This Molina Clinical Policy (MCP) is intended to facilitate the Utilization Management process. It expresses Molina's determination as to whether certain services or supplies are medically necessary, experimental, investigational, or cosmetic for purposes of determining appropriateness of payment. The conclusion that a particular service or supply is medically necessary does not constitute a representation or warranty that this service or supply is covered (i.e., will be paid for by Molina) for a particular member. The member's benefit plan determines coverage. Each benefit plan defines which services are covered, which are excluded, and which are subject to dollar caps or other limits. Members and their providers will need to consult the member's benefit plan to determine if there are any exclusion(s) or other benefit limitations applicable to this service or supply. If there is a discrepancy between this policy and a member's plan of benefits, the benefits plan will govern. In addition, coverage may be mandated by applicable legal requirements of a State, the Federal government or CMS for Medicare and Medicaid members. CMS's Coverage Database can be found on the CMS website. The coverage directive(s) and criteria from an existing National Coverage Determination (NCD) or Local Coverage Determination (LCD) will supersede the contents of this Molina Clinical Policy (MCP) document and provide the directive for all Medicare members.

**DESCRIPTION OF PROCEDURE/SERVICE/PHARMACEUTICAL**

Electroencephalography (EEG) is the recording of the brain's spontaneous electrical activity over a short period of time, usually 20–40 minutes, as recorded from multiple electrodes placed on the scalp. At times, a routine EEG is not sufficient, particularly when it is necessary to record an individual while they are having a seizure.

Ambulatory electroencephalography (AEEG) monitoring allows prolonged electroencephalographic (EEG) recording in the home setting, with a device slightly larger than a portable cassette player that continuously records brain wave patterns during 24 hours of a patient's routine daily activities and sleep. The device has the ability to record continuously for up to 72 hours increasing the chance of recording a seizure (i.e. ictal event, or the time between seizures, known as the interictal period). The interictal period is often used by neurologists when diagnosing epilepsy since an EEG trace will often show small interictal spiking and other abnormalities known by neurologists as subclinical seizures. The monitoring equipment consists of an electrode set, preamplifiers, and a cassette recorder. The electrodes attach to the scalp, and their leads are connected to a recorder, usually worn on a belt.

Video EEG monitoring is the simultaneous recording of clinical behavior and EEG. The purpose of the monitoring is to record seizure activity. The EEG is recorded continuously while a video camera records behavior. Video electroencephalography (EEG) monitoring is the synchronous recording and display of EEG patterns and video-recorded clinical behavior. Short recordings of several hours can be performed as an outpatient in an EEG laboratory, while longer recordings of 24 hours or more are generally done in a hospital inpatient setting. Since seizure medicine is often reduced
or stopped in order to provoke a seizure, the hospital setting is preferable so as to ensure the safety of the patient undergoing the seizure. The average hospital stay in the video EEG monitoring unit ranges from 3 to 4 days.

Video EEG monitoring is conducted for two main reasons: (1) for diagnostic monitoring when it is not clear from the clinical evaluation and routine EEG whether the patient has epileptic seizures or non-epileptic (psychiatric) events; and (2) for identifying the area of the brain from which seizures arise, especially for patients whose seizures are not controlled with antiepileptic medications and for whom surgery for epilepsy is being considered.

**INITIAL COVERAGE CRITERIA**

1. **Outpatient ambulatory encephalography (EEG) monitoring** may be considered medically necessary for up to 48 hours when all of the following criteria are met: [ALL]
   - Routine EEG, history and physical exam are inconclusive or non-diagnostic; and
   - Non-neurological causes of symptoms (e.g., syncope, cardiac arrhythmias) have been ruled out; and
   - Used to diagnose the following suspected conditions: [ONE]
     - Seizures or seizure like activity occurring ≥ 3 times per week;
     - To differentiate epileptic from non-epileptic events (i.e. a pseudo-seizure such as a psychogenic non-epileptic event, syncope or other);
     - To characterize the frequency or location of seizures in a nonclinical setting;
     - To document epilepsy response to treatment or to medication adjustment

2. **Out-patient video encephalography (EEG) monitoring** may be considered medically necessary when all of the following criteria are met: [ALL]
   - Used to diagnose the following suspected conditions: [ONE]
     - Know seizure disorder: [ALL]
       - recurrent refractory seizures despite treatment with ≥ 2 anticonvulsant medications; and
       - no current seizure provoking medications
     - Suspected non-epileptic seizure: [ALL]
       - recurrent symptoms not classic for seizures; and
       - history or lab results are non-diagnostic for seizure etiology; and
       - routine EEG non-specific; and
       - No sudden cessation of heavy alcohol use within 48 hours of seizure activity; and
       - No intoxication due to abuse of drugs within 48 hours of seizure activity
     - Prior to epilepsy surgery or intracranial electrode implantation and surgery to localize the seizure focus in members with documented medically refractory seizures.
   - If Anticonvulsant medication withdrawal is deemed unsafe in the outpatient setting (documentation is required), and ALL criteria above are met, then *inpatient video encephalography (EEG) monitoring may be authorized; and
   - Out-patient Video EEG length of stay is 23-72 hour observation and if the event being monitored does not occur in this time frame, then *inpatient video encephalography (EEG) monitoring may be authorized

*NOTE: For inpatient video encephalography (EEG) monitoring please use appropriate criteria for inpatient reviews that may include but is not limited to McKesson InterQual and/or Milliman MCG.*

**CONTINUATION OF THERAPY**

- Outpatient ambulatory EEG may be used for 48 hours.
Out-patient Video EEG length of stay is 23-72 hour observation but if the event being monitored does not occur in this time frame admission may be necessary for further monitoring or for preoperative localization of seizure foci.

Once the cause of seizures and specific type of epilepsy has been established, continued video EEG monitoring is considered not medically necessary.

**COVERAGE EXCLUSIONS**

Out-patient video encephalography (EEG) monitoring is considered NOT medically necessary in the home setting and is excluded due to insufficient evidence in the peer reviewed published literature.

**SUMMARY OF MEDICAL EVIDENCE**

**Ambulatory EEG Monitoring:**

The majority of the body of evidence published in the peer reviewed literature consists of retrospective and prospective cohort studies, retrospective studies and, retrospective reviews. The available literature shows the use of ambulatory EEG (AEEG) for diagnosis of epilepsy. Several studies focused on pediatric populations and/or video monitoring in addition to, or compared with, ambulatory EEG. Sample size ranged from 60 to >300 and included both children and adults. Follow up ranged from 1-3 years. The primary reasons for the AEEGs were to differentiate between seizures and non-epileptic events; to determine the frequency of seizures and epileptiform discharges; to characterize seizure type or localization; and to potentially diagnose epilepsy. Overall the majority of studies changed the diagnosis and refined the diagnosis by classifying the epilepsy into focal or generalized and confirmed the diagnostic utility of outpatient ambulatory EEG in the diagnosis of paroxysmal events.

**Video EEG Monitoring:**

The majority of the body of evidence published in the peer reviewed literature consists of retrospective and prospective cohort studies, retrospective studies and, retrospective reviews. Sample size ranged from 29 to >400 and mean age ranged from 35 to 52 years. Indications for referral included seizures requiring diagnostic clarification and presurgical evaluation. The duration of seizures across a lifetime was noted in 3 studies, with median/mean values of approximately 15 years with wide ranges. One study included patients that had a new-onset seizure. The studies reported that VEEG altered the diagnosis for some patients. The studies reported changes in antiepileptic drugs (AEDs) as a result of VEEG. The available evidence shows that video electroencephalogram (VEEG) monitoring can provide valuable information to guide the diagnosis and treatment of patients who report seizures, when a standard electroencephalogram (EEG) has not provided a clear diagnosis and in patients with refractory epilepsy. Studies have shown that VEEG changed the diagnosis from epileptic seizures to psychogenic non-epileptic seizures (PNES), and the vice versa. In addition, some of the studies reported elimination or reductions in medication after diagnostic changes.

**Professional Organizations**

*American Academy of Neurology, the Child Neurology Society, and the American Epilepsy Society,* published guidelines in 2000 (reaffirmed in 2014) for the evaluation of non-febrile seizures in children. This guideline recommends EEG testing as part of the neurodiagnostic evaluation of a child with an apparent first unprovoked seizure.

*American Academy of Neurology and the Practice Committee of the Child Neurology Society* published guidelines in 2006 (reaffirmed in 2013) for the diagnostic assessment of the child with status epilepticus (SE). This guideline has the following recommendations:
- An EEG may be considered in a child presenting with new onset SE as it may determine whether there are focal or generalized abnormalities that may influence diagnostic and treatment decisions.
- Although non-convulsive status epilepticus (NCSE) occurs in children who present with SE, there are insufficient data to support or refute recommendations regarding whether an EEG should be obtained to establish this diagnosis.
- An EEG may be considered in a child presenting with SE if the diagnosis of pseudo status epilepticus is suspected.

**CODING INFORMATION:** THE CODES LISTED IN THIS POLICY ARE FOR REFERENCE PURPOSES ONLY. LISTING OF A SERVICE OR DEVICE CODE IN THIS POLICY DOES NOT IMPLY THAT THE SERVICE DESCRIBED BY THIS CODE IS COVERED OR NON-COVERED. COVERAGE IS DETERMINED BY THE BENEFIT DOCUMENT. THIS LIST OF CODES MAY NOT BE ALL INCLUSIVE.

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<td>95950</td>
<td>Monitoring for identification and lateralization of cerebral seizure focus, electroencephalographic (eg. 8 channel EEG) recording and interpretation, each 24 hours</td>
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<td>95951</td>
<td>Monitoring for localization of cerebral seizure focus by cable or radio, 16 or more channel telemetry, combined electroencephalographic (EEG) and video recording and interpretation (eg, for presurgical localization ), each 24 hours</td>
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<td>95953</td>
<td>Monitoring for localization of cerebral seizure focus by computerized portable 16 or more channel EEG, electroencephalographic (EEG) recording and interpretation, each 24 hours, unattended</td>
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<td>95956</td>
<td>Monitoring for localization of cerebral seizure focus by cable or radio, 16 or more channel telemetry, electroencephalographic (EEG) recording and interpretation, each 24 hours</td>
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<td>Localization-related (focal) (partial) symptomatic epilepsy and epileptic syndromes with simple partial seizures, not intractable, with status epilepticus</td>
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<td>Localization-related (focal) (partial) symptomatic epilepsy and epileptic syndromes with simple partial seizures, not intractable, without status epilepticus</td>
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<tr>
<td>G40.311</td>
<td>Generalized idiopathic epilepsy and epileptic syndromes, intractable, with status epilepticus</td>
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18. Guldenb K., Baykan B. et al. The evaluation of the agreements of different epilepsy classifications in seizures recorded with video EEG monitoring. Journal of Neurological Sciences. 29 (2) (pp 201-211), 2012.

Other Resources
32. Hayes Heath Technology Assessment. Hayes, Inc. Winifred Hayes Inc. Lansdale, PA
   - Video Electroencephalogram (VEEG) for Diagnosis and Management of Epilepsy in Adults. Winifred. Updated Sept 2017.
   - Video Electroencephalogram (VEEG) for Diagnosis and Management of Epilepsy in Children. Winifred Hayes, Inc. Winifred Hayes Inc. Lansdale, PA. Updated Aug 2017. [archived Nov, 2018]
   - Hayes Search & Summary. Ambulatory Electroencephalography (EEG) for Diagnosis of Epilepsy.
   - Video Electroencephalograph (VEEG) Monitoring in the Home Setting for the
   - Diagnosis of Epilepsy in Children. May, 2016.
33. McKesson InterQual 2018:
   - Procedures Video Electroencephalographic (EEG) Monitoring, Adult & Pediatric.
   - Acute Pediatric Epilepsy Criteria
• Acute Adult Epilepsy Criteria
  35. Advanced Medical Review:
  • Policy reviewed by MD board certified in pediatrics. 2013
  • Policy reviewed by MD board certified in neurology. 2013
  • Policy reviewed by practicing physician board certified in neurology. May 6, 2018.

Revision/Review History:
3/1/16: This policy was reviewed and the criteria for ambulatory and video EEG was updated. Routine EEG, history and physical exam, and outpatient sleep study with EEG monitoring are inconclusive or non-diagnostic was included under ambulatory EEG criteria and MRI criteria for video EEG changed to only be required in cases of a suspected non-epileptic seizure when history and lab results are either normal or non-diagnostic for etiology of symptoms or findings. MRI is not required prior to epilepsy surgery or in a case of known seizure disorder.
5/3/16: The only revision was that the following sentence was added into the exclusion section of the policy: Out-patient video encephalography (EEG) monitoring is considered NOT medically necessary in the home setting and is excluded due to insufficient evidence in the peer reviewed published literature.
9/19/16: Changed video EEG criteria to require treatment with > 2 anticonvulsant medications instead of requiring therapeutic levels of > anticonvulsant medications and changed Out-patient Video EEG length of stay to up to 72 hour observation instead of 48 hours.
6/22/17: Policy reviewed, clinical criteria has not changed.
3/8/18: Policy reviewed, clinical criteria has not changed.
7/10/18: Policy reviewed and clinical criteria has changed according to AMR review: Under Ambulatory EEG criteria removed the criteria for outpatient sleep study with EEG monitoring and defined a non-epileptic events to include the following language: (i.e. a pseudo-seizure such as a psychogenic non-epileptic event, syncope or other).
9/18/19: Policy reviewed, clinical criteria has not changed.