan of the heart may be reasonable and appropriate for a Stress-Rest myocardial perfusion scan as part of the patient's risk assessment after coronary artery bypass graft (CABG) or percutaneous coronary intervention (PCI) when the patient's medical record demonstrates the following:
  - BMI is greater than 35; and **ANY** of the following:
    - Patient is being evaluated for new onset chest pain;
    - Study is being performed for assessment of an asymptomatic patient 5 years post CABG;
- Study is being performed for assessment of an asymptomatic patient 1-year post PCI.
REFERENCES:


7. Patrycja Galazka, MD1; Marcelo F. Di Carli, MD, Cardiac PET/CT and Prognosis. Cardiovascular Imaging Program, Departments of Radiology and Medicine, the Division of Nuclear Medicine and Molecular Imaging, Department of Radiology, and the Cardiovascular Division, Department of Medicine, Brigham and Women’s Hospital, Harvard Medical School, Boston, MA, USA, 18 November 2016

Cardiac Resynchronization Therapy

Utilization of a cardiac resynchronization therapy may be medically appropriate and supported by evidence to improve patient outcomes for the following indications.

Cardiac Resynchronization Therapy: Defibrillator (CRT-D)

Removal of a CRT-D may be reasonable and appropriate when the patient’s medical record demonstrates **EITHER** of the following:

- Device is at the end of its lifespan, implantation of device was greater than or equal to five (5) years ago; and **EITHER** of the following:
  - Request is for an upgrade of a single or dual chamber defibrillator to a device with biventricular pacing capability;
  - Request is for replacement with the same biventricular device.
- Device replacement is required, implantation greater than one (1) year ago and earlier than the normal replacement interval of five (5) years ago; and **ANY** of the following:
  - Device migration has occurred, excessive external device manipulation;
  - Complication, implantation related or chronic infection;
  - Complication, erosion of device through the skin;
  - Complication, device related pain;
  - Complication, leakage of other cardiac device or implant.

Lead replacement may be reasonable may be reasonable and appropriate when the patient’s medical record demonstrates **EITHER** of the following:

- Replacement with the same device due to end of lifespan, implantation of device was greater than or equal to five (5) years ago with or without an upgrade of a single or dual chamber defibrillator to a device with biventricular pacing capability.
- Replacement is needed earlier than replacement interval, implantation was less than five (5) years ago with implantation more than a year ago.
and this request is for an upgrade of a single or dual chamber defibrillator to a device with biventricular pacing capability. (59)

Replacement of CRT-D due to device battery error messaging may be reasonable and appropriate when the patient’s medical record demonstrates EITHER of the following:
  o Replacement of the current battery with the same battery; and EITHER of the following:
    ▪ Device is at the end of its lifespan, implantation of device was greater than or equal to five (5) years ago with or without an upgrade of a single or dual chamber defibrillator to a device with biventricular pacing capability;
    ▪ Device replacement is required, implantation greater than one (1) year ago and earlier than the normal replacement interval of five (5) years ago with or without an upgrade of a single or dual chamber defibrillator to a device with biventricular pacing capability.

Replacement of CRT-D due to complication may be reasonable and appropriate when the patient’s medical record demonstrates EITHER of the following:
  o Implantation of device was more than a year ago; and ANY of the following:
    ▪ Complication, lead fracture; (59)
    ▪ Complication, implantation related chronic infection; (59)
    ▪ Complication, erosion of device through the skin; (59)
    ▪ Complication, device related pain. (59)
    ▪ Complication, leakage of other cardiac device or implant. (59)
    ▪ Device migration has occurred, excessive external device manipulation. (59)

Upgrade from a single chamber device to dual chamber device with biventricular pacing capability may be reasonable and appropriate when the patient’s medical record demonstrates ALL of the following:
  o Current device was implanted more than a year ago; (59)
  o Patient has QRS complex width of greater than 120msec; (60)
  o Patient’s condition requires pacing, such as chronotropic incompetence; (60)
  o Patient is exhibiting progression of congestive heart failure to New York Heart Association functional class of II, III or IV. (60)
Cardiac Resynchronization Therapy: Pacemaker (CRT-P)

Removal of a CRT-P may be reasonable and appropriate when the patient's medical record demonstrates EITHER of the following:

- Device is at the end of its lifespan, implantation of device was greater than or equal to five (5) years ago; and EITHER of the following:
  - Request is for an upgrade of a single or dual chamber defibrillator to a device with biventricular pacing capability;
  - Request is for replacement with the same biventricular device.
- Device replacement is required, implantation greater than one (1) year ago and earlier than the normal replacement interval of five (5) years ago; and ANY of the following:
  - Device migration has occurred, excessive external device manipulation;
  - Complication, implantation related or chronic infection;
  - Complication, erosion of device through the skin;
  - Complication, device related pain;
  - Complication, leakage of other cardiac device or implant.

Lead replacement may be reasonable may be reasonable and appropriate when the patient’s medical record demonstrates EITHER of the following:

- Replacement with the same device due to end of lifespan, implantation of device was greater than or equal to five (5) years ago with or without an upgrade of a single or dual chamber defibrillator to a device with biventricular pacing capability.
- Replacement is needed earlier than replacement interval, implantation was less than five (5) years ago with implantation more than a year ago and this request is for an upgrade of a single or dual chamber defibrillator to a device with biventricular pacing capability. (59)
Replacement of CRT-P due to device battery error messaging may be reasonable and appropriate when the patient's medical record demonstrates **EITHER** of the following:

- Replacement of the current battery with the same battery; and **EITHER** of the following:
  - Device is at the end of its lifespan, implantation of device was greater than or equal to five (5) years ago with or without an upgrade of a single or dual chamber defibrillator to a device with biventricular pacing capability;
  - Device replacement is required, implantation greater than one (1) year ago and earlier than the normal replacement interval of five (5) years ago with or without an upgrade of a single or dual chamber defibrillator to a device with biventricular pacing capability.

Replacement of CRT-P due to complication may be reasonable and appropriate when the patient's medical record demonstrates **EITHER** of the following:

- Implantation of device was more than a year ago; and **ANY** of the following:
  - Complication, lead fracture; (59)
  - Complication, implantation related chronic infection; (59)
  - Complication, erosion of device through the skin; (59)
  - Complication, device related pain. (59)
  - Complication, leakage of other cardiac device or implant. (59)
- Device migration has occurred, excessive external device manipulation. (59)

Upgrade from a single chamber device to dual chamber device with biventricular pacing capability may be reasonable and appropriate when the patient's medical record demonstrates **ALL** of the following:

- Current device was implanted more than a year ago;
- Patient has QRS complex width of greater than 120msec; (60)
- Patient's condition requires pacing, such as chronotropic incompetence; (60)
- Patient is exhibiting progression of congestive heart failure to New York Heart Association functional class of II, III or IV. (60)
REFERENCES:

Cardiac Single Photon Emission Computerized Tomography

SPECT scan of the heart may be medically appropriate and supported by evidence to improve patient outcomes for the following indications.

Evaluation of a patient who has had acute coronary syndrome (ACS) in the past twelve (12) months but is currently asymptomatic may be reasonable and appropriate when the patient's medical record demonstrates ANY of the following:

- It has been greater than three (3) months since the ACS event and there was no percutaneous coronary intervention (PCI) or coronary artery bypass graft (CABG) performed at the time of the event and ANY of the following:
  - Patient has not had any previous coronary angiography for the evaluation of coronary artery disease (CAD);
  - Patient has had previous CAD evaluation within the last twenty four (24) months which demonstrated clinically significant cardiac stenosis of greater than 50% or a moderate to large stress induced perfusion defect;
  - Patient has had previous CAD evaluation greater than twenty four (24) months ago which demonstrated clinically significant cardiac stenosis of greater than 50% or a moderate or large stress induced perfusion defect or the previous study findings were equivocal.
- Patient had an incomplete vascularization after revascularization with PCI or CABG at the time of the ACS event;
- Patient had a complete vascularization after revascularization with PCI or CABG at the time of the ACS event and the patient has had a PCI more than two (2) years ago or a CABG more than five (5) years ago.

Evaluation of new onset atrial fibrillation with an unclear etiology demonstrated by standard workup in an asymptomatic patient may be reasonable and appropriate when the patient's medical record demonstrates ALL of the following:

- No metabolic or other causes of atrial fibrillation identified; (76)
o Framingham Risk Criteria greater than 10%. (76)

Evaluation of an asymptomatic patient with an elevated troponin level may be reasonable and appropriate when the patient's medical record demonstrates ANY of the following:
  o Patient has a coronary artery calcium (CAC) score of greater than five hundred (500); (76)
  o Patient is experiencing documented atrial fibrillation.

Evaluation of an asymptomatic patient who has had a previous revascularization via PCI or CABG may be reasonable and appropriate when the patient's medical record demonstrates ANY of the following:
  o Patient has had an incomplete revascularization and additional revascularization is feasible;
  o It has been greater than five (5) years since the patient's last CABG; (76)
  o It has been greater than two (2) years since the patient's last PCI.

Evaluation of an asymptomatic patient who has had a current diagnosis of CAD but no previous PCI or CABG may be reasonable and appropriate when the patient's medical record demonstrates ANY of the following:
  o Patient has had a previous CAD evaluation with stress echocardiogram which illustrated a severe wall motion abnormality; (76)
  o Patient has had a previous CAD evaluation with heart catheterization or CCTA demonstrating cardiac stenosis of greater than 50% in major vessel less than or equal to twenty four (24) months ago. (76)
  o Patient has had a previous CAD evaluation with cardiac SPECT demonstrating a moderate or large stress induced perfusion defect greater than or equal to twenty four (24) months ago. (76)

Evaluation of an asymptomatic patient who does not have a diagnosis of CAD may be reasonable and appropriate when the patient's medical record demonstrates the following:
  o It has been greater than twenty four (24) months since the patient's last CAD evaluation; and EITHER of the following:
    ▪ Patient is experiencing documented atrial fibrillation;
- Framingham Risk Criteria greater than 20%.\(^{(76)}\)

Evaluation of a patient with a high-risk occupation, i.e., police, firefighter, pilot or bus driver may be reasonable and appropriate when the patient’s medical record demonstrates **EITHER** of the following:
  - Framingham Risk Criteria of greater than 20%;\(^{(76)}\)
  - Framingham Risk Criteria of greater than 10% and patient is currently experiencing documented atrial fibrillation.

Evaluation of cardiac viability via a rest redistribution thallium scan may be reasonable and appropriate when the patient's medical record demonstrates **BOTH** of the following:
  - Patient is eligible for revascularization;\(^{(76)}\)
  - Patient has ischemic cardiomyopathy or myocardial infarct with possibility of viable myocardium. \(^{(76)}\)

Evaluation of worsening chest pain or ischemic equivalent in a patient who has not had PCI or CABG and is currently undergoing medical management of CAD may be reasonable and appropriate when the patient's medical record demonstrates the following:
  - Worsening cardiac symptoms suggesting ischemic equivalent despite treatment with maximal medical management and the patient's last CAD evaluation was within the past twelve (12) months; and **EITHER** of the following:
    - Last CAD testing was via heart catheterization or cardiac computed tomography angiography (CCTA) within the last twelve (12) months demonstrating cardiac stenosis of greater than 20% in any major vessel; \(^{(76)}\)
    - Last CAD testing was via cardiac SPECT or stress echocardiogram within the last twelve (12) months demonstrating an abnormal result of a mild or more severe stress induced perfusion defect or definite stress induced wall motion abnormality and if these findings are unchanged, the ordering physician has no intent to proceed to a heart catheterization. \(^{(76)}\)
Evaluation of worsening cardiac symptoms in a patient who has a diagnosis of CAD and has had a PCI or CABG may be reasonable and appropriate when the patient’s medical record demonstrates the following:

- Worsening cardiac symptoms suggesting cardiac ischemia in a patient who is on maximal medical management; and **ANY** of the following:
  - Heart Catheterization or CCTA was performed within the last twelve (12) months ago with clinically significant findings on last Heart Catheterization or CCTA that could explain patient’s ischemic equivalent symptoms; (76)
  - Heart Catheterization or CCTA was performed greater than twelve (12) months ago or patient has not had previous Heart Catheterization or CCTA have been performed; (76)
  - Cardiac SPECT or Stress Echocardiogram was performed within the last twelve (12) months and illustrated mild to severe stress induced perfusion defect or definite wall motion abnormality and if the perfusion defect or wall motion abnormality defect previous detected has not changed the patient will not undergo a heart cath. (76)

Evaluation of stable chest pain or ischemic equivalent in a patient who has not had PCI or CABG and is currently undergoing medical management of CAD may be reasonable and appropriate when the patient’s medical record demonstrates the following:

- Last CAD testing was within the past twenty four (24) months demonstrating cardiac stenosis of less than 50% in any major vessel or moderate to severe stress induced perfusion defect or wall motion abnormality. (76)

Evaluation of stable chest pain or ischemic equivalent in a patient who has a diagnosis of CAD and has had PCI or CABG may be reasonable and appropriate when the patient’s medical record demonstrates **EITHER** of the following:

- Last CAD evaluation was performed less than or equal to twenty four (24) months ago and demonstrated a lesion in a major coronary vessel or bypass graft with the potential to produce ischemia or a moderate to severe perfusion defect or wall motion abnormality; (76)
- Last CAD evaluation was performed greater than twenty four (24) months ago and if this Cardiac SPECT illustrates a new perfusion defect or increase
in a previous defined perfusion defect, the patient will undergo a heart catheterization. (76)

Evaluation of patient diagnosed with CAD and having high risk findings may be reasonable and appropriate when the patient's medical record demonstrates EITHER of the following:

- Last CAD testing was within the last twenty four (24) months; and ANY of the following:
  - Previous evaluation was via Stress Echocardiogram which demonstrated severe wall motion abnormality;
  - Patient has a history of silent ischemia on previous testing;
  - Previous evaluation was via Heart Catheterization or CCTA and demonstrated cardiac stenosis of greater than 50% in a major vessel. (76)

- Last CAD testing via Stress Echocardiogram was greater than or equal to twenty four (24) months ago and demonstrated severe wall motion abnormality.

Evaluation of a patient with cardiac symptoms who has no known history of CAD and no previous CAD testing in the past twenty four (24) months may be reasonable and appropriate when the patient's medical record demonstrates ANY of the following:

- Male patient 40 years of age or older with typical angina, (substernal chest pain or discomfort that is provoked by exertion or stress and is relieved by rest and/or administration of nitroglycerin);

- Male patient 60 years of age or older with atypical angina, (defined as chest pain that lacks one of the characteristics of typical chest pain, i.e., substernal chest pain or discomfort that is provoked by exertion or stress and is relieved by rest and/or administration of nitroglycerin);

- Female patient 60 years of age or older with typical angina, (substernal chest pain or discomfort that is provoked by exertion or stress and is relieved by rest and/or administration of nitroglycerin); (77)

- Patient with typical angina, (substernal chest pain or discomfort that is provoked by exertion or stress and is relieved by rest and/or administration of nitroglycerin) where a baseline EKG is not interpretable and the patient is unable to exercise on a treadmill. (77)
Evaluation of CAD in an asymptomatic patient who has a new diagnosis of heart failure or left ventricular systolic dysfunction may be reasonable and appropriate when the patient's medical record demonstrates **ANY** of the following:

- Framingham Risk Criteria of between 10 and 20% and patient is currently experiencing documented atrial fibrillation;
- Framingham Risk Criteria of greater than 20%; and **EITHER** of the following:
  - Patient's last CAD evaluation was performed greater than twelve (12) months ago;
  - Patient has not had a previous CAD evaluation.
- Patient has had a previous Heart Catheterization or CCTA was performed within the last twelve (12) months demonstrating cardiac stenosis of greater than 50% in a major vessel.

Cardiac evaluation in the presence of an abnormal or uninterpretable electrocardiogram (ECG) in a patient with no previous PCI or CABG may be reasonable and appropriate when the patient's medical record demonstrates **ANY** of the following:

- Framingham Risk Criteria of greater than 20% in a patient who is asymptomatic with no ischemic equivalent; (76)
- Patient is experiencing an ischemic equivalent with a possible acute coronary syndrome; (76)
- Patient is experiencing ischemic equivalent symptoms chronically and is unable to walk perform a walking treadmill stress test due to physical limitation such as orthopedic or vascular issues, this does not include dyspnea on exertion if it is considered to be the angina equivalent. (76)

Cardiac evaluation in the presence of an equivocal or uninterpretable Stress Echocardiogram in a patient with no previous PCI or CABG may be reasonable and appropriate when the patient's medical record demonstrates **ANY** of the following:

- Patient is experiencing new or worsening cardiac symptoms or ischemic equivalent; (77)
- Patient is experiencing documented atrial fibrillation;
- Patient has a diagnosis of CAD and their last evaluation was greater than two (2) years ago.
Stress-Rest Myocardial Cardiac Risk Assessment in an asymptomatic patient who is post-surgical intervention with CABG or PCI may be reasonable and appropriate when the patient’s medical record demonstrates **ANY** of the following:

- Incomplete revascularization and additional revascularization is feasible;
- It has been greater than five (5) years since the patient has had a CABG; (76)
- It has been greater than two (2) years since the patient has had a PCI;
- There are new changes on the patient’s ECG consistent with ischemia;
- There is new documentation illustrating worsening of left ventricular function; (76)
- Patient has had a PCI of the left main coronary artery in the past.

Pre-operative Cardiac Risk Evaluation of a stable patient may be reasonable and appropriate when the patient’s medical record demonstrates the following:

- The patient is to undergo vascular, open thoracic or open abdominal surgery or organ transplant and it has been greater than one (1) year since the patient has had a previous myocardial perfusion study, diagnostic heart catheterization or myocardial revascularization procedure; and **ANY** of the following: (75)
  - Patient is unable to walk perform a walking treadmill stress test due to physical limitation such as orthopedic or vascular issues, this does not include dyspnea on exertion if it is considered to be the angina equivalent and they have **ANY** of the following documented risk factors:
    - History of CAD;
    - History of prior or compensated congestive heart failure (CHF);
    - History of cerebrovascular disease;
    - History of diabetes mellitus (DM) requiring insulin;
    - History of renal insufficiency with a creatinine level of greater than 2.0.
  - Patient has a recent ECG obtained at rest demonstrating a left bundle branch block (LBBB) or ST-T wave depression greater than 1mm and an exercise tolerance score of less than 4 MET;
  - Patient has an exercise tolerance score of less than 4 MET. (76)

Pre-operative Cardiac Risk Evaluation of a symptomatic or unstable patient may be reasonable and appropriate when the patient’s medical record demonstrates the following:
Patient with active, new or unstable cardiovascular symptoms felt to represent ischemic CAD or CHF who is to undergo vascular, open thoracic or open abdominal surgery or organ transplant; and EITHER of the following:

- Patient is unable to walk perform a walking treadmill stress test due to physical limitation such as orthopedic or vascular issues, this does not include dyspnea on exertion if it is considered to be the angina equivalent;
- Patient has an exercise tolerance score of less than 4 MET.

Evaluation of syncope in a patient with no chest pain and no ischemic equivalent who has had a full evaluation for possible non-cardiac causes, which was indeterminate, may be reasonable and appropriate when the patient’s medical record demonstrates the following:

- Last CAD testing was greater than twelve (12) months ago; and EITHER of the following:
  - Patient does not have a diagnosis of CAD; and ANY of the following:
    - Framingham Risk Score of greater than 20%;
    - Framingham Risk Score of 10-20% and is currently experiencing documented atrial fibrillation
  - Patient has a diagnosis of CAD.

Evaluation of a patient with ventricular tachycardia who is not having any cardiac symptoms but does have a diagnosis of CAD may be reasonable and appropriate when the patient’s medical record demonstrates the EITHER of following:

- Last CAD testing was greater than twelve (12) months ago;
- Last CAD testing via Heart Catheterization or CCTA was less than twelve (12) months ago and demonstrated cardiac stenosis of greater than 50% in a major vessel.

Evaluation of a patient with ventricular tachycardia who is not having any cardiac symptoms but does not have a diagnosis of CAD may be reasonable and appropriate when the patient’s medical record demonstrates the EITHER of following:

- Last CAD testing was greater than twenty four of (24) months ago and the patient has a Framingham Score of greater than 20%;
o Last CAD testing was less than or equal to twenty four (24) months ago; and

EITHER of the following:

 Last CAD testing was via Heart Catheterization or CCTA and demonstrated cardiac stenosis of greater than 50% in a major vessel;
 Last CAD testing was via Stress Echocardiogram and demonstrated severe wall motion abnormality.
REFERENCES:


Coronary Computed Tomography Angiography

Utilization of a cardiac coronary tomography (CT) angiography may be medically appropriate and supported by evidence to improve patient outcomes for the ANY of the following indications.

Evaluation of chest pain may be reasonable and appropriate when the patient's medical record demonstrates ALL of the following:

- No significant arrhythmia or tachycardia (heart rate > 100);
- Recent stress test (treadmill ECG, myocardial perfusion study, or echocardiogram) negative or equivocal;
- At least two (2) of the following risk factors are present:
  - Male over 50;
  - Diabetes mellitus;
  - Hypertension;
  - Hypercholesterolemia;
  - Family history of coronary artery disease;
  - History of tobacco use;
  - Recent ECG, which was uninterpretable or demonstrates significant changes.

Evaluation of shortness of breath with suspected underlying coronary cause may be reasonable and appropriate when the patient's medical record demonstrates ALL of the following:

- No significant arrhythmia or tachycardia (heart rate > 100);
- Recent stress test (treadmill ECG, myocardial perfusion study, or echocardiogram) negative or equivocal;
- At least two (2) of the following risk factors are present:
  - Male over 50;
  - Diabetes mellitus;
  - Hypertension;
  - Hypercholesterolemia;
  - Family history of coronary artery disease;
- History of tobacco use;
- Recent ECG, which was uninterpretable or demonstrates significant changes.

Follow-up study for patient who has had a previous cardiac catheterization, which demonstrated coronary artery stenosis.

Follow-up study for a patient who has had a previous Percutaneous Coronary Intervention (PCI/STENT) procedure within the last three (3) months.
REFERENCES


Echocardiogram

Utilization of an echocardiogram may be medically appropriate and supported by evidence to improve patient outcomes for the following indications.

Transthoracic Echocardiogram (TTE)

Evaluation of an abnormal electrocardiogram (ECG) finding may be reasonable and appropriate when the patient’s medical record demonstrates ANY of the following:

- No prior TTE; and EITHER of the following:
  - Lightheadedness or near loss of consciousness with documented underlying heart disease;
  - Arrhythmia documented by telemetry, Holter monitoring or ECG; (86)
- Prior TTE performed less than twelve (12) months ago; and EITHER of the following:
  - Syncope or near syncope with clinical change;
  - Arrhythmia documented by telemetry, Holter monitoring or ECG with or without clinical change; (91)
- Prior TTE was performed greater than twelve (12) months ago; and EITHER of the following:
  - Syncope or near syncope with clinical change;
  - Arrhythmia documented by telemetry, Holter monitoring or ECG with or without clinical change; (120)
- Syncope or near syncope with occasional atrial or ventricular premature complexes (APC) (VPC) without evidence of heart disease; (119)
- Frequent or exercise induced VPCs. (120)

Evaluation of the aorta or aortic root may be reasonable and appropriate when the patient’s medical record demonstrates ANY of the following:

- Change in symptoms or clinical findings in a patient who has documented ascending aortic disease demonstrated on a previous TTE; (87)
- Known or suspected Marfan’s Syndrome; (115, 133)
Patient is under consideration for surgical intervention to repair a known ascending aorta or aortic valve condition; (93)

Post-surgical evaluation after repair of an ascending aorta or aortic valve condition; (128)

Prior TTE was greater than one (1) year ago and patient has documented ascending aortic disease demonstrated on a previous TTE.

Evaluation of atrial fibrillation may be reasonable and appropriate when the patient's medical record demonstrates ANY of the following:

- No prior TTE; and EITHER of the following:
  - Patient is diagnosed with atrial fibrillation; (121)
  - Patient has sustained or non-sustained atrial fibrillation documented by telemetry, Holter monitor or ECG.
- Prior TTE performed less than twelve (12) months ago; and EITHER of the following:
  - Patient is diagnosed with atrial fibrillation and exhibiting new cardiac symptoms;
  - Patient has sustained or non-sustained atrial fibrillation documented by telemetry, Holter monitor or ECG with a clinical change.
- Prior TTE was performed greater than twelve (12) months ago; and EITHER of the following:
  - Patient has sustained or non-sustained atrial fibrillation documented by telemetry, Holter monitor or ECG;
  - Patient is diagnosed with atrial fibrillation and exhibiting new cardiac symptoms.
- Patient is suspected of having an embolus originating from the heart or has had a transient ischemic attack(s) (TIA) or cerebrovascular accident (CVA). (89,90,94)

Evaluation of dyspnea in a patient with known cardiac disease may be reasonable and appropriate when the patient's medical record demonstrates that non-cardiac causes have been ruled out and ANY of the following:

- No prior TTE; and ANY of the following:
  - Suspicion of congestive heart failure (CHF); (134)
  - Known or suspected diastolic dysfunction; (125)
  - Known valvular heart disease. (122)
Prior TTE was performed greater than twelve (12) months ago; and ANY of the following:
  - Suspicion of congestive heart failure (CHF);
  - Known or suspected diastolic dysfunction;
  - Known valvular heart disease. (122)

Suspected CHF with clinical change/deterioration. (92)

Evaluation of dyspnea in a patient without known cardiac disease may be reasonable and appropriate when the patient's medical record demonstrates that non-cardiac causes have been ruled out and EITHER of the following:
  - No prior TTE and cardiac etiology of dyspnea is suspected based on clinical findings;
  - Prior TTE was performed greater than twelve (12) months ago and cardiac etiology of dyspnea is suspected based on clinical findings.

Evaluation of cardiac rhythm abnormality may be reasonable and appropriate when the patient has ANY of the following:
  - No prior echo and EITHER of the following:
    - Lightheadedness or near loss of consciousness with documented underlying heart disease such as cardiomyopathy, aortic stenosis or congestive heart failure along with heart palpitations. (137)
    - Arrhythmia documented by Telemetry, Holter Monitoring or ECG (i.e. Atrial fibrillation, SVT or VT) (52)
  - Prior echo and clinical change with arrhythmia documented by Telemetry, Holter monitoring or ECG.
  - Prior echo less than twelve (12) months ago and EITHER of the following:
    - Syncope or near syncope (loss of consciousness or near loss of consciousness) with a clinical change and heart palpitations.
    - Clinical change and arrhythmia documented by Telemetry, Holter Monitoring or ECG (i.e. atrial fibrillation, SVT or VT).
  - Prior echo greater than twelve (12) months ago and EITHER of the following:
    - No clinical change and arrhythmia documented by Telemetry, Holter monitoring or ECG i.e. Atrial fibrillation, SVT or VT.
- Clinical change with syncope or near syncope (loss of consciousness or near loss of consciousness).
  - Syncope or near syncope (loss of consciousness or near loss of consciousness) with occasional Atrial Premature Complexes (APC) or Occasional Ventricular Premature Complexes (VPC) without evidence of heart disease. (127)
  - Frequent or exercise-induced Ventricular Premature Complexes (VPC). (113,115)

Evaluation of cardiac structure and function may be reasonable and appropriate when the patient has not had a prior echo and have had abnormal diagnostic testing suggesting a new cardiac abnormality (e.g., abnormal ECG or chest C-ray, or myocardial perfusion imaging study). (137)

Evaluation of cardiac transplantation or left ventricular device (LVAD) candidacy may be reasonable and appropriate when the patient's medical record demonstrates EITHER of the following:
  - If the request is for Initial evaluation for LVAD candidacy. (115)
  - If the request is for follow up for suspected complication or to optimize settings. (115)

Evaluation of cardiac transplantation may be reasonable and appropriate when the patient's medical record demonstrates necessity for echo to assess donor suitability to guide therapy. (95,115)

Evaluation of cardiac transplantation in regards to pre and post-transplant evaluation may be reasonable and appropriate when the patient's medical record demonstrates EITHER of the following:
  - An echo is necessary to determine candidacy of transplant recipient pre-transplant (98, 115)
  - An echo is necessary to monitor for rejection in post-transplant recipient. (97)
Evaluation of cardiomyopathy as follow up for a stable patient may be reasonable and appropriate when the patient has had a prior echo greater than twelve (12) months ago and the medical record demonstrates EITHER of the following:

- Known cardiomyopathy demonstrated by Echo, Cath, CCTA, PET or MUGA. (115)
- The study is for periodic surveillance. (115)

Evaluation of cardiomyopathy as a known condition with change in clinical status or deterioration may be reasonable and appropriate when the diastolic dysfunction has been confirmed in prior imaging (Echo, Stress Echo, MUGA, SPECT or Catheterization) and the medical record demonstrates ANY of the following:

- New symptoms or worsening symptoms attributable to cardiomyopathy (e.g. Physical Limitation, Chest Pain, shortness of breath, lightheadedness, syncope, and peripheral edema) have been found on physical exam. (115)
- Abnormal ECG changes have been recently documented. (115)
- New clinical deterioration of left ventricular function (e.g. change in left ventricular ejection fraction) or myocardial perfusion defect has been documented. (115)

Evaluation of cardiomyopathy as suspected or initial evaluation may be reasonable and appropriate when the patient has had no prior echocardiogram or evaluation and the medical record demonstrates ANY of the following:

- Suspected cardiomyopathy based on clinical findings (96, 115)
- Infiltrated cardiomyopathy: (e.g., heart disease associated with sarcoid, amyloidosis, or hemochromatosis). (96, 115)
- The patient has a first-degree relative with an inherited cardiomyopathy such as hypertrophic cardiomyopathy. (96,99)
Evaluation of cerebrovascular accident (CVA) or systemic embolus may be reasonable and appropriate when the patient’s medical record demonstrates cerebrovascular accident (CVA), transient ischemic attack (TIA) or systemic embolization though to be of cardiac origin. (115)

Evaluation of chest pain may be reasonable and appropriate when the patient’s medical record demonstrates ANY of the following:

- Acute chest pain with suspected myocardial infarction (MI) and non-diagnostic ECG (resting echo during pain). (101,115)
- Evaluation of patient without chest pain but other symptoms of ischemic equivalent or lab markers suggesting ongoing MI. (115)
- Suspected complication of MI such as aortic mitral regurgitation, ventricular septal defect, free-wall rupture/tamponade, shock, right ventricular involvement, HF or thrombus. (115)
- Severe deceleration injury or chest trauma when valve injury, pericardial effusion, cardiac injury are possible or suspected. (115)

Evaluation of Congenital Heart Disease (CHD) in the adult may be reasonable and appropriate when the patient’s medical record demonstrates ANY of the following:

- No prior echo and congenital heart disease is suspected (115, 133)
- Prior echo over one (1) year ago and congenital heart disease is an established diagnosis (with or without a history of prior surgical repair). (115, 133)
- Pt has new or worsening symptoms and congenital heart disease is an established diagnosis (with or without a history of prior surgical repair). (106,133)
Evaluation of congestive heart failure (CHF) prior to surgery may be reasonable and appropriate when the patient's medical record demonstrates that the patient is in NYHA Class I, II, III, or IV heart failure and cardiac surgery is indicated in this patient to treat Heart Failure.  (133)

Evaluation of congestive heart failure (CHF) as follow up in a stable patient may be reasonable and appropriate when the patient's medical record demonstrates documented CHF symptoms and a prior echo has been done greater than twelve (12) months ago.  (132)

Evaluation of congestive heart failure (CHF) with change in clinical statue or deterioration may be reasonable and appropriate when the patient has a history of Patient is in NYHA Class I, II, III, or IV heart failure and the medical record demonstrates ANY of the following:
- New or worsening symptoms attributable to CHF (e.g. Physical limitation, Chest Pain, shortness of breath, lightheadedness, pre-syncope, syncope or peripheral edema) have been found on physical exam.  (105)
- Abnormal ECG changes have been recently documented.  (102, 105)
- New clinical deterioration of left ventricular function (e.g. Change in Left Ventricular Ejection Fraction) or myocardial perfusion defect has been documented.  (105)

Evaluation of congestive heart failure (CHF) as suspected or initial evaluation may be reasonable and appropriate when the patient's medical record demonstrates no prior echo with evaluation of congestive heart failure (CHF) symptoms.  (118)

Evaluation of coronary artery disease (CAD) may be reasonable and appropriate when the patient has known Coronary Artery Disease (CAD) has new symptoms or has prior Myocardial Infarction, and structure/function of the heart is being evaluated.  (104)
Evaluation of edema in known heart disease may be reasonable and appropriate when the patient has new symptoms of edema or change in clinical edema thought to be of cardiac origin medical record demonstrates ANY of the following:

- History of congestive heart failure (CHF) (134)
- History of Restrictive/Constrictive Cardiomyopathy
- History of other cardiomyopathy
- History of Coronary Artery Disease (CAD)
- History of Valvular Disease (134)

Evaluation of edema without known heart disease may be reasonable and appropriate when the patient's medical record demonstrates ANY of the following:

- New peripheral edema and clinical findings suggestive of cardiac origin of edema. (103,130)
- No echo in the last 12 months and chronic peripheral edema as documented from prior workup.
- Prior echo greater than twelve (12) months and chronic peripheral edema.
- Elevated venous pressure.
- Prior echo greater than twelve (12) months and clinical findings suggest cardiac origin of edema.

Evaluation of endocarditis (known or suspected in native or prosthetic valves) when evaluation is needed may be reasonable and appropriate when the patient's medical record demonstrates ANY of the following:

- Endocarditis is a confirmed diagnosis and echo is for follow up after course of therapy or not responding to therapy. (107,133)
- Endocarditis is suspected, a new murmur has developed with positive blood cultures. (107,133)
- Endocarditis is a confirmed diagnosis and there is a change in murmur or clinical status. (107,133)
- Endocarditis is suspected and there is a change in murmur or clinical status with or without positive blood cultures. (107,133)
Evaluation of hypertension may be reasonable and appropriate when the patient's medical record demonstrates ANY of the following:

- Last echo is greater than twelve (12) months ago and ventricular hypertrophy on ECG
- No previous echo, assessment is needed to evaluate cardiac structure and function and patient has clinical symptom (i.e. shortness of breather, syncope/fainting/lightheadedness, etc.) suggestive cardiac origin. (133)
- Patient has low to moderate hypertension that is medically managed and patient has clinical symptom (i.e. shortness of breather, syncope/fainting/lightheadedness, etc.) suggestive cardiac origin. (109)
- Pt has moderate to severe hypertension that is medically managed and patient has clinical symptom (i.e. shortness of breath/syncope/fainting/lightheadedness, etc.) suggestive cardiac origin. (109)

Evaluation of pulmonary hypertension may be reasonable and appropriate when the patient's medical record demonstrates ANY of the following:

- No prior echocardiogram with suspected pulmonary hypertension (115)
- Prior echocardiogram performed greater than or equal to one (1) year ago with known pulmonary hypertension. (115)
- Less than one year since last echocardiogram and EITHER of the following:
  - Known pulmonary hypertension and change in clinical status from the last cardiac evaluation. (115)
  - Known pulmonary hypertension and the echocardiogram is needed to guide therapy. (115)
Evaluation of pulmonary hypertension with change in clinical status or cardiac exam may be reasonable and appropriate when the patient has documented pulmonary hypertension and the medical record demonstrates **ANY** of the following:

- Severe elevated right ventricular systolic pressure on previous resting echo (127, 133)
- New or worsening symptoms suggest change in clinical status (e.g. Shortness of Breath, Lightheadedness, Presyncope, Syncope, Edema, Cyanosis, Palpitations) and Echo is needed to guide therapy OR evaluate response to therapy. (115, 133)
- New or worsening symptoms suggest change in clinical status (e.g. Shortness of Breath, Lightheadedness, Presyncope, Syncope, Edema, Cyanosis, Palpitations) and the patient has normal-borderline elevated right ventricular systolic pressure on previous resting echo. (127, 133)

Evaluation of suspected pulmonary hypertension with no prior echo may be reasonable and appropriate when the patient's medical record demonstrates no previous echo, pulmonary hypertension is suspected and echo is needed to evaluate right ventricular function and estimate pulmonary artery pressure. (133)

Evaluation of implantable cardio-defibrillator (ICD) candidacy may be reasonable and appropriate when the patient's medical record demonstrates that echo is needed to evaluate cardiac structure and function to assess candidacy of ICD implantation. (108, 115)

Evaluation of lightheadedness or pre-syncope may be reasonable and appropriate when the patient's medical record demonstrates **ANY** of the following:

- Clinical signs/symptoms suggestive of cardiac diagnosis known to cause lightheadedness (i.e. aortic stenosis, hypertrophic cardiomyopathy, or HF). (91, 122)
o Lightheadedness suspected to be of cardiac origin in-patient with known cardiac condition. (91)
o Patient has had repeated episodes of syncope. (133)

Evaluation of cardiac mass – tumor or thrombus - may be reasonable and appropriate when the patient’s medical record demonstrates ANY of the following:
o No prior echo with cardiac mass suspected. (36)
o Prior echo greater than one (1) year ago and cardiac mass is a documented diagnosis.
o Prior echo less than one (1) year ago, cardiac mass is a documented diagnosis and the patient has new symptoms or findings needed to guide therapy.
o No prior echo and patient has known primary cancer and cardiac involvement for staging is needed.
o Follow-up after surgical removal of a cardiac mass known to have high recurrence (i.e. myxoma). (110)
o Prior echocardiogram greater than one (1) year and patient with known primary cancer and cardiac involvement for staging is needed.

Evaluation of murmur as a new finding and is thought to be clinically significant may be reasonable and appropriate when the patient has valvular or structural heart disease reasonably suspected or confirmed and the medical record demonstrates ANY of the following:
o Continuous or late systolic or diastolic murmurs
o Grade 3 or greater mid-systolic murmurs
ECG OR chest X-ray findings suggest an abnormality (e.g. Ventricular hypertrophy, Myocardial infarction, LBBB/RBBB) and abnormal symptomatic findings such as frequent palpitations or auscultation. (116)

Evaluation of palpitations may be reasonable and appropriate when the patient's medical record demonstrates ANY of the following:

- No prior echocardiogram and recurrent and frequent palpitations (136)
- Prior echocardiogram LESS than twelve (12) months ago, recurrent and frequent palpitations with clinical change.
- Prior echocardiogram and arrhythmia documented by Telemetry, Holter monitoring or ECG (i.e. Atrial fibrillation, SVT or VT).
- Prior echocardiogram (TEE) Less than twelve (12) months ago with clinical change and arrhythmia documented by Telemetry, Holter monitoring or ECG (i.e. atrial fibrillation, SVT or VT).
- Prior echocardiogram less than twelve (12) months ago and palpitations, recurrent and frequent without clinical change.
- Prior echocardiogram greater than twelve (12) months ago with recurrent and frequent palpitations and clinical change present.
- Prior echocardiogram greater than twelve (12) months ago without clinical change and arrhythmia documented by Telemetry, Holter monitoring or ECG (i.e. Atrial fibrillation, SVT or VT).
- Frequent or exercise-induced Ventricular Premature Complexes (VPC) (129).

Evaluation of percutaneous non-coronary procedure, Trans-catheter aortic valve replacement (TAVR). Atrial septal defect (ASD), Post Atrial Fibrillation Ablation may be reasonable and appropriate when the patient's medical record demonstrates EITHER of the following:

- Transthoracic echocardiogram (TTE) for pre-procedural guidance. (111)
o Initial post-procedural transthoracic echocardiogram (TTE). (111,52)

Evaluation of pericardial disease may be reasonable and appropriate when the patient's medical record demonstrates EITHER of the following:
  o Suspected pericarditis or pericardial effusion. (92)
  o Pericarditis or a pericardial effusion is documented and a repeat TTE is needed to guide further therapy. (92)

Evaluation of pre-operative patient (non-cardiac surgery) when assessment is needed may be reasonable and appropriate when the patient's medical record demonstrates ANY of the following:
  o Prior echo greater than twelve (12) months ago and Heart disease is a documented condition. (116)
  o No prior echocardiogram or evaluation and heart disease is suspected based on symptoms, clinical findings, or diagnostic testing results AND needs evaluation prior to non-cardiac surgery. (92)
  o No prior echocardiogram or evaluation and Heart disease is suspected based on symptoms, clinical findings, or diagnostic testing results AND needs evaluation prior to non-cardiac surgery. (92)
  o No prior echocardiogram or evaluation and Evaluation of cardiac structure and function prior to non-cardiac solid organ transplantation. (135)
  o Prior echo GREATER than twelve (12) months ago and heart disease is a documented condition. (117)

Evaluation of respiratory problems – respiratory failure or pulmonary embolism - may be reasonable and appropriate when the patient's medical record demonstrates ANY of the following:
  o Known acute pulmonary embolism to guide therapy (e.g. thrombectomy and thrombolitics). (115)
Respiratory failure or hypoxemia of uncertain or suspected cardiac etiology. (115)

Routine surveillance of prior pulmonary embolism and prior assessment indicates change in right ventricular function and pulmonary artery pressure. (115)

Reevaluation of known pulmonary embolism after thrombolysis or thrombectomy and prior assessment indicates normal right ventricular function and/or pulmonary artery pressure. (115)

Evaluation of syncope may be reasonable and appropriate when the patient's medical record demonstrates ANY of the following:

- Last CAD testing over twelve (12) months ago and patient has had repeated episodes of syncope.
- Syncope in a patient with known cardiac condition. (91)
- Clinical signs/symptoms suggestive of cardiac diagnosis known to cause syncope (i.e. aortic stenosis, hypertrophic cardiomyopathy, or HF). (91)

Evaluation of moderate or severe valvular disease with clinical deterioration may be reasonable and appropriate when the patient has moderate or severe valvular or structural heart disease, which has been documented by previous imaging, and the medical record demonstrates ANY of the following:

- Moderate or severe valvular stenosis and ECG or chest X-ray findings suggest an abnormality (e.g. Ventricular hypertrophy, Myocardial infarction, LBBB/RBBB). (126)
- Moderate or severe valvular regurgitation and ECG or chest X-ray findings suggest an abnormality (e.g. Ventricular hypertrophy, Myocardial infarction, LBBB/RBBB). (126)
- New or worsening symptoms related to valvular disease (e.g. Dyspnea, Syncope, Angina, A fib, Hypertension). (126)
Evaluation of valvular function in evaluation of heart murmur may be reasonable and appropriate when the patient's medical record demonstrates ANY of the following:

- Valvular heart disease has been documented by prior echocardiography and new symptoms or examination findings are documented and suggest a change in the severity of previously documented valvular heart disease. (122)
- Valvular heart disease has been documented by prior echocardiography, last echo over 3 years ago, mild valvular insufficiency or mild valvular stenosis has been documented, and valvular heart disease has been documented and the echocardiogram is a periodic surveillance study. (115)
- Valvular heart disease has been documented by prior echocardiography, last echo greater than twelve (12) months ago, valvular heart disease has been documented and the echocardiogram is a periodic surveillance study, and moderate or severe valvular insufficiency or stenosis has been documented. (114)
- No prior echocardiogram or evaluation, initial evaluation of highly suspected valvular or structural heart disease, and new symptoms or examination findings are documented and suggest a change in the severity of previously documented valvular heart disease. (122)
- Initial evaluation of highly suspected valvular or structural heart disease and new symptoms or examination findings are documented and suggest a change in the severity of previously documented valvular heart disease. (122)
- Last echo greater than twelve (12) months ago and moderate to severe valvular regurgitation without change in clinical status or cardiac exam. (114)

Evaluation of prosthetic or repaired valve may be reasonable and appropriate when the patient's medical record demonstrates ANY of the following:

- Echocardiogram has not been performed since the patient's hospital procedure discharge. (114, 133)
o Surveillance echocardiogram: Greater than three (3) years since the one-year post-procedural milestone echocardiogram. (133)

o Echo is for surveillance one (1) year post procedure milestone. (114,133)

o There is a change or suspected malfunction of the repaired or replaced valve. (133)

o Reevaluation of known prosthetic valve when it would change management or guide therapy. (133)

o Initial postoperative evaluation of prosthetic valve for establishment of baseline. (114,133)

Evaluation of ventricular function – left ventricle to evaluate pre or post-cardio toxic chemotherapy may be reasonable and appropriate when the patient's medical record demonstrates EITHER of the following:

o When baseline echo is needed (112)

o When echo is for follow-up and ANY of the following:
  - Chemotherapy complete - initial follow up echo to assess cardiac function. (112)
  - Ongoing chemotherapy GREATER than two (2) months since last echo (112).
  - Chemotherapy complete- follow-up GREATER than one (1) year, restaging primary cancer. (131)
Evaluation of ventricular tachycardia (sustained or un-sustained), multifocal PVCs may be reasonable and appropriate when the patient’s medical record includes ECG or Holter monitoring which demonstrates Ventricular Tachycardia characterized by Multifocal Pre-Ventricular Contractions (PVC). (113)

Evaluation of aortic valve disease by transthoracic echo (TTE) may be reasonable and appropriate when the patient’s medical record demonstrates ANY of the following:

- No prior echocardiogram (TTE), initial evaluation of highly suspected aortic valve or structural heart disease, and new symptoms or examination highly suggestive of aortic valvular heart disease or cardiac structure abnormality. (133)
- Aortic valvular heart disease has been documented by prior echocardiography and new symptoms or examination findings are documented and suggest a change in the severity of previously documented aortic valvular heart disease. (133)
- Last echo over three (3) years ago, aortic valvular heart disease has been documented by prior echocardiography, mild aortic valvular insufficiency or mild aortic valvular stenosis has been documented, Aortic valvular heart disease has been documented and the echocardiogram is a periodic surveillance study. (133)
- Last echo between 1 and 3 years ago or over three (3) years ago, aortic valvular heart disease has been documented by prior, aortic valvular heart disease has been documented and the echocardiogram is a periodic surveillance study, moderate or severe aortic valvular insufficiency or stenosis has been documented.(133)
- Initial evaluation of highly suspected aortic valve or structural heart disease and new symptoms or examination findings are documented and suggest a
change in the severity of previously documented aortic valvular heart
disease.
  o Between 1 and 3 years ago or greater than 3 years and moderate to severe
  aortic valvular regurgitation without change in clinical status or cardiac
  exam.  (133)

Evaluation of mitral valve disorders by transthoracic echo (TTE) may be reasonable
and appropriate when the patient's medical record demonstrates ANY of the
following:
  o No prior echocardiogram or evaluation, initial evaluation of highly suspected
    mitral valve or structural heart disease, and new symptoms or examination
    findings are documented and suggest a change in the severity of previously
    documented mitral valvular heart disease.  (133)
  o Mitral valvular heart disease has been documented by prior
    echocardiography and new symptoms or examination findings are
    documented and suggest a change in the severity of previously documented
    mitral valvular heart disease.  (133)
  o Last echo greater than three (3) years ago and mitral valvular heart disease
    has been documented by prior echocardiography, mild mitral valvular
    insufficiency or mild aortic valvular stenosis has been documented and
    mitral valvular heart disease has been documented and the echocardiogram
    is a periodic surveillance study.  (133)
  o Last echo between 1 and 3 years ago; or greater than three (3) years ago,
    mitral valvular heart disease has been documented by prior
    echocardiography, mitral valvular heart disease has been documented and
    the echocardiogram is a periodic surveillance study, moderate or severe
    mitral valvular insufficiency or stenosis has been documented.  (133)
  o Initial evaluation of highly suspected mitral valve or structural heart disease
    and new symptoms or examination findings are documented and suggest a
change in the severity of previously documented mitral valvular heart
disease. (133)

- Last echo between 1 and 3 years ago or greater than three (3) years ago and
  moderate to severe mitral valvular regurgitation without change in clinical
  status or cardiac exam. (133)
Transesophageal Echocardiogram (TEE)

Evaluation of endocarditis by transesophageal echo TEE may be reasonable and appropriate when the patient's medical record demonstrates EITHER of the following:

- TTE and/or blood cultures are non-diagnostic and intent to diagnose infective endocarditis with a MODERATE or HIGH pretest probability. (107)
- Intent to diagnose infective endocarditis with a MODERATE or HIGH pretest probability and additional information that may impact clinical care is anticipated with TEE. (107,115)

Evaluation of aortic disease by transesophageal echo TEE may be reasonable and appropriate when the patient's medical record demonstrates suspected acute aortic pathology including but not limited to dissection/transection. (116)

Evaluation of atrial fibrillation/atrial flutter by transesophageal echo (TEE) may be reasonable and appropriate when ordered as valuation of atrial fibrillation to facilitate clinical decision making with regard to anticoagulation, cardioversion, and/or radiofrequency ablation. (88, 89, 94,115)

Evaluation of cardiac structure and function for general evaluation by transesophageal echo (TEE) may be reasonable and appropriate when the patient’s medical record demonstrates ANY of the following:

- There is a high probability of a non-diagnostic transthoracic echo (TTE) due to patient characteristics or inadequate visualization of relevant structures (115)
- Patient has chest wall injuries and previous transthoracic echo is non-diagnostic, inadequate, or insufficient in information provided.(124)
o Reevaluation of prior TEE finding for interval change when a change in therapy is anticipated. (115)

o Patient has cardiomegaly. (125)

o Patient has suspected mitral valve disorder or insufficiency and EITHER:
  ▪ Previous transthoracic echo is non-diagnostic, inadequate, or insufficient in information provided; (122)
  ▪ Pre-operative TEE evaluation needed (with prior insufficient TTE). (122)

o Patient has ostium secundum atrial septal defect and previous transthoracic echo is non-diagnostic, inadequate, or insufficient in information provided. (129)

o Patient has cardiomegaly and previous transthoracic echo is non-diagnostic, inadequate, or insufficient in information provided. (125)

Evaluation of cerebral ischemia or infarction by transesophageal echo (TEE) may be reasonable and appropriate when the patient’s medical record demonstrates EITHER of the following:

- Patient has had full workup of transient ischemic attack (non-cardiac, cardiac, carotid) and no clear source has been identified; (100)
- Patient has had a full workup and no clear source of cerebral artery occlusion has been identified. (100)

Evaluation of embolism by transesophageal echo (TEE) may be reasonable and appropriate when the request is for evaluation for cardiovascular source of embolus with no identified non-cardiac source. (100, 115)

Evaluation of guidance of percutaneous trans-catheter procedure by transesophageal echo (TEE) may be reasonable and appropriate when the patient is to undergo percutaneous non-coronary cardiac interventions, which require
guidance including, but not limited to, closure device placement, radiofrequency ablation, and percutaneous valve procedures. (93, 114, 115)

Evaluation of intraoperative TEE may be reasonable and appropriate when the patient's medical record demonstrates **EITHER** of the following:

- Patient to undergo any open heart or thoracic aortic surgical procedure that requires TEE (i.e. valve repair, valve replacement, and aortic dissection). (114)
- Patient is scheduled for coronary artery bypass graft surgery and a transthoracic echo (TTE) has a low probability of guiding surgery. (122)

Evaluation of valvular disease by transesophageal echo (TEE) may be reasonable and appropriate when the patient’s medical record demonstrates **ANY** of the following:

- Patient has suspected mitral valve disorder or insufficiency and there is a high probability of a non-diagnostic transthoracic echo (TTE) due to patient characteristics or inadequate visualization of relevant structures. (122)
- Patient has suspected aortic valve disorder or insufficiency and there is a high probability of a non-diagnostic transthoracic echo (TTE) due to patient characteristics or inadequate visualization of relevant structures. (122)
- There is a high probability of a non-diagnostic transthoracic echo (TTE) due to patient characteristics or inadequate visualization of relevant structures with evaluation of valvular structure and function to assess suitability for, and assist in planning of, an intervention. (122)
- Evaluation of paravalvular abscesses (both native and prosthetic valves) and prosthetic heart valves. (122)
- Patient has suspected mitral valve disorder or insufficiency and previous transthoracic echo is non-diagnostic. (122)
- Patient has suspected aortic valve disorder or insufficiency and previous transthoracic echo is non-diagnostic. (122)
- Previous transthoracic echo is non-diagnostic and Evaluation of valvular structure and function to assess suitability for, and assist in planning of, an intervention. (93)
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Implantable Loop Recorder

Placement of an Implantable Loop Recorder may be medically appropriate and supported by evidence to improve patient outcomes for the following indications.

Utilization in a patient who is over the age of 18 and has had a recent myocardial infarction (MI) may be reasonable and appropriate when the patient's medical record demonstrates **EITHER** of the following:

- Patient has a New York Heart Association (NYHA) Functional Classification of I, II, or III and has no valvular heart disease present; and **ALL** of the following:
  - Patient's life expectancy is greater than one (1) year;
  - Patient has had an echocardiogram performed in the past twenty-one (21) days demonstrating a left ventricular ejection fraction (LVEF) of 40 percent or less and cardiac wall motion of 1.3 or less; (1)
  - Patient has not had a coronary bypass graft (CABG) or previous insertion or explanations of an implantable cardioverter defibrillator (ICD) or cardiac pacemaker;
  - Patient is not currently pregnant or expecting to become pregnant;
  - Patient is currently in normal sinus rhythm;
  - Patient has a baseline electrocardiogram (ECG) or Holter monitor which demonstrates evidence of non-sustained ventricular tachycardia; (1)
  - Patient has a turbulence slope value of less than 0 or greater than 2.5ms/per normal to normal R-R interval on Holter monitoring;
  - Patient is at risk for sudden cardiac death (SCD) as evidenced by **ANY** of the following arrhythmias or conduction abnormalities:
    - New onset atrial fibrillation (A-fib);
    - Second- or third-degree atrioventricular (AV) block;
    - Bradycardia with or without beta-blocking therapy;
  - Patient has received a thirty (30) day ECG event monitoring previously. (1)
Patient has a NYHA Functional Classification of I, II, or III and mild to moderate valvular heart disease present; and ALL of the following:

- Patient’s life expectancy is greater than one (1) year;
- Patient has had an echocardiogram performed in the past twenty-one (21) days demonstrating an LVEF of 40 percent or less and cardiac wall motion of 1.3 or less;
- Patient has not had a CABG or previous insertion or explanations of an ICD or cardiac pacemaker;
- Patient is not currently pregnant or expecting to become pregnant;
- Patient is currently in normal sinus rhythm;
- Patient has a baseline ECG or Holter monitor which demonstrates evidence of non-sustained ventricular tachycardia;
- Patient has a turbulence slope value of less than 0 or greater than 2.5ms/per normal to normal R-R interval on Holter monitoring;
- Patient is at risk for SCD as evidenced by ANY of the following arrhythmias or conduction abnormalities:
  - New onset A fib; (1)
  - Second- or third-degree AV block; (1)
  - Bradycardia with or without beta-blocking therapy;
- Patient has received a thirty (30) day ECG event monitoring previously. (1)

Utilization in a patient who is 40 years of age or older and who has had a recent stroke or transient ischemic attack (TIA) but who has not had a recent MI may be reasonable and appropriate when the patient’s medical record demonstrates ALL of the following:

- Stroke or TIA occurred within the past ninety (90) days; (3)
- Cryptogenic stroke which has had all non-cardiac causes ruled out; (2,3)
- Patient does not have a history of A-fib; (2,3)
- Patient does not have an indication for and does not currently have an ICD or cardiac pacemaker; (2,3)
- Patient is a candidate for anti-coagulant therapy; (2,3)
• Patient has received a thirty (30) day ECG event monitoring previously which did not detect A-fib or any other cardiac arrhythmia.

Utilization in a patient who is under the age of 40 and who has had a recent stroke or TIA but who has not had a recent MI may be reasonable and appropriate when the patient’s medical record demonstrates ALL of the following:

• Stroke or TIA occurred within the past ninety (90) days;
• Cryptogenic stroke which has had all non-cardiac causes ruled out;
• Patient does not have a history of A-fib;
• Patient does not have an indication for and does not currently have an ICD or cardiac pacemaker;
• Patient is a candidate for anti-coagulant therapy;
• Patient has received a thirty (30) day ECG event monitoring previously which did not detect A-fib or any other cardiac arrhythmia.

Utilization in a patient who has not had a recent MI, stroke, or TIA but has been diagnosed with A-fib may be reasonable and appropriate when the patient’s medical record demonstrates EITHER of the following:

• Patient is receiving anti-coagulant therapy; and ALL of the following:
  ▪ Patient has undergone catheter ablation or Maze surgery which produced successful pulmonary vein isolation;
  ▪ Patient has A-fib that terminates spontaneously or with intervention within seven (7) days of onset;
  ▪ Transesophageal echocardiogram (TEE) was performed but it was non-diagnostic for structural causes of the patient’s symptoms;
  ▪ Discontinuation of anti-coagulants is being considered;
  ▪ Patient has received a thirty (30) day ECG event monitoring previously which was non-diagnostic for A-fib or any other cardiac arrhythmia;

• Patient is not receiving anti-coagulant therapy; and ALL of the following:
• Patient has undergone catheter ablation or Maze surgery which produced successful pulmonary vein isolation;
• Patient has A-fib that terminates spontaneously or with intervention within seven (7) days of onset;
• Patient is a candidate for anti-coagulation therapy;
• Patient has received a thirty (30) day ECG event monitoring previously which was non-diagnostic for A-fib or any other cardiac arrhythmia.

Utilization in a patient who has transient loss of consciousness but has not had a recent MI, stroke, TIA, or A-fib diagnosis may be reasonable and appropriate when the patient's medical record demonstrates ANY of the following:

  o Syncope or Pre-syncope are present with at least two (2) episodes of arrhythmogenic symptoms in the past six (6) months; and ALL of the following:
    • Unknown likelihood that episodes are being caused by a cardiac arrhythmia;
    • TWO or MORE of the following are present:
      • Palpitations present before syncopal episode;
      • Abnormal ECG and/or a diagnosis of heart disease is present;
      • Arrhythmogenic symptoms occur during effort;
      • Arrhythmogenic symptoms occur while the patient is supine;
    • Patient has had recent non-invasive cardiac imaging which has ruled out structural cardiac abnormalities and malignant ventricular arrhythmia as possible causes for arrhythmogenic symptoms;
    • Patient is not at high risk for SCD.
  o Syncope or Pre-syncope are present with at least two (2) episodes of arrhythmogenic symptoms in the past six (6) months; and ALL of the following:
    • Unknown likelihood that episodes are being caused by a cardiac arrhythmia;
• **TWO or MORE** of the following are present:
  - Palpitations present before syncopal episode;
  - Abnormal ECG and/or a diagnosis of heart disease is present;
  - Arrhythmogenic symptoms occur during effort;
  - Arrhythmogenic symptoms occur while the patient is supine;

• Patient has had recent non-invasive cardiac imaging which has ruled out structural cardiac abnormalities and malignant ventricular arrhythmia as possible causes for arrhythmogenic symptoms;

• Patient is at high risk for SCD;

• Electrophysiology study (EPS) has been performed and the patient is negative for ventricular tachycardia.

  o Syncope or Pre-syncope are present and patient has had one (1) episode every two (2) days for the past thirty (30) days and a thirty (30) day ECG event monitor has been unsuccessful in correlating the arrhythmogenic symptoms with a cardiac arrhythmia; and **ALL** of the following:
    - Unknown likelihood that episodes are being caused by a cardiac arrhythmia;

• **TWO or MORE** of the following are present:
  - Palpitations present before syncopal episode; (5)
  - Abnormal ECG and/or a diagnosis of heart disease is present;
  - Arrhythmogenic symptoms occur during effort;
  - Arrhythmogenic symptoms occur while the patient is supine;

• Patient has had recent non-invasive cardiac imaging which has ruled out structural cardiac abnormalities and malignant ventricular arrhythmia as possible causes for arrhythmogenic symptoms;

• Patient is not at high risk for SCD.

  o Syncope or Pre-syncope are present and patient has had one (1) episode every two (2) days for the past thirty (30) days and a thirty (30) day ECG event monitor has been unsuccessful in correlating the arrhythmogenic symptoms with a cardiac arrhythmia; and **ALL** of the following:
• Unknown likelihood that episodes are being caused by a cardiac arrhythmia;

• **TWO or MORE** of the following are present:
  • Palpitations present before syncopal episode;
  • Abnormal ECG and/or a diagnosis of heart disease is present;
  • Arrhythmogenic symptoms occur during effort;
  • Arrhythmogenic symptoms occur while the patient is supine;

• Patient has had recent non-invasive cardiac imaging which has ruled out structural cardiac abnormalities and malignant ventricular arrhythmia as possible causes for arrhythmogenic symptoms;

• Patient is at high risk for SCD;

• EPS has been performed and the patient is negative for ventricular tachycardia.

  o Syncope or Pre-syncope are present with at least two (2) episodes of arrhythmogenic symptoms in the past six (6) months; and **ALL** of the following:
    • High likelihood that episodes are being caused by a cardiac arrhythmia; (4,5)
    • Neurological factors have been ruled out as cause for arrhythmogenic symptoms; (4,5)
    • Orthostatic hypertension has been ruled out as cause for syncopal episodes;
    • Patient has had recent non-invasive cardiac imaging which has ruled out structural cardiac abnormalities and malignant ventricular arrhythmia as possible causes for arrhythmogenic symptoms;
    • Patient is not at high risk for SCD.

  o Syncope or Pre-syncope are present with at least two (2) episodes of arrhythmogenic symptoms in the past six (6) months; and **ALL** of the following:
    • High likelihood that episodes are being caused by a cardiac arrhythmia; (4)
Neurological factors have been ruled out as cause for arrhythmogenic symptoms;

Orthostatic hypertension has been ruled out as cause for arrhythmogenic symptoms;

Patient has had recent non-invasive cardiac imaging which has ruled out structural cardiac abnormalities and malignant ventricular arrhythmia as possible causes for arrhythmogenic symptoms;

Patient is at high risk for SCD;

EPS has been performed and the patient is negative for ventricular tachycardia.

- Syncope or Pre-syncope are present and patient has had one (1) episode every two (2) days for the past thirty (30) days and a thirty (30) day ECG event monitor has been unsuccessful in correlating the arrhythmogenic symptoms with a cardiac arrhythmia; and **ALL** of the following:
  - High likelihood that episodes are being caused by a cardiac arrhythmia;
  - Neurological factors have been ruled out as cause for arrhythmogenic symptoms;
  - Orthostatic hypertension has been ruled out as cause for arrhythmogenic symptoms;
  - Patient has had recent non-invasive cardiac imaging which has ruled out structural cardiac abnormalities and malignant ventricular arrhythmia as possible causes for arrhythmogenic symptoms;
  - Patient is not at high risk for SCD.

- Syncope or Pre-syncope are present and patient has had one (1) episode every two (2) days for the past thirty (30) days and a thirty (30) day ECG event monitor has been unsuccessful in correlating the arrhythmogenic symptoms with a cardiac arrhythmia; and **ALL** of the following:
  - High likelihood that episodes are being caused by a cardiac arrhythmia;
  - Neurological factors have been ruled out as cause for arrhythmogenic symptoms;
Orthostatic hypertension has been ruled out as cause for arrhythmogenic symptoms;

Patient has had recent non-invasive cardiac imaging which has ruled out structural cardiac abnormalities and malignant ventricular arrhythmia as possible causes for arrhythmogenic symptoms;

Patient is at high risk for SCD;

EPS has been performed and the patient is negative for ventricular tachycardia.

Seizure activity is present and patient has had one (1) episode every two (2) days for the past thirty (30) days and a thirty (30) day ECG event monitor has been unsuccessful in correlating the arrhythmogenic symptoms with a cardiac arrhythmia; and ALL of the following:

Abnormal ECG;

Patient has had non-invasive cardiac imaging demonstrating a LVEF of greater than 35 percent and no structural abnormalities, heart failure, or pulmonary hypertension identified;

Patient has had tilt table testing resulting in either maintenance of a normal heart rate and blood pressure or reduced heart rate of 10 to 15 beats per minute with normal blood pressure maintained;

Patient is not at high risk for SCD.

Seizure activity is present and patient has had one (1) episode every two (2) days for the past thirty (30) days and a thirty (30) day ECG event monitor has been unsuccessful in correlating the arrhythmogenic symptoms with a cardiac arrhythmia; and ALL of the following:

Abnormal ECG;

Patient has had non-invasive cardiac imaging demonstrating a LVEF of greater than 35 percent and no structural abnormalities, heart failure, or pulmonary hypertension identified;

Patient has had tilt table testing resulting in either maintenance of a normal heart rate and blood pressure or reduced heart rate of 10 to 15 beats per minute with normal blood pressure maintained;
● Patient is at high risk for SCD;
● EPS has been performed and the patient is negative for ventricular tachycardia.

○ Seizure activity is present with at least two (2) episodes of arrhythmogenic symptoms in the past six (6) months; and **ALL** of the following:
  - Abnormal ECG;
  - Patient has had non-invasive cardiac imaging demonstrating a LVEF of greater than 35 percent and no structural abnormalities, heart failure, or pulmonary hypertension identified;
  - Patient has had tilt table testing resulting in either maintenance of a normal heart rate and blood pressure or reduced heart rate of 10 to 15 beats per minute with normal blood pressure maintained;
  - Patient is not at high risk for SCD.

○ Seizure activity is present with at least two (2) episodes of arrhythmogenic symptoms in the past six (6) months; and **ALL** of the following:
  - Abnormal ECG;
  - Patient has had non-invasive cardiac imaging demonstrating a LVEF of greater than 35 percent and no structural abnormalities, heart failure, or pulmonary hypertension identified;
  - Patient has had tilt table testing resulting in either maintenance of a normal heart rate and blood pressure or reduced heart rate of 10 to 15 beats per minute with normal blood pressure maintained;
  - Patient is at high risk for SCD;
  - EPS has been performed and the patient is negative for ventricular tachycardia.

Utilization in a patient who is experiencing palpitations but has not had a recent MI, stroke, TIA, A-fib diagnosis, or transient loss of consciousness may be reasonable and appropriate when the patient's medical record demonstrates **ANY** of the following:

○ Patient has had a baseline ECG or Holter monitor with **ANY** of the following:
- Normal ECG; and **ALL** of the following: (6)
  - Normal result on prior non-invasive cardiac imaging; (6)
  - LVEF of greater than 35 percent; (6)
  - Patient has reported only one (1) episode of palpitations in the past seven (7) days. (6)
  - Patient has had previous 30-day event monitoring which did not correlate palpitations to a cardiac arrhythmia. (6)

- Normal ECG; and **ALL** of the following:
  - Normal result on prior non-invasive cardiac imaging; (6)
  - LVEF of greater than 35 percent; (6)
  - Patient has reported only one (1) episode of palpitations in the past seven (7) days; (6)
  - Patient has not had previous 30 day event monitoring. (6)

- Normal ECG; and **ALL** of the following:
  - Normal result on prior non-invasive cardiac imaging; (6)
  - LVEF of greater than 35 percent; (6)
  - Patient has reported only one (1) episode of palpitations more than seven (7) days ago. (6)

- Palpitations which have a high likelihood of arrhythmic origin and no family history of SCD; and **ALL** of the following:
  - Palpitations which have a high likelihood of arrhythmic origin and no family history of SCD; and **ALL** of the following: (6)
• Normal result on prior non-invasive cardiac imaging; (6)
• LVEF of greater than 35 percent; (6)
• Patient has reported only one (1) episode of palpitations in the past seven (7) days; (6)
• Patient has not had previous 30 day event monitoring. (6)

- Palpitations which have a high likelihood of arrhythmic origin and no family history of SCD; and ALL of the following:
  • Normal result on prior non-invasive cardiac imaging; (6)
  • LVEF of greater than 35 percent; (6)
  • Patient has reported only one (1) episode of palpitations more than seven (7) days ago. (6)
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Leadless Intracardiac Pacemaker

Utilization of a leadless intracardiac pacemaker may be medically appropriate and supported by evidence to improve patient outcomes for the following indications.

Insertion of a leadless intracardiac pacemaker may be reasonable and appropriate when the patient's medical record demonstrates **BOTH** of the following:

- Patient does NOT demonstrate any of the following:
  - Implanted mechanical tricuspid valve; (3,4,5,6,7,8)
  - Implanted inferior vena cave filter; (3,4,5,6,7,8)
  - Hypersensitivity to Dexamethasone Acetate (5,7,8)
  - Unfavorable femoral venous anatomy; (1,2)
  - Morbid obesity that prevents the implanted device from obtaining telemetry communication within 12.5 cm. (5,7,8)
  - Intolerance to heparin; (8)
  - Sensitivity to contrast media that cannot be adequately pre-medicated. (8)

- Patient has clinical factors that render atrial lead placement difficult, high risk or not deemed necessary for effective therapy. (1,2,3,6,8)
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Left Atrial Appendage Closure

Performing closure of the left atrial appendage (LAA) may be medically appropriate and supported by evidence to improve patient outcomes for the following indications.

LAA closure in patients with non-valvular atrial fibrillation may be reasonable and appropriate when the patient's medical record demonstrates ALL of the following:

- CHADS₂ of two (2) or greater or CHA₂DS₂ –VASc score of three (3) or greater has been determined; (1,2,3,5,6,8,10)
- Patient has had a diagnostic workup for atrial fibrillation (A-fib) using transesophageal echocardiography (TEE) which was negative for any structural heart disease. (4,10)
- Patient is currently on oral anti-coagulant therapy but is unsuitable for long-term use of this therapy due to a HAS-BLED score of three (3) or greater. (3,4,6,9,10)
- Patient is enrolled in, and the patient’s medical care team and facility are participating in a prospective, nation and audited registry for LAA closure. (3,7)
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MRA Heart

Utilization of a MRA of the heart may be medically appropriate and supported by evidence to improve patient outcomes for the following indications.

Utilization of a MRA of the heart prior to a cardiac ablation procedure is considered medically necessary and is supported by evidence-based medicine.

Utilization of a MRA of the heart to further evaluate a cardiac mass discovered by ultrasound is considered medically necessary and is supported by evidence-based medicine.
REFERENCES:


MRI Heart

Utilization of a MRI of the heart may be medically appropriate and supported by evidence to improve patient outcomes for the following indications.

Utilization of a MRI of the heart prior to a cardiac ablation procedure is considered medically necessary and is supported by evidence-based medicine.

Utilization of a MRI of the heart to further evaluate a cardiac mass discovered by ultrasound is considered medically necessary and is supported by evidence-based medicine.
REFERENCES:


Percutaneous Coronary Interventions

Utilization of a percutaneous coronary intervention (PCI) may be medically appropriate and supported by evidence to improve patient outcomes for the following indications.

PCI use in the presence of a previous coronary artery bypass graft (CABG) may be reasonable and appropriate when the patient’s medical record demonstrates EITHER of the following:

- Non-Invasive cardiac testing has been performed; and EITHER of the following:
  - Symptoms of cardiac ischemia are present; and ANY of the following:
    - Non-Invasive cardiac testing revealed an intermediate to high risk of coronary artery disease (CAD); and EITHER of the following:
      - Coronary angiogram illustrating coronary vessel stenosis of greater than 70% in a grafted vessel; (4,5,6)
      - Coronary angiogram illustrating coronary vessel stenosis of greater than 70% in a non-grafted vessel. (1,2,3,4,5,6)
    - Non-Invasive cardiac testing revealed a low risk of CAD; and EITHER of the following:
      - Moderate to severe ischemic symptoms, Canadian Class Score of 3-4; and EITHER of the following:
        - Coronary angiogram illustrating coronary vessel stenosis of greater than 70%;(1,2,3,4,5,6)
        - Coronary angiogram illustrating coronary vessel stenosis of greater than 70% in a non-grafted vessel. (1,2,3,4,5,6)
      - Coronary angiogram illustrating coronary vessel stenosis of greater than 70% in a non-grafted vessel.
- Non-Invasive cardiac testing has not been performed and patient is having moderate to severe symptoms of cardiac ischemia, Canadian Class Score of 3-4 while on maximal medical therapy for CAD; and EITHER of the following:
• Coronary angiogram illustrating coronary vessel stenosis of greater than 70% in a grafted vessel; (4,5,6)
• Coronary angiogram illustrating coronary vessel stenosis of greater than 70% in a non-grafted vessel. (1,2,3,4,5,6)

Initial use of PCI in the presence of symptomatic Ischemic CAD of a native vessel may be reasonable and appropriate when the patient's medical record demonstrates that a coronary angiogram has already been performed and EITHER of the following:

- Non-Invasive cardiac testing has been performed but a Fractional Flow Reserve (FFR) was not performed as part of a prior left heart catheterization; and EITHER of the following:
  • Non-Invasive cardiac testing revealed an intermediate to high risk of coronary artery disease (CAD); and ANY of the following:
    • Coronary angiogram illustrating any single proximal vessel with greater than 70% stenosis and left main stenosis of less than 50%, FFR was not performed due to facility or physician training limitations; (1)
    • Coronary angiogram illustrating left anterior descending (LAD) vessel with greater than 70% stenosis or greater than 70% stenosis in 2 or 3 vessels, FFR was not performed due to facility or physician training limitations;
    • Isolated left main stenosis of less than 50%; (2,3,6)
    • Isolated left main stenosis of greater than or equal to 50%. (2,3)
  • Non-Invasive cardiac testing revealed a low risk of CAD; and ANY of the following:
    • Coronary angiogram illustrating any single proximal vessel with greater than 70% stenosis and left main stenosis of less than 50%, FFR was not performed due to facility or physician training limitations; (1)
    • Coronary angiogram illustrating left anterior descending (LAD) vessel with greater than 70% stenosis or greater than 70% stenosis in 2 or 3 vessels, FFR was not performed due to facility or physician training limitations;
    • Isolated left main stenosis of less than 50%; (2,3,6)
    • Isolated left main stenosis of greater than or equal to 50%. (1,2,3,6)
o Non-Invasive cardiac testing has not been performed and patient is having moderate to severe symptoms of cardiac ischemia, Canadian Class Score of 3–4 while on maximal medical therapy for CAD; and EITHER of the following:
  ▪ Coronary angiogram illustrating any single proximal vessel with greater than 70% stenosis and left main stenosis of less than 50%, FFR was not performed due to facility or physician training limitations;
  ▪ Coronary angiogram illustrating left anterior descending (LAD) vessel with greater than 70% stenosis or greater than 70% stenosis in 2 or 3 vessels, FFR was not performed due to facility or physician training limitations.

Repeated use of PCI in the presence of symptomatic Ischemic CAD of a native vessel may be reasonable and appropriate when the patient's medical record demonstrates that a coronary angiogram has already been performed and ANY of the following:
  o This planned PCI will be conducted on the a vessel which was previously treated with PCI; and EITHER of the following:
    ▪ Non-Invasive cardiac testing revealed an intermediate to high risk of coronary artery disease (CAD); and ANY of the following:
      • Isolated left main stenosis of greater than or equal to 50%; (1,5,6)
      • Coronary angiogram illustrating any single proximal vessel with greater than 70% stenosis and left main stenosis of less than 50%; (2,3,5,6)
      • Coronary angiogram illustrating left anterior descending (LAD) vessel with greater than 70% stenosis or greater than 70% stenosis in 2 or 3 vessels as well as left main stenosis of less than 50% and moderate to severe symptoms of cardiac ischemia, Canadian Class Score of 3–4. (2,3,5,6)
    ▪ Non-Invasive cardiac testing revealed a low risk of CAD; and ANY of the following:
      • Isolated left main stenosis of greater than or equal to 50%;
• Coronary angiogram illustrating any single proximal vessel with greater than 70% stenosis and left main stenosis of less than 50% and moderate to severe symptoms of cardiac ischemia, Canadian Class Score of 3-4 are present; (2,3,5,6)

• Coronary angiogram illustrating left anterior descending (LAD) vessel with greater than 70% stenosis or greater than 70% stenosis in 2 or 3 vessels as well as left main stenosis of less than 50% and moderate to severe symptoms of cardiac ischemia, Canadian Class Score of 3-4. (2,3,5,6)

- Non-Invasive cardiac testing was not performed; and **ANY** of the following:
  - Coronary angiogram illustrating any single proximal vessel with greater than 70% stenosis and left main stenosis of less than 50% and moderate to severe symptoms of cardiac ischemia, Canadian Class Score of 3-4 are present; (2,3,5,6)
  - Coronary angiogram illustrating left anterior descending (LAD) vessel with greater than 70% stenosis or greater than 70% stenosis in 2 or 3 vessels as well as left main stenosis of less than 50% and mild symptoms of cardiac ischemia, Canadian Class Score of 1-2. (2,3,5,6)
  - Coronary angiogram illustrating borderline stenosis of 50-70% in any proximal vessel with no other stenosis of greater than 70% identified and mild symptoms of cardiac ischemia, Canadian Class Score of 1-2. (2,3,5,6)

  - This planned PCI will be conducted on a vessel other than that which was previously treated with PCI and Fractional Flow Reserve (FFR) was not performed as part of a prior left heart catheterization; and **ANY** of the following:
    - Non-Invasive cardiac testing revealed an intermediate to high risk of coronary artery disease (CAD); and **ANY** of the following:
      - Isolated left main stenosis of greater than or equal to 50%; (2,3,5,6)
      - Coronary angiogram illustrating any single proximal vessel with greater than 70% stenosis and left main stenosis of less than 50%; (2,3,5,6)
      - Coronary angiogram illustrating left anterior descending (LAD) vessel with greater than 70% stenosis or greater than 70% stenosis in 2 or 3 vessels.
stenosis in 2 or 3 vessels as well as left main stenosis of less than 50% and moderate to severe symptoms of cardiac ischemia, Canadian Class Score of 3-4, FFR was not performed due to facility or physician training limitations. (8)

- Non-Invasive cardiac testing revealed a low risk of CAD; and ANY of the following:
  - Isolated left main stenosis of greater than or equal to 50%; (1,2,3,5,6)
  - Coronary angiogram illustrating any single proximal vessel with greater than 70% stenosis and left main stenosis of less than 50% and moderate to severe symptoms of cardiac ischemia, Canadian Class Score of 3-4 are present; (2,3,5,6)
  - Coronary angiogram illustrating left anterior descending (LAD) vessel with greater than 70% stenosis or greater than 70% stenosis in 2 or 3 vessels as well as left main stenosis of less than 50% and moderate to severe symptoms of cardiac ischemia, Canadian Class Score of 3-4, FFR was not performed due to facility or physician training limitations. (8)

- Non-Invasive cardiac testing was not performed; and ANY of the following:
  - Coronary angiogram illustrating any single proximal vessel with greater than 70% stenosis and left main stenosis of less than 50% and moderate to severe symptoms of cardiac ischemia, Canadian Class Score of 3-4 are present;
  - Coronary angiogram illustrating left anterior descending (LAD) vessel with greater than 70% stenosis or greater than 70% stenosis in 2 or 3 vessels as well as left main stenosis of less than 50% and mild symptoms of cardiac ischemia, Canadian Class Score of 1-2. (2,3,4,5,6)
  - Coronary angiogram illustrating borderline stenosis of 50-70% in any proximal vessel with no other stenosis of greater than 70% identified and mild symptoms of cardiac ischemia, Canadian Class Score of 1-2. (2,3,4,5,6)

- Ad Hoc PCI where cardiac catheterization findings suggest 50-79% stenosis and the physician will perform an FFR prior to PCI and where PCI will only be performed if FFR is less than 0.80. (7)
PCI use for patient who is asymptomatic but based on Non-Invasive cardiac testing is suspected to have Ischemic CAD may be reasonable and appropriate when the patient's medical record demonstrates that a coronary angiogram has already been performed and EITHER of the following:

- Non-Invasive testing reveals low risk for CAD; and EITHER of the following:
  - Left main stenosis less than 50% and proximal LAD stenosis greater than 70%, patient is on maximal medical management for CAD and a FFR was not performed as part of the prior left heart catheterization due to facility or physician training limitations. (8)
  - Isolated left main stenosis of greater than or equal to 50% and a FFR was not performed as part of the prior left heart catheterization. (8)

- Non-Invasive testing reveals intermediate to high risk for CAD; and ANY of the following:
  - Left main stenosis less than 50% and proximal LAD stenosis greater than 70%, patient is on maximal medical management for CAD and a FFR was not performed as part of the prior left heart catheterization due to facility or physician training limitations. (8)
  - Left main stenosis less than 50% and proximal LAD stenosis greater than 70%, patient is not on maximal medical management for CAD and a FFR was not performed as part of the prior left heart catheterization due to facility or physician training limitations. (8)
  - Left main stenosis less than 50% and any single proximal vessel with greater than 70% stenosis, patient is on maximal medical management for CAD and a FFR was not performed as part of the prior left heart catheterization due to facility or physician training limitations.
  - Left main stenosis of less than 50% and an FFR was performed as part of the prior left heart catheterization to assess for lesion severity and the FFR was less than or equal to 0.80. (2,3,5,6,9)
  - Isolated left main stenosis of greater than or equal to 50% and a FFR was not performed as part of the prior left heart catheterization. (8)
REFERENCES:


7. (7) Ad Hoc Percutaneous Coronary Intervention: A Consensus Statement From the Society for Cardiovascular Angiography and Interventions; James C. Blankenship,1*MD, Osvaldo S. Gigliotti,2MD, Dmitriy N. Feldman,3 MD,Timothy A. Mixon,4MD, Rajan A.G.Patel,5 MD, Paul Sorajja,6MD,Steven J. Yakubov,7MD, and Charles E. Chambers,8 MD

8. (8) Current Use of Fractional Flow Reserve: A Nationwide Survey; Bashar Hannawi, MD Wilson W. Lam, MD Suwei Wang, PhD George A. Younis, MD

9. (9) Fractional Flow Reserve versus Angiography for Guiding Percutaneous Coronary Intervention Pim A.L. Tonino, M.D., Bernard De Bruyne, M.D., Ph.D., Nico H.J. Pijls, M.D., Ph.D., Uwe Siebert, M.D., M.P.H., Sc.D., Fumiai Ikono, M.D., Marcel van’t Veer, M.Sc., Volker Klauss, M.D., Ph.D., Ganesh Manoharan, M.D., Thomas Engström, M.D., Ph.D., Keith G. Oldroyd, M.D., Peter N. Ver Lee, M.D., Philip A. McCarthy, M.D., Ph.D., and William F. Fearon, M.D., for the FAME Study Investigators
Percutaneous Ventricular Assistive Device

Utilization of a percutaneous ventricular assistive device (PVAD) may be medically appropriate and supported by evidence to improve patient outcomes for the following indications.

PVAD to be utilized as a bridge to cardiac transplant or permanent left ventricular assistive device (LVAD) may be reasonable and appropriate when the patient’s medical record demonstrates the following:

- Patient is scheduled to have a cardiac transplant or permanent LVAD placement within the next fourteen (14) days; and
- **ANY** of the following:
  - Patient is status post-cardiotomy; and
  - **ANY** of the following:
    - Patient has experienced cardiogenic shock refractory to intra-aortic balloon pump (IABP); (1,2,4,8,9)
    - Patient is diagnosed with an aortic dissection; (10)
    - Patient has aortic stenting; (2,9)
    - Patient has severe aortic insufficiency. (9)
  - Patient is status post-acute myocardial infarction (MI); and
  - **ANY** of the following:
    - Patient has experienced cardiogenic shock refractory to IABP; (2,3,4,5,8)
    - Patient is diagnosed with an aortic dissection; (10)
    - Patient has aortic stenting; (2)
    - Patient has severe aortic insufficiency. (9)
  - Patient is high risk and needs ancillary support for an invasive cardiovascular procedure; and **ANY** of the following:
    - Patient has experienced cardiogenic shock refractory to IABP; (1,2,4,5,7)
    - Patient is diagnosed with an aortic dissection; (6)
    - Patient has aortic stenting; (2)
    - Patient has severe aortic insufficiency (6,7,9)


PVAD for use in a patient with cardio myogenic shock for IABP may be reasonable and appropriate when the patient's medical record demonstrates ANY of the following:

- Patient is status post-cardiotomy; and ANY of the following:
  - Patient has experienced cardiogenic shock refractory to intra-aortic balloon pump (IABP); (1,2,3,5,9,10)
  - Patient is diagnosed with an aortic dissection; (3,6,10)
  - Patient has aortic stenting; (2)
  - Patient has severe aortic insufficiency. (7,9,10)

- Patient is status post-acute myocardial infarction (MI); and ANY of the following:
  - Patient has experienced cardiogenic shock refractory to IABP; (1,2,5,8,9)
  - Patient is diagnosed with an aortic dissection; (6)
  - Patient has aortic stenting; (2)
  - Patient has severe aortic insufficiency. (6,9)

- Patient is high risk and needs ancillary support for an invasive cardiovascular procedure; and ANY of the following:
  - Patient has experienced cardiogenic shock refractory to IABP; (1,2,4,5,8)
  - Patient is diagnosed with an aortic dissection; (10)
  - Patient has aortic stenting; (2)
  - Patient has severe aortic insufficiency. (6)
REFERENCES:


4. Clinical Experience with the TandemHeart Percutaneous Ventricular Assist Device as a Bridge to Cardiac Transplantation, Brian A. Bruckner, MD, Leon P. Jacob, MD, Igor D. Gregoric, MD, Pranav Loyalka, MD, Biswajit Kar, MD, William E. Cohn, MD, Saverio La Francesca, MD, Branislav Radovancevic, MD, and O. H. Frazier, MD

5. Trends in the Use of Percutaneous Ventricular Assist Devices; Analysis of National Inpatient Sam{le Data, 2007 Through 2012, Rohan Khera, MD1; Peter Cram, MD, MBA2,3; Xin Lu, MS4; et al

6. Bridge to Transplantation with the TandemHeart, Bending the Indications in a Chronic Aortic Dissection Patient with Postcardiotomy Shock; Stephane Reverdin, MD Igor D. Gregoric, MD Biswajit Kar, MD Pranav Loyalka, MD Mark C. Bieniarz, MD Scott A. Lemaire, MD Joseph S. Coselli, MD O.H. Frazier, MD

7. Use of a Percutaneous Ventricular Assist Device for Treatment of Cardiogenic Shock due to Critical Aortic Stenosis; Christopher M. Frank, MD Nanthini Palanichamy, MD Biswajit Kar, MD James M. Wilson, MD Igor D. Gregoric, MD Pranav Loyalka, MD Andrew B. Civitello, MD.


9. Mechanical circulatory support in acute cardiogenic shock; Mubashar H Khan, Brian J Corbett and Steven M Hollenberg*

10. Percutaneous Circulatory Support in Cardiogenic Shock, Interventional Bridge to Recovery; Biswajit Kar, MD; Sukhdeep S. Basra, MD, MPH; Nishant R. Shah, MD; Pranav Loyalka, MD.
Trans-catheter Aortic Valve Replacement

Performing a Trans-catheter Aortic Valve Replacement (TAVR) may be medically appropriate and supported by evidence to improve patient outcomes for the treatment of aortic valve stenosis when the patient's medical record demonstrates **EITHER** of the following:

- Patient has activities of daily living (ADL) score of three (3) or more with asymptomatic Stage C ventricular heart disease and aortic velocity of greater than 5.0m/s or mean pressure gradient greater than 40mmHg; and **ANY** of the following:
  - Patient has a predicted post-TAVR survival rate of greater than twelve (12) months; and **ANY** of the following:
    - Left ventricular ejection fraction (LVEF) of less than 50% and greater than 50% predicted post-operative surgical risk morbidity from all causes at one (1) year; (1,4,5)
    - Left ventricular ejection fraction (LVEF) of less than 50% and three (3) or more organ systems which are compromised and not expected to improve post-operatively;
    - Left ventricular ejection fraction (LVEF) of less than 50% and patient has a severe procedure specific impediment; (4)
    - Patient has a decreased exercise tolerance or a drop in blood pressure is exhibited with exercise and greater than 50% predicted post-operative surgical risk morbidity from all causes at one (1) year; (8)
    - Patient has a decreased exercise tolerance or a drop in blood pressure is exhibited with exercise and three (3) or more organ systems which are compromised and not expected to improve post-operatively;
    - Patient has a decreased exercise tolerance or a drop in blood pressure is exhibited with exercise and patient has a severe procedure specific impediment;
o Left ventricular ejection fraction (LVEF) of less than 50% and patient has a Society of Thoracic Surgeons Predicted Risk of Mortality (STS-PROM) greater than 8%; (3,5)

o Left ventricular ejection fraction (LVEF) of less than 50% and two (2) or more organ systems which are compromised and not expected to improve post-operatively;

o Left ventricular ejection fraction (LVEF) of less than 50% and patient has a possible procedure specific impediment; (5)

o Left ventricular ejection fraction (LVEF) of less than 50% and patient has moderate to severe frailty; (1)

o Patient has a decreased exercise tolerance or a drop in blood pressure is exhibited with exercise and the patient has a Society of Thoracic Surgeons Predicted Risk of Mortality (STS-PROM) greater than 8%; (3)

o Patient has a decreased exercise tolerance or a drop in blood pressure is exhibited with exercise and two (2) or more organ systems which are compromised and not expected to improve post-operatively;

o Patient has a decreased exercise tolerance or a drop in blood pressure is exhibited with exercise and patient has a possible procedure specific impediment. (7)

o Patient has a decreased exercise tolerance or a drop in blood pressure is exhibited with exercise and patient has moderate to severe frailty. (7)

Patient has activities of daily living (ADL) score of three (3) or more with symptomatic Stage D ventricular heart disease; and **ALL** of the following:

o Patient has a predicted post-TAVR survival rate of greater than twelve (12) months; (6)

o Left ventricular ejection fraction (LVEF) of less than 50%;

o Greater than 50% predicted post-operative surgical risk morbidity from all causes at one (1) year; (7)

o Aortic velocity of greater than 5.0m/s or mean pressure gradient greater than 40mmHg; (2,9)
• Patient has low-flow, low-grade aortic stenosis.

As well as ANY of the following:

• Greater than 50% predicted post-operative surgical risk morbidity from all causes at one (1) year; (1,6)
• Patient has two (2) or more organ systems which are compromised and not expected to improve post-operatively (4,6,9)
• Patient has a possible or severe procedure specific impediment; (8,9)
• Patient has a Society of Thoracic Surgeons Predicted Risk of Mortality (STS-PROM) greater than 8%; (3,5,7,8,9)
• Patient has moderate to severe frailty. (2,9)
REFERENCES:

Clinical Frailty Scale:

Very Fit – People who are robust, active, energetic and motivated. These people commonly exercise regularly. They are among the fittest for their age.

Well – People who have no active disease symptoms but are less fit than category 1. Often, they exercise or are very active occasionally, e.g. seasonally.

Managing Well – People whose medical problems are well controlled, but are not regularly active beyond routine walking.

Vulnerable – While not dependent on others for daily help, often symptoms limit activities. A common complaint is being "slowed up", and/or being tired during the day.

Mildly Frail – These people often have more evident slowing, and need help in high order IADLs (finances, transportation, heavy housework, medications). Typically, mild frailty progressively impairs shopping and walking outside alone, meal preparation and housework.

Moderately Frail – People need help with all outside activities and with keeping house. Inside, they often have problems with stairs, need help with bathing, and might need minimal assistance (cuing, standby) with dressing.

Severely Frail – Completely dependent for personal care, from whatever cause (physical or cognitive). Even so, they seem stable and not at high risk of dying (within ~ 6 months).

Very Severely Frail – Completely dependent, approaching the end of life. Typically, they could not recover even from a minor illness.

Terminally Ill - Approaching the end of life. This category applies to people with a life expectancy <6 months, who are not otherwise evidently frail.

Scoring frailty in people with dementia:

The degree of frailty corresponds to the degree of dementia.
Common symptoms in mild dementia include forgetting the details of a recent event, though still remembering the event itself, repeating the same question/story and social withdrawal.

In moderate dementia, recent memory is very impaired, even though they seemingly can remember their past life events well. They can do personal care with prompting.

In severe dementia, they cannot do personal care without help.


Trans-catheter Mitral Valve Replacement

Performing a Trans-catheter Mitral Valve Repair (TMVR) may be medically appropriate and supported by evidence to improve patient outcomes for the treatment of mitral valve regurgitation when the patient’s medical record demonstrates ALL of the following:

- Patient has a high surgical risk; (1,3,4,5,6,7,8,9,11,12,13)
- Patient has no other indication for cardiac surgery, i.e. Coronary bypass graft, aortic valve repair etc.; (2)
- Patient is a candidate for surgical mitral valve repair; (1,2,3,5,7,13,15)
- Patient has primary mitral valve regurgitation resulting in anatomical alteration of the valve leaflets or sub-valvular structures; (2,12,13)
- Patient is exhibiting severe symptoms despite guideline directed medical therapy (GDMT) for heart failure; (2,3,7,8,13,15)
- Left ventricular ejection fraction (LVEF) is between 20 and 60%; (2,6,11,13)
- Care will be overseen utilizing a heart team approach with surgical representation to determine the most appropriate plan of care; (7,9,10,13)
- The facility where the procedure is to be conducted has a surgical team with at least twenty-five (25) total mitral valve procedures for severe mitral regurgitation per year and at least ten (10) of these must be mitral valve repairs;
- The facility where the procedure is to be conducted has an interventional cardiology program which performs at least one thousand (1000) cardiac catheterizations per year which includes at least four hundred (400) percutaneous coronary interventions (PCI) per year with acceptable outcomes compared to the National Cardiovascular Data Registry (NCDR) benchmarks with each interventional cardiologist performing at least fifty (50) structural...
procedures per year including atrial septal defects (ASD), patent foramen ovale (PFO) and trans-septal punctures;

- This request is going to be reported to the single national database; (7,8,10)
- The heart team and facility must be participating in a prospective, national, audited registry that collects data to address how outcomes, adverse events, device durability and patient demographics compare to pivotal clinical studies in statistical populations and subpopulations.

As well as **ANY** of the following:

- Patient is diagnosed with mitral regurgitation severity of a three (3) and New York Heart Association (NYHA) class three (3) heart failure; (1,2,5,7,8,9,11,12,14)
- Patient is diagnosed with mitral regurgitation severity of a four (4) and New York Heart Association (NYHA) class three (3) heart failure; (1,2,5,7,8,9,11,12,14)
- Patient is diagnosed with mitral regurgitation severity of a three (3) and New York Heart Association (NYHA) class four (4) heart failure; (1,2,5,7,8,9,11,12,14)
- Patient is diagnosed with mitral regurgitation severity of a four (4) and New York Heart Association (NYHA) class four (4) heart failure. (1,2,5,7,8,9,11,12,14)
REFERENCES:


Wearable Cardiac Defibrillator

Utilization of a wearable cardiac defibrillator (WCD) may be medically appropriate and supported by evidence to improve patient outcomes for the following indications where the patient has no contraindications:

WCD status post coronary artery bypass graft (CABG) or percutaneous cardiac intervention (PCI) may be reasonable and appropriate when the patient's medical record demonstrates ALL of the following:

- New York Heart Association (NYHA) functional class score of 2 or 3
- Left ventricular ejection fraction (LVEF) of less than or equal to 35% documented on previous cardiac study
- The patient is expected to receive a permanent implanted cardiac defibrillator (ICD) within the next ninety (90) days or after any temporary contraindication or complication to receiving a permanent ICD is resolved with or without concurrent disease, which prevents immediate implantation.

WCD status post cardiac rhythm abnormality or cardiac arrest may be reasonable and appropriate when the patient's medical record demonstrates ALL of the following:

- The patient has no contraindication to an automatic implantable cardioverter defibrillator;
- The patient has had an episode of cardiac arrest due to irreversible ventricular fibrillation;
- The patient is expected to receive a permanent implanted cardiac defibrillator (ICD) within the next ninety (90) days or after any temporary contraindication or complication to receiving a permanent ICD is resolved with or without concurrent disease which prevents immediate implantation;
- The patient has had sustained ventricular tachyarrhythmia either induced during electrophysiology study or spontaneously for greater than thirty (30) seconds.
REFERENCES:


### APPENDIX A: PROCEDURE CODES ASSOCIATED WITH THIS POLICY

Any procedure codes that have been associated with this HealthHelp Clinical Guideline are for informational use only. The inclusion of a code in this guideline does not guarantee coverage or reimbursement by the individual health plan.

#### CARDIOLOGY SERVICES

<table>
<thead>
<tr>
<th>CTA</th>
<th>CODES:</th>
</tr>
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<tbody>
<tr>
<td>Computed tomography, heart, without contrast material, with quantitative evaluation of coronary calcium</td>
<td>75571</td>
</tr>
<tr>
<td>Computed tomography, heart, with contrast material, for evaluation of cardiac structure and morphology (including 3D image postprocessing, assessment of cardiac function, and evaluation of venous structures, if performed)</td>
<td>75572</td>
</tr>
<tr>
<td>Computed tomography, heart, with contrast material, for evaluation of cardiac structure and morphology in the setting of congenital heart disease (including 3D image postprocessing, assessment of LV cardiac function, RV structure and function and evaluation of venous structures, if performed)</td>
<td>75573</td>
</tr>
<tr>
<td>Computed tomographic angiography, heart, coronary arteries and bypass grafts (when present), with contrast material, including 3D image postprocessing (including evaluation of cardiac structure and morphology, assessment of cardiac function, and evaluation of venous structures, if performed)</td>
<td>75574</td>
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<th>MRA</th>
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<tr>
<td>MRI</td>
<td>CODES:</td>
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<tr>
<td>Cardiac magnetic resonance imaging for morphology and function without contrast material;</td>
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<tr>
<td>Cardiac magnetic resonance imaging for morphology and function without contrast material; with stress imaging</td>
<td>75559</td>
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<tr>
<td>Cardiac magnetic resonance imaging for morphology and function without contrast material(s), followed by contrast material(s) and further sequences;</td>
<td>75561</td>
</tr>
<tr>
<td>Cardiac magnetic resonance imaging for morphology and function without contrast material(s), followed by contrast material(s) and further sequences; with stress imaging</td>
<td>75563</td>
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<tr>
<th>PET</th>
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<tr>
<td>Myocardial imaging, positron emission tomography (PET), perfusion; single study at rest or stress</td>
<td>78491</td>
</tr>
<tr>
<td>Myocardial imaging, positron emission tomography (PET), perfusion; multiple studies at rest and/or stress</td>
<td>78492</td>
</tr>
<tr>
<td>Myocardial imaging, positron emission tomography (PET), perfusion; single study at rest or stress</td>
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<tr>
<td>Myocardial imaging, positron emission tomography (PET), perfusion; multiple studies at rest and/or stress</td>
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<td>Service Description</td>
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<td>Myocardial imaging, positron emission tomography (PET), perfusion; single study at rest or stress</td>
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<tr>
<td>Myocardial imaging, positron emission tomography (PET), perfusion; multiple studies at rest and/or stress</td>
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<td><strong>Cardiac Nuclear Medicine</strong></td>
<td>CODES:</td>
</tr>
<tr>
<td>Myocardial perfusion imaging, tomographic (SPECT) (including attenuation correction, qualitative or quantitative wall motion, ejection fraction by first pass or gated technique, additional quantification, when performed); single study, at rest or stress (exercise or pharmacologic)</td>
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<tr>
<td>Myocardial perfusion imaging, tomographic (SPECT) (including attenuation correction, qualitative or quantitative wall motion, ejection fraction by first pass or gated technique, additional quantification, when performed); multiple studies, at rest and/or stress (exercise or pharmacologic) and/or redistribution and/or reinjection</td>
<td>78452</td>
</tr>
<tr>
<td>Myocardial perfusion imaging, planar (including qualitative or quantitative wall motion, ejection fraction by first pass or gated technique, additional quantification, when performed); single study, at rest or stress (exercise or pharmacologic)</td>
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</tr>
<tr>
<td>Myocardial perfusion imaging, planar (including qualitative or quantitative wall motion, ejection fraction by first pass or gated technique, additional quantification, when performed); multiple studies, at rest and/or stress (exercise or pharmacologic) and/or redistribution and/or reinjection</td>
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<td>Myocardial imaging, infarct avid, planar; qualitative or quantitative</td>
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<tr>
<td>Myocardial imaging, infarct avid, planar; with ejection fraction by first pass technique</td>
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<tr>
<td>Myocardial imaging, infarct avid, planar; tomographic SPECT with or without quantification</td>
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</tr>
<tr>
<td>Cardiac blood pool imaging, gated equilibrium; planar, single study at rest or stress (exercise and/or pharmacologic), wall motion study plus ejection fraction, with or without additional quantitative processing</td>
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<tr>
<td>Cardiac blood pool imaging, gated equilibrium; multiple studies, wall motion study plus ejection fraction, at rest and stress (exercise and/or pharmacologic), with or without additional quantification</td>
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<tr>
<td>Cardiac blood pool imaging (planar), first pass technique; single study, at rest or with stress (exercise and/or pharmacologic), wall motion study plus ejection fraction, with or without quantification</td>
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<tr>
<td>Cardiac blood pool imaging (planar), first pass technique; multiple studies, at rest and with stress (exercise and/or pharmacologic), wall motion study plus ejection fraction, with or without quantification</td>
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<tr>
<td>Cardiac blood pool imaging, gated equilibrium, SPECT, at rest, wall motion study plus ejection fraction, with or without quantitative processing</td>
<td>78494</td>
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<tr>
<td><strong>Cardiac Catheterization</strong></td>
<td>CODES:</td>
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<tr>
<td>Right heart catheterization including measurement(s) of oxygen saturation and cardiac output, when performed</td>
<td>93451</td>
</tr>
<tr>
<td>Left heart catheterization including intraprocedural injection(s) for left ventriculography, imaging supervision and interpretation, when performed</td>
<td>93452</td>
</tr>
<tr>
<td>Code</td>
<td>Description</td>
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<tr>
<td>93453</td>
<td>Combined right and left heart catheterization including intraprocedural injection(s) for left ventriculography, imaging supervision and interpretation, when performed</td>
</tr>
<tr>
<td>93454</td>
<td>Catheter placement in coronary artery(s) for coronary angiography, including intraprocedural injection(s) for coronary angiography, imaging supervision and interpretation;</td>
</tr>
<tr>
<td>93455</td>
<td>Catheter placement in coronary artery(s) for coronary angiography, including intraprocedural injection(s) for coronary angiography, imaging supervision and interpretation; with catheter placement(s) in bypass graft(s) (internal mammary, free arterial venous grafts) including intraprocedural injection(s) for bypass graft angiography</td>
</tr>
<tr>
<td>93456</td>
<td>Catheter placement in coronary artery(s) for coronary angiography, including intraprocedural injection(s) for coronary angiography, imaging supervision and interpretation; with right heart catheterization</td>
</tr>
<tr>
<td>93457</td>
<td>Catheter placement in coronary artery(s) for coronary angiography, including intraprocedural injection(s) for coronary angiography, imaging supervision and interpretation; with catheter placement(s) in bypass graft(s) (internal mammary, free arterial, venous grafts) including intraprocedural injection(s) for bypass graft angiography and right heart catheterization</td>
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<tr>
<td>93458</td>
<td>Catheter placement in coronary artery(s) for coronary angiography, including intraprocedural injection(s) for coronary angiography, imaging supervision and interpretation; with left heart catheterization including intraprocedural injection(s) for left ventriculography, when performed</td>
</tr>
<tr>
<td>93459</td>
<td>Catheter placement in coronary artery(s) for coronary angiography, including intraprocedural injection(s) for coronary angiography, imaging supervision and interpretation; with left heart catheterization including intraprocedural injection(s) for left ventriculography, when performed, catheter placement(s) in bypass graft(s) (internal mammary, free arterial, venous grafts) with bypass graft angiography</td>
</tr>
<tr>
<td>93460</td>
<td>Catheter placement in coronary artery(s) for coronary angiography, including intraprocedural injection(s) for coronary angiography, imaging supervision and interpretation; with right and left heart catheterization including intraprocedural injection(s) for left ventriculography, when performed</td>
</tr>
<tr>
<td>93461</td>
<td>Catheter placement in coronary artery(s) for coronary angiography, including intraprocedural injection(s) for coronary angiography, imaging supervision and interpretation; with right and left heart catheterization including intraprocedural injection(s) for left ventriculography, when performed, catheter placement(s) in bypass graft(s) (internal mammary, free arterial, venous grafts) with bypass graft angiography</td>
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<tr>
<td>93530</td>
<td>Right heart catheterization, for congenital cardiac anomalies</td>
</tr>
<tr>
<td>93531</td>
<td>Combined right heart catheterization and retrograde left heart catheterization, for congenital cardiac anomalies</td>
</tr>
<tr>
<td>93532</td>
<td>Combined right heart catheterization and transseptal left heart catheterization through intact septum with or without retrograde left heart catheterization, for congenital cardiac anomalies</td>
</tr>
<tr>
<td>93533</td>
<td>Combined right heart catheterization and transseptal left heart catheterization through existing septal opening, with or without retrograde left heart catheterization, for congenital cardiac anomalies</td>
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</tbody>
</table>
Indicator dilution studies such as dye or thermal dilution, including arterial and/or venous catheterization; with cardiac output measurement (separate procedure) 93561

Indicator dilution studies such as dye or thermal dilution, including arterial and/or venous catheterization; subsequent measurement of cardiac output (separate procedure) 93562

<table>
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<tr>
<th><strong>Cardiac Devices - Pacemakers</strong></th>
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<tr>
<td>Insertion or replacement of permanent pacemaker with transvenous electrode(s); atrial</td>
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<tr>
<td>Insertion or replacement of permanent pacemaker with transvenous electrode(s); ventricular</td>
<td>33207</td>
</tr>
<tr>
<td>Insertion or replacement of permanent pacemaker with transvenous electrode(s); atrial and ventricular</td>
<td>33208</td>
</tr>
<tr>
<td>Insertion or replacement of temporary transvenous single chamber cardiac electrode or pacemaker catheter (separate procedure)</td>
<td>33210</td>
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<tr>
<td>Insertion or replacement of temporary transvenous dual chamber pacing electrodes (separate procedure)</td>
<td>33211</td>
</tr>
<tr>
<td>Insertion or replacement of pacemaker pulse generator only; single chamber, atrial or ventricular</td>
<td>33212</td>
</tr>
<tr>
<td>Insertion or replacement of pacemaker pulse generator only; dual chamber</td>
<td>33213</td>
</tr>
<tr>
<td>Upgrade of implanted pacemaker system, conversion of single chamber system to dual chamber system (includes removal of previously placed pulse generator, testing of existing lead, insertion of new lead, insertion of new pulse generator)</td>
<td>33214</td>
</tr>
<tr>
<td>Removal of permanent pacemaker pulse generator with replacement of pacemaker pulse generator; single lead system</td>
<td>33227</td>
</tr>
<tr>
<td>Removal of permanent pacemaker pulse generator with replacement of pacemaker pulse generator; dual lead system</td>
<td>33228</td>
</tr>
<tr>
<td>Removal of permanent pacemaker pulse generator</td>
<td>33233</td>
</tr>
<tr>
<td>Removal of transvenous pacemaker electrode(s); single lead system, atrial or ventricular</td>
<td>33234</td>
</tr>
<tr>
<td>Removal of transvenous pacemaker electrode(s); dual lead system</td>
<td>33235</td>
</tr>
</tbody>
</table>

**Cardiac Devices - Automatic Implantable Cardioverter Defibrillator (AICD)**

| CODES: |
|--------------------------------|-----------|
| Insertion of a single transvenous electrode, permanent pacemaker or implantable defibrillator | 33216 |
| Insertion of 2 transvenous electrodes, permanent pacemaker or implantable defibrillator | 33217 |
| Insertion of pacing implantable defibrillator pulse generator only; with existing dual leads | 33230 |
| Insertion of implantable defibrillator pulse generator only; with existing single lead | 33240 |
| Removal of implantable defibrillator pulse generator only | 33241 |
| Removal of single or dual chamber implantable defibrillator electrode(s); by transvenous extraction | 33244 |
| Insertion or replacement of permanent implantable defibrillator system with transvenous lead(s), single or dual chamber | 33249 |
| Removal of implantable defibrillator pulse generator with replacement of implantable defibrillator pulse generator; single lead system | 33262 |
| Removal of implantable defibrillator pulse generator with replacement of implantable defibrillator pulse generator; dual lead system | 33263 |
Insertion or replacement of permanent subcutaneous implantable defibrillator system, with subcutaneous electrode, including defibrillation threshold evaluation, induction of arrhythmia, evaluation of sensing for arrhythmia termination, and programming or reprogramming of sensing or therapeutic parameters, when performed

- Insertion of subcutaneous implantable defibrillator electrode
- Repositioning of previously implanted subcutaneous implantable defibrillator electrode

### Cardiac Devices - Cardiac Resynchronization Therapy - Pacemaker (CRT-P) CODES:

- Insertion or replacement of permanent pacemaker with transvenous electrode(s); ventricular
- Insertion or replacement of permanent pacemaker with transvenous electrode(s); atrial and ventricular
- Upgrade of implanted pacemaker system, conversion of single chamber system to dual chamber system (includes removal of previously placed pulse generator, testing of existing lead, insertion of new lead, insertion of new pulse generator)
- Insertion of pacemaker pulse generator only; with existing multiple leads
- Insertion of pacing electrode, cardiac venous system, for left ventricular pacing, with attachment to previously placed pacemaker or implantable defibrillator pulse generator (including revision of pocket, removal, insertion, and/or replacement of existing generator)
- Removal of permanent pacemaker pulse generator with replacement of pacemaker pulse generator; multiple lead system
- Removal of permanent pacemaker pulse generator
- Removal of transvenous pacemaker electrode(s); single lead system, atrial or ventricular

### Cardiac Devices - Cardiac Resynchronization Therapy - Defibrillator (CRT-D) CODES:

- Insertion of a single transvenous electrode, permanent pacemaker or implantable defibrillator
- Insertion of 2 transvenous electrodes, permanent pacemaker or implantable defibrillator
- Insertion of pacing electrode, cardiac venous system, for left ventricular pacing, with attachment to previously placed pacemaker or implantable defibrillator pulse generator (including revision of pocket, removal, insertion, and/or replacement of existing generator)
- Insertion of pacing implantable defibrillator pulse generator only; with existing multiple leads
- Insertion of implantable defibrillator pulse generator only; with existing single lead
- Removal of implantable defibrillator pulse generator only
- Removal of single or dual chamber implantable defibrillator electrode(s); by transvenous extraction
- Insertion or replacement of permanent implantable defibrillator system with transvenous lead(s), single or dual chamber
- Removal of implantable defibrillator pulse generator with replacement of implantable defibrillator pulse generator; multiple lead system

### Cardiac Devices - Wearable CODES:
Initial set-up and programming by a physician of wearable cardioverter-defibrillator includes initial programming of system, establishing baseline electronic ECG, transmission of data to data repository, patient instruction in wearing system and patient reporting of problems or events.

<table>
<thead>
<tr>
<th>Cardiac Devices - HCPCS</th>
<th>CODES:</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cardioverter-defibrillator, dual chamber (implantable)</td>
<td>C1721</td>
</tr>
<tr>
<td>Cardioverter-defibrillator, single chamber (implantable)</td>
<td>C1722</td>
</tr>
<tr>
<td>Lead, cardioverter-defibrillator, endocardial single coil (implantable)</td>
<td>C1777</td>
</tr>
<tr>
<td>Lead, pacemaker, transvenous VDD single pass</td>
<td>C1779</td>
</tr>
<tr>
<td>Pacemaker, dual chamber, rate-responsive (implantable)</td>
<td>C1785</td>
</tr>
<tr>
<td>Pacemaker, single chamber, rate-responsive (implantable)</td>
<td>C1786</td>
</tr>
<tr>
<td>Cardioverter-defibrillator, other than single or dual chamber (implantable)</td>
<td>C1882</td>
</tr>
<tr>
<td>Lead, cardioverter-defibrillator, endocardial dual coil (implantable)</td>
<td>C1895</td>
</tr>
<tr>
<td>Lead, cardioverter-defibrillator, other than endocardial single or dual coil (implantable)</td>
<td>C1896</td>
</tr>
<tr>
<td>Lead, pacemaker, other than transvenous VDD single pass</td>
<td>C1898</td>
</tr>
<tr>
<td>Lead, pacemaker/cardioverter-defibrillator combination (implantable)</td>
<td>C1899</td>
</tr>
<tr>
<td>Lead, left ventricular coronary venous system</td>
<td>C1900</td>
</tr>
<tr>
<td>Pacemaker, dual chamber, nonrate-responsive (implantable)</td>
<td>C2619</td>
</tr>
<tr>
<td>Pacemaker, single chamber, nonrate-responsive (implantable)</td>
<td>C2620</td>
</tr>
<tr>
<td>Pacemaker, other than single or dual chamber (implantable)</td>
<td>C2621</td>
</tr>
<tr>
<td>Automatic external defibrillator, with integrated electrocardiogram analysis, garment type</td>
<td>K0606</td>
</tr>
<tr>
<td>Replacement battery for automated external defibrillator, garment type only, each</td>
<td>K0607</td>
</tr>
<tr>
<td>Replacement garment for use with automated external defibrillator, each</td>
<td>K0608</td>
</tr>
<tr>
<td>Replacement electrodes for use with automated external defibrillator, garment type only, each</td>
<td>K0609</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Cardiac Devices - Transcatheter Aortic Valve Replacement (TAVR)</th>
<th>CODES:</th>
</tr>
</thead>
<tbody>
<tr>
<td>Transcatheter aortic valve replacement (TAVR/TAVI) with prosthetic valve; percutaneous femoral artery approach</td>
<td>33361</td>
</tr>
<tr>
<td>Transcatheter aortic valve replacement (TAVR/TAVI) with prosthetic valve; open femoral artery approach</td>
<td>33362</td>
</tr>
<tr>
<td>Transcatheter aortic valve replacement (TAVR/TAVI) with prosthetic valve; open axillary artery approach</td>
<td>33363</td>
</tr>
<tr>
<td>Transcatheter aortic valve replacement (TAVR/TAVI) with prosthetic valve; open iliac artery approach</td>
<td>33364</td>
</tr>
<tr>
<td>Transcatheter aortic valve replacement (TAVR/TAVI) with prosthetic valve; transaortic approach (eg, median sternotomy, mediastinotomy)</td>
<td>33365</td>
</tr>
<tr>
<td>Transcatheter aortic valve replacement (TAVR/TAVI) with prosthetic valve; transapical exposure (eg, left thoracotomy)</td>
<td>33366</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Cardiac Devices - Transcatheter Mitral Valve Repair (TMVR, MitraClip)</th>
<th>CODES:</th>
</tr>
</thead>
<tbody>
<tr>
<td>Transcatheter mitral valve repair, percutaneous approach, including transseptal puncture when performed; initial prosthesis</td>
<td>33418</td>
</tr>
<tr>
<td>Transcatheter mitral valve repair percutaneous approach via the coronary sinus</td>
<td>0345T</td>
</tr>
<tr>
<td>Cardiac Devices - Ventricular Assist Device (VAD)</td>
<td>CODES:</td>
</tr>
<tr>
<td>--------------------------------------------------</td>
<td>--------</td>
</tr>
<tr>
<td>Insertion of ventricular assist device, percutaneous including radiological supervision and interpretation; arterial access only</td>
<td>33990</td>
</tr>
<tr>
<td>Insertion of ventricular assist device, percutaneous including radiological supervision and interpretation; both arterial and venous access, with transseptal puncture</td>
<td>33991</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Cardiac Devices - Leadless Pacemakers (e.g., Micra Transcatheter Pacing System [TPS])</th>
<th>CODES:</th>
</tr>
</thead>
<tbody>
<tr>
<td>Transcatheter insertion or replacement of permanent leadless pacemaker, right ventricular, including imaging guidance (eg, fluoroscopy, venous ultrasound, ventriculography, femoral venography) and device evaluation (eg, interrogation or programming), when performed</td>
<td>33274</td>
</tr>
<tr>
<td>Transcatheter removal of permanent leadless pacemaker, right ventricular</td>
<td>33275</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Transthoracic Echocardiography (TTE)</th>
<th>CODES:</th>
</tr>
</thead>
<tbody>
<tr>
<td>Transthoracic echocardiography for congenital cardiac anomalies; complete</td>
<td>93303</td>
</tr>
<tr>
<td>Transthoracic echocardiography for congenital cardiac anomalies; follow-up or limited study</td>
<td>93304</td>
</tr>
<tr>
<td>Echocardiography, transthoracic, real-time with image documentation (2D), includes M-mode recording, when performed, complete, with spectral Doppler echocardiography, and with color flow Doppler echocardiography</td>
<td>93306</td>
</tr>
<tr>
<td>Echocardiography, transthoracic, real-time with image documentation (2D) with or without M-mode recording; complete</td>
<td>93307</td>
</tr>
<tr>
<td>Echocardiography, transthoracic, real-time with image documentation (2D) with or without M-mode recording; follow-up or limited study</td>
<td>93308</td>
</tr>
<tr>
<td>Transthoracic echocardiography with contrast, or without contrast followed by with contrast, for congenital cardiac anomalies; complete</td>
<td>C8921</td>
</tr>
<tr>
<td>Transthoracic echocardiography with contrast, or without contrast followed by with contrast, for congenital cardiac anomalies; follow-up or limited study</td>
<td>C8922</td>
</tr>
<tr>
<td>Transthoracic echocardiography with contrast, or without contrast followed by with contrast, real-time with image documentation (2D), includes M-mode recording, when performed, complete, without spectral or color doppler echocardiography</td>
<td>C8923</td>
</tr>
<tr>
<td>Transthoracic echocardiography with contrast, or without contrast followed by with contrast, real-time with image documentation (2D), includes M-mode recording, when performed, follow-up or limited study</td>
<td>C8924</td>
</tr>
<tr>
<td>Transthoracic echocardiography with contrast, or without contrast followed by with contrast, real-time with image documentation (2D), includes M-mode recording, when performed, complete, with spectral doppler echocardiography, and with color flow doppler echocardiography</td>
<td>C8929</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Transesophageal Echocardiography (TEE)</th>
<th>CODES:</th>
</tr>
</thead>
<tbody>
<tr>
<td>Echocardiography, transesophageal, real-time with image documentation (2D) (with or without M-mode recording); including probe placement, image acquisition, interpretation and report</td>
<td>93312</td>
</tr>
<tr>
<td>Echocardiography, transesophageal, real-time with image documentation (2D) (with or without M-mode recording); placement of transesophageal probe only</td>
<td>93313</td>
</tr>
</tbody>
</table>
### Echocardiography, transesophageal, real-time with image documentation (2D) (with or without M-mode recording); image acquisition, interpretation and report only

93314

### Transesophageal echocardiography for congenital cardiac anomalies; including probe placement, image acquisition, interpretation and report

93315

### Transesophageal echocardiography for congenital cardiac anomalies; placement of transesophageal probe only

93316

### Transesophageal echocardiography for congenital cardiac anomalies; image acquisition, interpretation and report only

93317

### Echocardiography, transesophageal (TEE) for monitoring purposes, including probe placement, real-time 2-dimensional image acquisition and interpretation leading to ongoing (continuous) assessment of (dynamically changing) cardiac pumping function and to therapeutic measures on an immediate time basis

93318

### Echocardiography, transesophageal (TEE) for guidance of a transcatheter intracardiac or great vessel(s) structural intervention(s) (eg, TAVR, transcatheter pulmonary valve replacement, mitral value repair, paravalvular regurgitation repair, left atrial appendage occlusion/closure, ventricular septal defect closure) (peri-and intraprocedural), real-time image acquisition and documentation, guidance with quantitative measurements, probe manipulation, interpretation and report, including diagnostic transesophageal echocardiography and, when performed, administration of ultrasound contrast, Doppler, color flow, and 3D

93355

### Transesophageal echocardiography (TEE) with contrast, or without contrast followed by with contrast, real time with image documentation (2D) (with or without M-mode recording); including probe placement, image acquisition, interpretation and report

C8925

### Transesophageal echocardiography (TEE) with contrast, or without contrast followed by with contrast, for congenital cardiac anomalies; including probe placement, image acquisition, interpretation and report

C8926

### Transesophageal echocardiography (TEE) with contrast, or without contrast followed by with contrast, for monitoring purposes, including probe placement, real time 2-dimensional image acquisition and interpretation leading to ongoing (continuous) assessment of (dynamically changing) cardiac pumping function and to therapeutic measures on an immediate time basis

C8927

### Stress Echocardiography

#### CODES:

**Echocardiography, transthoracic, real-time with image documentation (2D), includes M-mode recording, when performed, during rest and cardiovascular stress test using treadmill, bicycle exercise and/or pharmacologically induced stress, with interpretation and report**

93350

**Echocardiography, transthoracic, real-time with image documentation (2D), includes M-mode recording, when performed, during rest and cardiovascular stress test using treadmill, bicycle exercise and/or pharmacologically induced stress, with interpretation and report; including performance of continuous electrocardiographic monitoring, with physician supervision**

93351
Transthoracic echocardiography with contrast, or without contrast followed by with contrast, real-time with image documentation (2D), includes M-mode recording, when performed, during rest and cardiovascular stress test using treadmill, bicycle exercise and/or pharmacologically induced stress, with interpretation and report

<table>
<thead>
<tr>
<th>Transthoracic echocardiography, with contrast, or without contrast followed by with contrast, real-time with image documentation (2D), includes M-mode recording, when performed, during rest and cardiovascular stress test using treadmill, bicycle exercise and/or pharmacologically induced stress, with interpretation and report; including performance of continuous electrocardiographic monitoring, with physician supervision</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Angioplasty (Percutaneous Coronary Intervention [PCI])</strong></td>
</tr>
<tr>
<td><strong>Stent (Percutaneous Coronary Intervention [PCI])</strong></td>
</tr>
<tr>
<td><strong>Interventional Cardiology - Electrophysiological Studies (EPS) - Arrhythmia Induction and Mapping</strong></td>
</tr>
<tr>
<td>Transthoracic echocardiography, with contrast, or without contrast followed by with contrast, real-time with image documentation (2D), includes M-mode recording, when performed, during rest and cardiovascular stress test using treadmill, bicycle exercise and/or pharmacologically induced stress, with interpretation and report; including performance of continuous electrocardiographic monitoring, with physician supervision</td>
</tr>
</tbody>
</table>

### Angioplasty (Percutaneous Coronary Intervention [PCI])

**CODES:**

- Percutaneous transluminal coronary angioplasty; single major coronary artery or branch 92920

### Stent (Percutaneous Coronary Intervention [PCI])

**CODES:**

- Percutaneous transcatheter placement of intracoronary stent(s), with coronary angioplasty when performed; single major coronary artery or branch 92928
- Percutaneous transluminal revascularization of or through coronary artery bypass graft (internal mammary, free arterial, venous), any combination of intracoronary stent, atherectomy and angioplasty, including distal protection when performed; single vessel 92937
- Percutaneous transluminal revascularization of chronic total occlusion, coronary artery, coronary artery branch, or coronary artery bypass graft, any combination of intracoronary stent, atherectomy and angioplasty; single vessel 92943
- Percutaneous transcatheter placement of drug eluting intracoronary stent(s), with coronary angioplasty when performed; single major coronary artery or branch C9600
- Percutaneous transluminal revascularization of or through coronary artery bypass graft (internal mammary, free arterial, venous), any combination of drug-eluting intracoronary stent, atherectomy and angioplasty, including distal protection when performed; single vessel C9604
- Percutaneous transluminal revascularization of chronic total occlusion, coronary artery, coronary artery branch, or coronary artery bypass graft, any combination of drug-eluting intracoronary stent, atherectomy and angioplasty; single vessel C9607

### Interventional Cardiology - Electrophysiological Studies (EPS) - Arrhythmia Induction and Mapping

**CODES:**

- Bundle of HIS recording 93600
- Intra-atrial recording 93602
- Right ventricular recording 93603
- Intra-atrial pacing 93610
- Intraventricular pacing 93612
- Induction of arrhythmia by electrical pacing 93618
- Comprehensive electrophysiologic evaluation with right atrial pacing and recording, right ventricular pacing and recording, His bundle recording, including insertion and repositioning of multiple electrode catheters, without induction or attempted induction of arrhythmia 93619
Comprehensive electrophysiologic evaluation including insertion and repositioning of multiple electrode catheters with induction or attempted induction of arrhythmia; with right atrial pacing and recording, right ventricular pacing and recording, His bundle recording | 93620

Electrophysiologic follow-up study with pacing and recording to test effectiveness of therapy, including induction or attempted induction of arrhythmia | 93624

Intra-operative epicardial and endocardial pacing and mapping to localize the site of tachycardia or zone of slow conduction for surgical correction | 93631

Electrophysiologic evaluation of single or dual chamber pacing cardioverter-defibrillator leads including defibrillation threshold evaluation (induction of arrhythmia, evaluation of sensing and pacing for arrhythmia termination) at time of initial implantation or replacement | 93640

Electrophysiologic evaluation of single or dual chamber pacing cardioverter-defibrillator leads including defibrillation threshold evaluation (induction of arrhythmia, evaluation of sensing and pacing for arrhythmia termination) at time of initial implantation or replacement; with testing of single or dual chamber pacing | 93641

Electrophysiologic evaluation of single or dual chamber transvenous pacing cardioverter-defibrillator (includes defibrillation threshold evaluation, induction of arrhythmia, evaluation of sensing and pacing for arrhythmia termination, and programming or reprogramming of sensing or therapeutic parameters) | 93642

Electrophysiologic evaluation of subcutaneous implantable defibrillator (includes defibrillation threshold evaluation, induction of arrhythmia, evaluation of sensing for arrhythmia termination, and programming or reprogramming of sensing or therapeutic parameters) | 93644

**Interventional Cardiology - Electrophysiological Studies (EPS) - Ablation**

<table>
<thead>
<tr>
<th>Codes:</th>
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</thead>
</table>

Intracardiac catheter ablation of atrioventricular node function, atrioventricular conduction for creation of complete heart block, with or without temporary pacemaker placement | 93650

Comprehensive electrophysiologic evaluation including insertion and repositioning of multiple electrode catheters with induction or attempted induction of an arrhythmia with right atrial pacing and recording, right ventricular pacing and recording (when necessary), and His recording (when necessary) with intracardiac catheter ablation of arrhythmogenic focus; with treatment of supraventricular tachycardia by ablation of fast or slow atrioventricular pathway, accessory atrioventricular connection, cavotricuspid isthmus or other single atrial focus or source of atrial re-entry | 93653

Comprehensive electrophysiologic evaluation including insertion and repositioning of multiple electrode catheters with induction or attempted induction of an arrhythmia with right atrial pacing and recording, right ventricular pacing and recording (when necessary), and His recording (when necessary) with intracardiac catheter ablation of arrhythmogenic focus; with treatment of ventricular tachycardia or focus of ventricular ectopy including intracardiac electrophysiologic 3D mapping, when performed, and left ventricular pacing and recording, when performed | 93654
Comprehensive electrophysiologic evaluation including transseptal catheterizations, insertion and repositioning of multiple electrode catheters with induction or attempted induction of an arrhythmia including left or right atrial pacing/recording when necessary, right ventricular pacing/recording when necessary, and His bundle recording when necessary with intracardiac catheter ablation of atrial fibrillation by pulmonary vein isolation

<table>
<thead>
<tr>
<th>Mobile Cardiovascular Telemetry (MCT)</th>
<th>CODES:</th>
</tr>
</thead>
<tbody>
<tr>
<td>External mobile cardiovascular telemetry with electrocardiographic recording, concurrent computerized real time data analysis and greater than 24 hours of accessible ECG data storage (retrievable with query) with ECG triggered and patient selected events transmitted to a remote attended surveillance center for up to 30 days; review and interpretation with report by a physician or other qualified health care professional</td>
<td>93228</td>
</tr>
<tr>
<td>External mobile cardiovascular telemetry with electrocardiographic recording, concurrent computerized real time data analysis and greater than 24 hours of accessible ECG data storage (retrievable with query) with ECG triggered and patient selected events transmitted to a remote attended surveillance center for up to 30 days; technical support for connection and patient instructions for use, attended surveillance, analysis and transmission of daily and emergent data reports as prescribed by a physician or other qualified health care professional</td>
<td>93229</td>
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</tbody>
</table>

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<thead>
<tr>
<th>Implantable Loop Recorder</th>
<th>CODES:</th>
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</thead>
<tbody>
<tr>
<td>Insertion, subcutaneous cardiac rhythm monitor, including programming</td>
<td>33285</td>
</tr>
<tr>
<td>Removal, subcutaneous cardiac rhythm monitor</td>
<td>33286</td>
</tr>
</tbody>
</table>