Vestibular Function Testing

Policy #  00556
Original Effective Date:  04/19/2017
Current Effective Date:  04/19/2017

Applies to all products administered or underwritten by Blue Cross and Blue Shield of Louisiana and its subsidiary, HMO Louisiana, Inc.(collectively referred to as the “Company”), unless otherwise provided in the applicable contract. Medical technology is constantly evolving, and we reserve the right to review and update Medical Policy periodically.

When Services May Be Eligible for Coverage
Coverage for eligible medical treatments or procedures, drugs, devices or biological products may be provided only if:

- Benefits are available in the member’s contract/certificate, and
- Medical necessity criteria and guidelines are met.

Based on review of available data, the Company may consider vestibular function testing using an electronystagmography (ENG) and videonystagmography (VNG) testing batteries, caloric testing, or rotational chair testing to be eligible for coverage when the following conditions have been met:

Patient Selection Criteria
Coverage eligibility will be considered when all of the following criteria have been met:

- The patient has symptoms of a vestibular disorder (eg, dizziness, vertigo, imbalance); AND
- A clinical evaluation, including maneuvers such as the Dix-Hallpike test if indicated, has failed to identify the cause of the symptoms.

When Services Are Considered Not Medically Necessary
Based on review of available data, the Company considers the use of vestibular function testing for the assessment of typical benign paroxysmal positional vertigo that can be diagnosed clinically to be not medically necessary.**

Based on review of available data, the Company considers repeat vestibular function testing when treatment resolves symptoms to be not medically necessary.**

When Services Are Considered Investigational
Coverage is not available for investigational medical treatments or procedures, drugs, devices or biological products.

Based on review of available data, the Company considers vestibular function testing in all other situations to be investigational.*

Based on review of available data, the Company considers vestibular evoked myogenic potential (VEMP) tests to be investigational.*

Based on review of available data, the Company considers all other laboratory-based vestibular function tests not described above to be investigational.*
Background/Overview

VERTIGO

The vestibular system controls balance. It includes 5 end organs, 3 semicircular canals sensitive to head rotations, and 2 otolith organs (saccule, utricle) that sense gravity and straight-line (forward, backward, left, right, downward or upward) accelerations. Vertigo is the primary symptom of vestibular dysfunction. It can be experienced as illusory movement such as spinning, swaying, or tilting. Vertigo may be associated with a feeling of being pushed or pulled to the ground, blurred vision, nausea and vomiting, or postural and gait instability. Vertigo may arise from damage or dysfunction of the vestibular labyrinth, vestibular nerve, or central vestibular structures in the brainstem.

Vertigo may be caused by loose particles (otoconia) from the otolith organs that pass into one of the semicircular canals, most frequently the posterior canal. Specific head movements cause the particle to stimulate the canal, causing brief benign paroxysmal positional vertigo (BPPV). BPPV can usually be diagnosed clinically based on history of positional vertigo, response to the Dix-Hallpike maneuver or lateral roll tests, and resolution of symptoms with canal repositioning maneuvers.

Testing

If vertigo cannot be attributed to BPPV based on history, symptoms, or response to the standard maneuvers, a number of laboratory-based tests can be used to determine whether the vertigo is due to loss of vestibular function. These tests are based on the vestibulo-ocular reflex, which is an involuntary beating movement of the eyes (nystagmus) in response to vestibular stimulation. Nystagmus induced by these tests can help to distinguish between central and peripheral etiologies, in addition to determining whether the deficit is unilateral or bilateral. The typical tests include the ENG or VNG test batteries, caloric testing, and rotational chair testing.

ENG/VNG Test Batteries

The ENG/VNG test batteries include oculomotor evaluation and positional testing. ENG uses electrodes at the canthus of the eyes to detect nystagmus while VNG uses infrared video monitoring with goggles to measure nystagmus.

Caloric Testing

Caloric testing evaluates unilateral vestibular function. In the caloric test, warm or cold water or warm or cold air, is introduced into each of the external ear canals. In some descriptions, caloric testing is conducted as part of ENG/VNG test batteries.

Rotational Chair Testing

The rotational chair test evaluates bilateral vestibular function. Rotational chair devices include a lightproof booth, computer-driven chair with a head restraint that rotates around a vertical axis, ENG recording, an infrared camera, and a 2-way communication system. Typically, the chair is rotated in 4 different patterns, constant acceleration followed by deceleration, rotating followed by a rapid stop, rotating at progressively increasing velocities, and alternating directions.
Passive rotational testing without a rotational chair may be performed when the rotational chair is not available. For the head impulse test, the patient is instructed to keep his or her eyes on a target. The examiner then turns the head rapidly by about 15°. With passive whole body testing the examiner rotates the whole body to the rhythm of a metronome.

**Vestibular Evoked Myogenic Potential Testing**

Vestibular evoked myogenic potential tests are newer techniques that use loud sound (eg, click, tone burst) or bone vibration (eg, tendon hammer tap to the forehead or mastoid) to assess otolith function. Both the saccule and utricle are sensitive to sound as well as vibration and movement.

Cervical VEMPs (cVEMPS) are measured by surface electrodes on the ipsilateral sternocleidomastoid (SCM) muscle in the neck and are thought to originate primarily in the saccule. The cVEMP response was first described more than 20 years ago. Abnormality in any part of the auditory cVEMP pathway (saccule, inferior vestibular nerve, vestibular nucleus, medial vestibulospinal tract, the accessory nucleus, the eleventh nerve, SCM) can affect the response.

Ocular VEMPs (oVEMPs) detect subtle activity of an extraocular muscle using surface electrodes under the contralateral eye during an upward gaze, and are thought to be due primarily to stimulation of the utricle. The vestibulo-ocular reflex stimulated by sound or vibration is very small, but synchronous bursts of activity of the extraocular muscles can be detected by electromyography. Lesions that affect the oVEMP may occur in the utricle, superior vestibular nerve, vestibular nucleus, and the crossed vestibulo-ocular reflex pathways.

**Dynamic Posturography**

Dynamic posturography may also be used to evaluate balance.

**Treatment**

The central vestibular system is able to compensate for loss of peripheral vestibular function. Thus, the primary therapy for peripheral vestibular dysfunction is exercise-based and includes exercises to promote gaze stability, habituate symptoms, and improve balance and gait. Medications such as vestibular suppressants or antiemetics may be used in the acute stage but are not recommended for chronic use. For patients who have recurrent symptoms uncontrolled by other methods, a surgical or ablative approach may be used. The objective of the ablative approach is to stabilize the deficit to allow central compensation.

**FDA or Other Governmental Regulatory Approval**

U.S. Food and Drug Administration (FDA)

Vestibular analysis devices are currently regulated by the FDA through the 510(k) pathway, under the product code LXV. The term “vestibular analysis devices” includes both diagnostic devices (eg, rotary chairs, multiaxial chairs) and therapeutic devices (eg, balance training and balance rehabilitation devices). Some devices indicated for diagnostic testing are included in Table 1.
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Table 1. Vestibular Analysis Devices Approved by the Food and Drug Administration

<table>
<thead>
<tr>
<th>Trade Name</th>
<th>Manufacturer (510k applicant)</th>
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<th>Date</th>
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<tbody>
<tr>
<td>ICS Impulse‡‡</td>
<td>Otometrics</td>
<td>K122550</td>
<td>2013</td>
</tr>
<tr>
<td>Sway Balance‡‡</td>
<td>Sway Medical (Capacity Sports)</td>
<td>K121590</td>
<td>2012</td>
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<td>Nydiag 200 Rotary Chair</td>
<td>Interacoustics A/S</td>
<td>K102364</td>
<td>2010</td>
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<tr>
<td>Epley Omninax‡‡</td>
<td>Vesticon</td>
<td>K071973</td>
<td>2008</td>
</tr>
<tr>
<td>VMT System</td>
<td>Target Health</td>
<td>K971549</td>
<td>1998</td>
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<tr>
<td>VORTEQ‡‡ (Vestibular Ocular Reflex Test Equipment)</td>
<td>Micromedical Technologies</td>
<td>K991008</td>
<td>1999</td>
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<td>Chair, Vestibular, Rotary, Computerized</td>
<td>Contraves</td>
<td>K781268</td>
<td>1987</td>
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<td>RVT-50 Rotary Chair for Vestibular Testing</td>
<td>ICS Medical</td>
<td>K872093</td>
<td>1987</td>
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<tr>
<td>EquiTest‡‡</td>
<td>Natus Medical (NeuroCom International)</td>
<td>K851744</td>
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An example of equipment used for vestibular evoked myogenic potentials is the Bio-Logic Nav-Pro (BioLogic Systems Corp), which in 2003 was cleared for marketing by FDA through the 510(k) process (K994149) for use in the recording and displaying human physiologic data, for auditory screening and assisting in evaluation of auditory and hearing-related disorders using auditory brainstem responses recorded from electroencephalography electrodes placed on the scalp.

Centers for Medicare and Medicaid Services (CMS)
There is no national coverage determination (NCD). In the absence of an NCD, coverage decisions are left to the discretion of local Medicare carriers.

**Rationale/Source**
Laboratory-based vestibular function testing is well-established and has a large evidence base. In a 2000 technology assessment, the American Academy of Neurology (AAN) evaluated tests that stimulate the vestibular system (see Table 2). AAN included caloric irrigation and rotational chair testing as established as effective, with passive examiner-generated head rotation testing and active head rotation as probably effective but not yet fully accepted by expert consensus. AAN noted that quantitative vestibular testing is not always necessary, and a number of bedside methods can be used to evaluate nystagmus.

**ELECTRONYSTAGMOGRAPHY AND VIDEONYSTAGMOGRAPHY TEST BATTERIES**
The basic ENG and VNG test batteries include a spontaneous nystagmus test that measures the ability of the eyes to maintain a fixed position, a positional nystagmus test that measures the ability of the eyes to maintain a static position when the head is in different positions, an optokinetic nystagmus test that measures nystagmus caused by viewing a series of targets moving to the right and then to the left, and an oscillating tracking test that evaluates patient ability to track a moving target. The basic ENG/VNG test batteries with these 4 tests are well-established for evaluating vestibular function in patients who have a suspected vestibular disorder.

**CALORIC TESTING**
Caloric testing is the most widely used vestibular function test and is considered the criterion standard for detecting unilateral vestibular loss. When warm or cold water or air is introduced into one of the external ear
canals, the temperature change is transmitted through the middle ear and bone, causing a thermal gradient in the semicircular canal and resulting in nystagmus. Cold water will cause a movement response of the eye opposite to the stimulation, while warm water will induce nystagmus in the direction of the ear being stimulated. These eye movements can be measured by electrodes at the canthus or by video monitoring. An asymmetrical response after stimulating both ears indicates unilateral vestibular dysfunction.

**ROTATIONAL CHAIR TESTING**
Rotational chair testing is considered the gold standard for detecting bilateral vestibular loss. Rotational chair devices include a lightproof booth, computer-driven chair with a head restraint that rotates around a vertical axis, ENG recording, an infrared camera, and a 2-way communication system. Typically, the chair is rotated in 4 different patterns, constant acceleration followed by deceleration, rotating followed by a rapid stop, rotating at progressively increasing velocities, and alternating directions. Each pattern is repeated in both directions several times, and the accompanying post-rotation nystagmus, including parameters of gain, phase, and symmetry, is measured and averaged. Although traditionally used to detect bilateral vestibular loss, this battery can identify a unilateral vestibular deficit and identify the site of the lesion.

**VESTIBULAR EVOKED MYOGENIC POTENTIAL TESTING**
Vestibular evoked myogenic potential tests use sound or vibration to stimulate the otolith organs. Cervical VEMP measures evoked electrical potentials in the ipsilateral sternocleidomastoid (SCM) muscle following stimulation of the saccule, while oVEMP measures electrical potentials in the extraocular muscles contralateral to the utricle. There is a large and rapidly growing literature on VEMPs for the assessment of otolith function, although most studies assess how cVEMP and oVEMP change with various disease states. VEMPs have been evaluated in superior canal dehiscence, vestibular neuritis, BPPV, vestibular schwannoma, Meniere disease, vestibular migraine, and central vestibular disorders. There are a number of concerns about using VEMPs to assess the otolith organs. One issue is that sound and bone conduction stimuli are likely to influence senses other than the saccule and utricle, and stimulation of structures other than the utricle can affect the VEMP. Another is that latency and amplitude measures are very sensitive to variables that can be introduced by the examiner. In addition, VEMP responses have been shown to decrease with age, with a high rate of absent responses in normal older adults. A 2014 expert consensus document on cVEMP methods has stated that the clinical use of VEMP “is evolving and questions still exist about its underlying physiology and measurement,” while a 2016 article on the impact of examiners concluded that the field should “work on a better standard for VEMP recordings”. Together, the available evidence has indicated that the use of VEMP tests to evaluate suspected vestibular disorders is at a very early stage of development. Standardization of procedures and studies on the diagnostic accuracy of these procedures are needed.

**LABORATORY-BASED VESTIBULAR FUNCTION TESTS FOR CLINICALLY DIAGNOSED BENIGN PAROXYSMAL POSITIONAL VERTIGO**
BPPV with a typical presentation is usually diagnosed clinically with a combination of a history of periods of brief positional vertigo, recurrence of symptoms with the Dix-Hallpike maneuver or lateral roll procedures, and/or alleviation of symptoms after canal repositioning maneuver. The Dix-Hallpike maneuver is the criterion standard for the diagnosis of posterior canal BPPV, limiting evaluation of its performance characteristics. The 2008 practice guidelines from the American Academy of Otolaryngology – Head and
Neck Surgery (AAO-HNS) gave a strong recommendation for the diagnosis of BPPV of the posterior canal when vertigo associated with nystagmus has been provoked by the Dix-Hallpike maneuver. If the Dix-Hallpike maneuver is negative, but the history is consistent with BPPV, a lateral roll test can be used to assess BPPV of the horizontal canal. In the event that both the Dix-Hallpike maneuver and lateral roll tests are negative, alleviation of symptoms with the canal repositioning maneuver supports a diagnosis of BPPV. AAO-HNS has recommended against vestibular testing in patients who meet clinical criteria for the diagnosis of BPPV. If the clinical presentation is atypical, if Dix-Hallpike testing elicits equivocal or unusual nystagmus findings, if symptoms do not resolve following treatment, or if there are additional symptoms or signs, vestibular function testing may be indicated.

**SUMMARY OF EVIDENCE**

For individuals who have a suspected vestibular disorder not clinically diagnosed as benign paroxysmal positional vertigo who receive ENG and VNG test batteries, caloric testing, or rotational chair testing, the evidence includes technology assessments of a large body of literature. Relevant outcomes are test accuracy, symptoms, functional outcomes, and quality of life. Based on review of controlled studies, caloric testing was given a level A recommendation that this test was predictive of loss of vestibular function. Based on a prospective study assessing a narrow spectrum of patients with the suspected vestibular dysfunction, or well-designed retrospective study compared with the criterion standard test, rotational chair testing was also given a level A recommendation. These tests are both considered criterion standard tests of vestibular function. ENG/VNG test batteries, which may include caloric testing, are also established methods of assessing loss of vestibular function. The evidence is sufficient to determine that the technology results in a meaningful improvement in the net health outcome.

For individuals who have a suspected vestibular disorder not clinically diagnosed as benign paroxysmal positional vertigo who receive a VEMP testing, the evidence includes mainly association studies. Relevant outcomes are test accuracy, symptoms, functional outcomes, and quality of life. There is a large and rapidly growing literature on VEMP tests for the assessment of otolith function, although most studies have assessed how the cervical VEMP and ocular VEMP change with various disease states. Studies on diagnostic accuracy and clinical utility of this technique for evaluating otolith organs and central pathways are needed. The evidence is insufficient to determine the effects of the technology on health outcomes.

For individuals who have clinically diagnosed BPPV with typical presentation who receive laboratory-based vestibular function tests, the evidence includes technology assessments and practice guidelines. Relevant outcomes are test accuracy, symptoms, functional outcomes, and quality of life. BPPV with a typical presentation can be diagnosed clinically based on history, the Dix-Hallpike maneuver, lateral roll test, and canalith repositioning procedures; thus, laboratory-based vestibular function tests do not add diagnostic information in such routine cases. The evidence is sufficient to determine that the technology is unlikely to improve the net health outcome.

**References**

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Policy # 00556
Original Effective Date: 04/19/2017
Current Effective Date: 04/19/2017


Policy History
Original Effective Date: 04/19/2017
Current Effective Date: 04/19/2017
04/06/2017 Medical Policy Committee review
04/19/2017 Medical Policy Implementation Committee approval. New policy.
Next Scheduled Review Date: 04/2018

Coding
The five character codes included in the Blue Cross Blue Shield of Louisiana Medical Policy Coverage Guidelines are obtained from Current Procedural Terminology (CPT®), copyright 2016 by the American Medical Association (AMA). CPT is developed by the AMA as a listing of descriptive terms and five character identifying codes and modifiers for reporting medical services and procedures performed by physician.

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Codes used to identify services associated with this policy may include (but may not be limited to) the following:

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<th>Code Type</th>
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<td>92537, 92538, 92540, 92541, 92542, 92544, 92545, 92546, 92547, 92700</td>
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<td>HCPCS</td>
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<tr>
<td>ICD-10 Diagnosis</td>
<td>A88.1, H81.01-H82.9, R42</td>
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*Investigational – A medical treatment, procedure, drug, device, or biological product is Investigational if the effectiveness has not been clearly tested and it has not been incorporated into standard medical practice. Any determination we make that a medical treatment, procedure, drug, device, or biological product is Investigational will be based on a consideration of the following:

A. Whether the medical treatment, procedure, drug, device, or biological product can be lawfully marketed without approval of the U.S. FDA and whether such approval has been granted at the time the medical treatment, procedure, drug, device, or biological product is sought to be furnished; or
B. Whether the medical treatment, procedure, drug, device, or biological product requires further studies or clinical trials to determine its maximum tolerated dose, toxicity, safety, effectiveness, or effectiveness as compared with the standard means of treatment or diagnosis, must improve health outcomes, according to the consensus of opinion among experts as shown by reliable evidence, including:
   1. Consultation with the Blue Cross and Blue Shield Association technology assessment program (TEC) or other nonaffiliated technology evaluation center(s);
   2. Credible scientific evidence published in peer-reviewed medical literature generally recognized by the relevant medical community; or
   3. Reference to federal regulations.

**Medically Necessary (or “Medical Necessity”) - Health care services, treatment, procedures, equipment, drugs, devices, items or supplies that a Provider, exercising prudent clinical judgment, would provide to a patient for the purpose of preventing, evaluating, diagnosing or treating an illness, injury, disease or its symptoms, and that are:
   A. In accordance with nationally accepted standards of medical practice;
   B. Clinically appropriate, in terms of type, frequency, extent, level of care, site and duration, and considered effective for the patient's illness, injury or disease; and
   C. Not primarily for the personal comfort or convenience of the patient, physician or other health care provider, and not more costly than an alternative service or sequence of services at least as likely to produce equivalent therapeutic or diagnostic results as to the diagnosis or treatment of that patient's illness, injury or disease.

For these purposes, “nationally accepted standards of medical practice” means standards that are based on credible scientific evidence published in peer-reviewed medical literature generally recognized by the relevant medical community, Physician Specialty Society recommendations and the views of Physicians practicing in relevant clinical areas and any other relevant factors.

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