PA.060.PW – Outpatient/Mobile Real Time Cardiac Surveillance Systems

This policy applies to the following lines of business:
✓ Piedmont Commercial

Piedmont WellStar HealthPlans considers outpatient/mobile real time cardiac surveillance medically necessary when all of the following indications are met (1-5):

1. The monitoring of a member is needed for **either** of the following:
2. Detection, characterization and documentation of symptomatic transient or paroxysmal dysrhythmia when the frequency of the symptoms is limited and the use of a 24-hour ambulatory ECG is documented in the medical record to be unlikely to capture and record the dysrhythmia, Prolonged monitoring is required specifically to ensure the absence of atrial fibrillation prior to the discontinuation of anticoagulation therapy Other testing and/or monitoring/recording/telemetry (e.g. ECG, 24 hour Holter, etc.) has been unrevealing There is a low likelihood of a potentially life-threatening cardiac event
3. It is anticipated that the results of this service would provide diagnostic and treatment information in the ongoing management of the member
4. Members needs cannot be met using an event recorder with loop memory and auto-trigger features

**Real-Time Cardiac Surveillance Requirements:**
1. Each member should have a recorder for their own exclusive use throughout the duration of the monitoring period. Recorders may not be shared between two members.
2. Monitoring is limited to once in a thirty day period and no more than one time in a twelve month period. In the event more time is needed, requests for longer intervals of monitoring must be accompanied by documentation that clearly supports the medical necessity of the continued surveillance.
3. This service should be ordered and interpreted by providers with experience in caring for these types of patients. These providers should also possess a thorough knowledge of the patient receiving the service.
4. Surveillance must occur continuously, 24 hours a day, 7 days a week while the member is wearing the device. This diagnostic test is ordered by the treating physician (or other treating practitioner acting within the scope of his or her license) who furnishes a consultation or who uses the results in the treatment and management of the member’s specific medical problem.

Services provided by an independent diagnostic testing facility (IDTF):

- The procedure must be performed under the general supervision of a physician specializing in cardiology or internal medicine.
- In general supervision, the procedure is furnished under the physician’s overall direction and control, but the physician’s presence is not required during the performance of the procedure.
- Under general supervision, the training of the non-physician personnel who actually perform the diagnostic procedure and the maintenance of the necessary equipment and supplies is the continuing responsibility of the physician.
- National or state-level training and certification requirements for non-physician personnel include:
  - Certified Cardiographic Technician (CCT) (Cardiovascular Credentialing International (CCI))
  - Registered nurse with current certification in Advanced Cardiac Life Support (ACLS)
  - Emergency Medical Technician (EMT) with current ACLS certification

Limitations

1. The concomitant use of cardiac surveillance, Holter monitoring, and /or event monitoring is considered not medically necessary

2. Real-time cardiac surveillance is not indicated for any of the following:
   a. Members with known or suspected dysrhythmias
   b. Outpatient monitoring of recently discharged post-infarct members
   c. Members at high risk of developing sustained ventricular tachycardia or ventricular fibrillation
   d. Members who would be more appropriately cared for in a hospital setting
   e. Use of cardiac surveillance and Holter or event monitoring for the same member on the same day
   f. Services performed for screening purposes
   g. Members with mild to moderate symptoms of palpitations, dizziness or weakness

3. Real-time cardiac surveillance is not considered medically necessary for all members with indications for cardiac monitoring. It should be used only in
circumstances where traditional Holter monitoring or cardiac event recording is not expected to provide adequate information.

Background
An electrocardiogram (EKG) is a graphic representation of electrical activity within the heart. Electrodes placed on the body in predetermined locations sense this electrical activity, which is then recorded by various means for review and interpretation. EKG recordings are used to diagnose a wide range of heart disease and other conditions that manifest themselves by abnormal cardiac electrical activity.

Ambulatory electrocardiography (AECG) refers to services rendered in an outpatient setting over a specified period of time, generally while a patient is engaged in daily activities, including sleep. AECG devices are intended to provide the physician with documented episodes of arrhythmia, which may not be detected using a standard 12-lead EKG. AECG is most typically used to evaluate symptoms that may correlate with intermittent cardiac arrhythmias and/or myocardial ischemia. Such symptoms include syncope, dizziness, chest pain, palpitations, or shortness of breath. Additionally, AECG is used to evaluate patient response to initiation, revision, or discontinuation of arrhythmic drug therapy. Descriptions of AECGs include:

- Dynamic electrocardiography devices that continuously record a real-time EKG, commonly known as Holter™ monitors, typically record over a 24-hour period
- An event monitor, or event recorder, is a patient-activated or event-activated EKG device that intermittently records cardiac arrhythmic events as they occur. These event monitors include pre-symptom memory loop and post-symptom recorders.

Real-time, outpatient cardiac telemetry involves the use of an automatically activated system that requires no patient intervention to either capture or transmit a dysrhythmia when it occurs. The purpose of this service is for real-time, continuous, long term (> 24 hours) cardiac surveillance of patients in order to identify and document a suspected and/or paroxysmal dysrhythmia.

Codes:

CPT Codes
PA.060.PW – Outpatient/Mobile Real Time Cardiac Surveillance Systems

Policy Number: PA.060.PW
Last Review Date: 09/10/2015
Effective Date: 01/01/2016
Renewal Date: 01/01/2017

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<tr>
<th>Code</th>
<th>Description</th>
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<tr>
<td>93228</td>
<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 (Report 93228 only once per 30 days)</td>
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<tr>
<td>93229</td>
<td>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 (Report 93229 only once per 30 days)</td>
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References


8. Joshi AK, Kowey PR, Prystowsky EN, et al. First experience with a mobile cardiac outpatient telemetry (MCOT) system for the diagnosis and management of cardiac arrhythmia. Am J Cardiol. 2005 Apr; 95(7):878-881. http://ac.els-cdn.com/S0002914905000147/1-s2.0-S0002914905000147-main.pdf?_tid=1bc95ae6-1b1b-11e3-a774-00000aacb361&acdnat=1378929079_46c4d32120d4a840fa0a81a834c7837b


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