Vestibular Neuritis

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Vestibular neuritis refers to a disorder characterized by acute, isolated, spontaneous vertigo due to unilateral vestibular deafferentiation.1 Even though the clinical features had been described previously, it was Dix and Hallpike who first coined the term vestibular neuronitis in 1952 to distinguish it from Ménière’s disease.2

Vestibular neuritis accounts for 3.2 to 9% of the patients visiting a dizziness center,3,4 and has an incidence of ~3.5 per 100,000 population.5 Characteristic symptoms of acute vestibular neuritis include vertigo, nausea/vomiting, oscillopsia, and unsteadiness.6 Patients with vestibular neuritis show spontaneous horizontal-torsional nystagmus beating away from the lesion side, abnormal head impulse test (HIT) for the involved semicircular canals, ipsilesional caloric paresis, decreased responses of vestibular-evoked myogenic potentials (VEMPs) during stimulation of the affected ear, and unsteadiness with a falling tendency toward the lesion side. Vestibular neuritis preferentially involves the superior vestibular labyrinth and its afferents. Accordingly, the function of the posterior semicircular canal and saccule, which constitute the inferior vestibular labyrinth, is mostly spared in vestibular neuritis. However, because the rare subtype of inferior vestibular neuritis lacks the typical features of vestibular neuritis, it may be misdiagnosed as a central vestibular disorder. Even in the patient with the typical pattern of spontaneous nystagmus observed in vestibular neuritis, brain imaging is indicated when the patient has unprecedented headache, negative head impulse test, severe unsteadiness, or no recovery within 1 to 2 days. Symptomatic medication is indicated only during the acute phase to relieve the vertigo and nausea/vomiting. Vestibular rehabilitation hastens the recovery. The efficacy of topical and systemic steroids requires further validation.

Abstract

Vestibular neuritis is the most common cause of acute spontaneous vertigo. Vestibular neuritis is ascribed to acute unilateral loss of vestibular function, probably due to reactivation of herpes simplex virus in the vestibular ganglia. The diagnostic hallmarks of vestibular neuritis are spontaneous horizontal-torsional nystagmus beating away from the lesion side, abnormal head impulse test for the involved semicircular canals, ipsilesional caloric paresis, decreased responses of vestibular-evoked myogenic potentials during stimulation of the affected ear, and unsteadiness with a falling tendency toward the lesion side. Vestibular neuritis preferentially involves the superior vestibular labyrinth and its afferents. Accordingly, the function of the posterior semicircular canal and saccule, which constitute the inferior vestibular labyrinth, is mostly spared in vestibular neuritis. However, because the rare subtype of inferior vestibular neuritis lacks the typical features of vestibular neuritis, it may be misdiagnosed as a central vestibular disorder. Even in the patient with the typical pattern of spontaneous nystagmus observed in vestibular neuritis, brain imaging is indicated when the patient has unprecedented headache, negative head impulse test, severe unsteadiness, or no recovery within 1 to 2 days. Symptomatic medication is indicated only during the acute phase to relieve the vertigo and nausea/vomiting. Vestibular rehabilitation hastens the recovery. The efficacy of topical and systemic steroids requires further validation.

Keywords

► vestibular neuritis
► vertigo
► nystagmus
► head impulse test
► vestibular evoked myogenic potential

Vestibular neuritis refers to a disorder characterized by acute, isolated, spontaneous vertigo due to unilateral vestibular deafferentiation.1 Even though the clinical features had been described previously, it was Dix and Hallpike who first coined the term vestibular neuronitis in 1952 to distinguish it from Ménière’s disease.2

Vestibular neuritis accounts for 3.2 to 9% of the patients visiting a dizziness center,3,4 and has an incidence of ~3.5 per 100,000 population.5 Characteristic symptoms of acute vestibular neuritis include vertigo, nausea/vomiting, oscillopsia, and unsteadiness.6 Patients with vestibular neuritis show spontaneous horizontal-torsional nystagmus beating away from the lesion side, abnormal head impulse test (HIT) for the involved semicircular canals, ipsilesional caloric paresis, decreased responses of vestibular-evoked myogenic potentials (VEMPs) during stimulation of the affected ear, and unsteadiness with a falling tendency toward the lesion side.7

The vestibular labyrinth may be subdivided into the superior and inferior divisions (►Fig. 1). The superior vestibular labyrinth comprises the anterior (AC) and horizontal semicircular canals (HC), the utricle, and their afferents. In contrast, the inferior vestibular labyrinth consists of the posterior semicircular canal (PC), the saccule, and their afferents. The head impulse test enables us to evaluate the function of each semicircular canal,8,9 and VEMPs that reflect the function of the utricle and saccule10,11 are now able to securely diagnose the three distinctive patterns of vestibular neuritis, the superior, inferior, and total (superior + inferior) types.

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Because vestibular neuritis preferentially affect the superior division, superior vestibular neuritis is most common (55–100%), followed by total (15–30%) and inferior vestibular neuritis (3.7–15%).

Although vestibular neuritis itself is not life threatening, distinguishing vestibular neuritis from other debilitating disorders such as strokes is essential. The management of vestibular neuritis includes (1) symptomatic care for nausea/vomiting and vertigo, (2) etiological treatments, and (3) vestibular rehabilitation to improve central compensation.

Clinical Features

Patients with vestibular neuritis mostly present with subacute or acute spontaneous vertigo with nausea/vomiting and unsteadiness. The symptoms may develop suddenly or evolve over several hours. Prodromal dizziness may be reported in ~8.6 to 24% of patients. Most (74%) of the preceding episodes are described as nonvertiginous dizziness, often associated with a feeling of nausea or unsteadiness. The episodes may have abrupt or gradual onset, and may last a few minutes to days. Patients may have precedent or concurrent viral illnesses.

The vertigo in vestibular neuritis may gradually increase over several hours and reach a peak within the first day. Vertigo is usually described as rotational, and is markedly increased by head movements. Patients usually prefer to lie in bed with their eyes closed in a side position with the healthy ear down. Most patients suffer from severe nausea and vomiting. Severe vertigo improves markedly over 1 to 2 days.

Because stimulation of a single semicircular canal induces eye movements approximately in the plane of that canal, the patterns of spontaneous nystagmus in vestibular neuritis would depend on relative involvement of each semicircular canal. Accordingly, when all three canals are completely damaged, the nystagmus is mixed horizontal-torsional and the fast component of the spontaneous nystagmus is away from the affected side. However, the spontaneous nystagmus invariably accompanies a vertical component that is mostly upbeat because AC is more commonly affected than PC. As the nystagmus is markedly suppressed by visual fixation, proper observation of the nystagmus requires a removal of visual fixation using Frenzel glasses or infrared video goggles. The nystagmus increases during the gaze in the direction of the nystagmus and decreases when looking into the other direction (Alexander’s law), but never changes directions. The nystagmus is mostly augmented by horizontal head shaking, vibratory stimuli on the mastoids or the brow, or hyperventilation.

The spontaneous nystagmus gives rise to illusion of apparent movement of the environment (oscillopsia). Because the illusory visual perception of rotation is produced by the slow phase of the nystagmus in the opposite direction, right-beating spontaneous nystagmus in left vestibular neuritis induces contraversive rightward (clockwise) rotation of the surroundings.

Patients tend to fall toward the affected side on standing with the feet together or on attempted walking. However, they should maintain sitting balance or stand unaided with the feet apart because the brain can still utilize the information for balance control via the visual and somatosensory systems.

Etiology

Although precedent or concurrent viral illnesses have been described in vestibular neuritis, evidence of systemic viral infection based on seroconversion remain unconvincing. The signs and symptoms confined to the vestibular nerve in most series do not support systemic viral infection either. Instead, increasing evidence suggests possible reactivation of
latent type 1 herpes simplex virus (HSV-1) as a cause of vestibular neuritis, likewise in idiopathic facial paresis. HSV-1 DNA is detected on autopsy in about two of three human vestibular ganglia along with the expression of CD8-positive T lymphocytes, cytokines, and chemokines. These findings indicate latent infection of the vestibular ganglia with HSV-1. Furthermore, inoculation of HSV induces vestibular dysfunction with infected vestibular ganglion cells in mice. Other possible mechanisms include autoimmune and microvascular ischemic insults to the vestibular labyrinth.

Vestibular neuritis preferentially affects the superior division. In other words, vestibular neuritis is most likely partial with a predominant involvement of AC, HC, and the utricle. This may be ascribed to the anatomical differences between the superior and inferior vestibular nerves. The bony canal of the superior vestibular nerve is seven times longer than the singular nerve canal. Furthermore, the superior vestibular nerve and arteriole travel through a relatively narrower passage than the singular nerve and its vascular supply because there are more bony spicules occupying the superior vestibular compared with the inferior vestibular or singular channels. Therefore, the superior nerve may be more susceptible to swelling or ischemic insults. Sparing of PC function in vestibular neuritis could also be explained by the relatively common double innervations of PC by two distinct nerves, which reach the posterior cupula through separate bony canals.

### Evaluation

During the acute phase, the spontaneous nystagmus may be seen even with visual fixation, but is better appreciated with removal of visual fixation. Nystagmus can be recorded and quantitatively analyzed using oculography. Three-dimensional video-oculography is commonly used, which can measure the direction and slow-phase velocity of spontaneous nystagmus. Nystagmus may be induced or modulated by various maneuvers such as head-shaking, vibration, or hyperventilation. Head-shaking nystagmus is mostly assessed by using a passive head-shaking maneuver. Vibration-induced nystagmus is recorded by applying a vibration stimulator to the forehead and both mastoids.

A head impulse test can evaluate the function of each semicircular canal bedside. A bedside head impulse test is feasible, and the sensitivity is clinically acceptable. However, a bedside head impulse test may be negative, especially when the vestibular deficits are partial or the corrective saccades are covert, i.e., the corrective saccades occur only during head impulse rather than after the head rotation. Furthermore, bedside head impulse tests in the planes of the vertical canals are more difficult to interpret than a horizontal head impulse test. In such cases, the quantitative head impulse test using video oculography or the magnetic search-coil technique would aid in detecting the deficits of the vestibulo-ocular reflex (VOR). A head impulse test has been suggested a predictor of symptom recovery in a previous study, in which 80% of patients with persistent dizziness had a positive head impulse test, whereas only 10% of patients without dizziness showed a positive head impulse test.

Unilateral caloric paresis has been the diagnostic hallmark of vestibular neuritis. However, caloric test can only evaluate the function of HC only in the lower frequency range (~0.003 Hz), and would be normal in vestibular neuritis sparing HC, likewise in inferior vestibular neuritis.

Evaluation for the oculotilt reaction (OTR) should include measurements of head tilt, skew deviation, and ocular torsion. Ocular torsion can be determined on fundus photos. The sensory manifestation of ocular torsion can be measured as a deviation of the subjective visual vertical (SVV). The SVV tilt can be determined bedside using a simple, self-made bucket, or be measured using a computerized program in the laboratory.

Even though the origins of cervical and ocular VEMPs remain controversial, cervical and ocular VEMPs appear to be valuable tools in evaluating the otolithic function in vestibular neuritis. Furthermore, the dissociated patterns of abnormalities in cervical and ocular VEMPs may provide important clues for determining the involved vestibular division in vestibular neuritis. In patients with vestibular neuritis and intact cervical VEMP evoked by air-conducted sounds (ACS), the severity of ACS-induced ocular VEMP abnormality correlated with HC dysfunction on caloric or head impulse testing. In a recent study, patients with superior vestibular neuritis showed abnormal ocular VEMP and normal cervical VEMP in response to ACS. This study supports the use of ocular and cervical VEMPs in determining each subtype of vestibular neuritis.

The rotatory chair test has a limited diagnostic value in vestibular neuritis because whole-body rotation modulates the activities of both labyrinths simultaneously, which renders evaluation of unilateral vestibular dysfunction difficult. The horizontal VOR elicited by sinusoidal rotation is asymmetrical, and the VOR gain is decreased during velocity step rotation toward the affected ear, especially during the acute phase. In contrast, the VOR gain during velocity step rotation toward the contralateral ear remains unchanged.

In vestibular neuritis, direct visualization of the affected vestibular nerve has been reported with gadolinium-enhanced 3-Tesla magnetic resonance imaging (3T MRI). However, imaging the affected nerve is of more academic interest than of practical utility in vestibular neuritis. In vestibular neuritis, neuroimaging is mostly indicated when a central cause is suspected.

### Diagnosis

Because no confirmatory diagnostic tests are available, vestibular neuritis is primarily a diagnosis of exclusion. Diagnosis of vestibular neuritis is mostly based on a constellation of bedside and laboratory findings. Even though the detailed findings may differ according to the
vestibular divisions affected, the key symptoms and signs of vestibular neuritis generally include acute onset of isolated sustained vertigo with nausea/vomiting, unidirectional horizontal-torsional nystagmus beating away from the lesion side, impaired semicircular function documented by head impulse or caloric tests, ipsiversive OTR and SVV tilt, decreased or abolished VEMP responses during stimulation of the affected ear, and unsteadiness with a falling tendency to the lesion side.

The recent development of the head impulse test and VEMPs has enabled us to evaluate the function all three semicircular canals, utricle, and saccule, and to securely define the subtypes of vestibular neuritis (►Table 2).

**Superior Vestibular Neuritis**
Superior vestibular neuritis is the most common type. The spontaneous nystagmus is mostly contraversive horizontal-torsional and upbeat (►Fig. 2A). Indeed, three-dimensional recording of eye movements in vestibular neuritis showed that the rotational axes of spontaneous nystagmus clustered along the axis of HC or between the axes of HC and AC (►Fig. 2B). Patients also show a positive head impulse test for the involved AC and HC. Laboratory findings include ipsiversive ocular torsion and tilt of the SVV, ipsilesional caloric paresis, and abnormal ocular VEMP with preservation of hearing and cervical VEMP (►Table 2).

**Inferior Vestibular Neuritis**
Vestibular neuritis rarely involves the inferior division only. Diagnosis of inferior vestibular neuritis is challenging because the usual signs of vestibular neuritis are absent in this disorder. As a result, isolated inferior vestibular neuritis may erroneously be ascribed to a central pathology if there is no scrutinized evaluation for the inferior vestibular function. The spontaneous nystagmus is contraversive torsional and downbeat (►Fig. 3A). The rotation axis of the spontaneous nystagmus is best aligned with that of the involved PC (►Fig. 3B). Other findings include a positive head impulse test only for the affected PC and an abnormal cervical VEMP in response to ACS in the presence of normally functioning HC and AC, as determined by a normal head impulse test and caloric test (►Table 2). Ocular torsion, SVV, and ocular VEMP are mostly within the normal range (►Fig. 3C–G). Some patients may have tinnitus and hearing loss in the involved side.

**Total Vestibular Neuritis**
The spontaneous nystagmus is mostly torsional-horizontal beating away from the affected ear. Otherwise, the patients with total vestibular neuritis would show all the

<table>
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<th>Table 1 Clinical features of vestibular neuritis</th>
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<td>Subacute or acute onset of spontaneous vertigo with nausea/vomiting</td>
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<td>Oscillopsia: Spinning of surroundings in the direction of nystagmus quick phase</td>
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<td>Impaired function of the semicircular canals as revealed by head impulse tests or caloric testing</td>
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<td>Ipsiversive ocular tilt reaction (head tilt, skew deviation, and ocular torsion) and ipsiversive tilt of the subjective visual vertical/horizontal</td>
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| Table 2 Comparison of the findings among the subtypes of vestibular neuritis |
|----------------|-----------------|----------------|
|    | Superior | Inferior | Total |
| SN | H(C)-T(C)-U | T(C)-D | H(C)-T(C) |
| HIT-AC | Impaired | Normal | Impaired |
| HIT-HC | Impaired | Normal | Impaired |
| HIT-PC | Normal | Impaired | Impaired |
| Caloric test | Abnormal | Normal | Abnormal |
| OTR | Ipsiversive | Normal | Ipsiversive |
| SVV | Ipsiversive | Normal | Ipsiversive |
| oVEMP | Abnormal | Normal | Abnormal |
| cVEMP | Normal | Abnormal | Abnormal |

Abbreviations: AC, anterior semicircular canal; C, contraversive (quick phase); cVEMP, cervical vestibular-evoked myogenic potential; D, downbeat; H, horizontal; HC, horizontal semicircular canal; HIT, head impulse test; OTR, ocular tilt reaction; oVEMP, ocular vestibular-evoked myogenic potential; PC, posterior semicircular canal; SN, spontaneous nystagmus; SVV, subjective visual vertical; T, torsional; U, upbeat.
Fig. 2 Clinical features of superior vestibular neuritis. (A) A patient with right vestibular neuritis involving the superior division shows left beat, upbeat, and counterclockwise (from the patient’s perspective) torsional nystagmus. (B) The rotational axes of the spontaneous nystagmus cluster between those of anterior and horizontal semicircular canal. (C) Rightward ocular torsion is noted on fundus photos (normal range: 0–12.6 degrees; negative values indicate intorsion.). (D–G) The patient also shows right caloric paresis (F) and decreased amplitude of ocular vestibular-evoked myogenic potential (VEMP) during right ear stimulation with air-conducted sounds (ACS) in the presence of normal hearing (D) and cervical VEMP to ACS (F). In (A), upward deflection indicates rightward, upward, and clockwise torsional eye motion. In (B), the rotational axis of each canal was calculated from the data of Della Santina et al.93 (LH, horizontal position of the left eye; LV, vertical position of the left eye; LT, torsional position of the left eye; RHC, right horizontal semicircular canal; RAC, right anterior semicircular canal; RPC, right posterior semicircular canal semicircular canal).
Fig. 3  Clinical features of inferior vestibular neuritis. (A) A patient with right inferior vestibular neuritis shows counterclockwise (from the patient’s perspective) torsional and downbeat nystagmus. (B) The rotational axes of the spontaneous nystagmus cluster around that of posterior semicircular canal. (C–G) The patient shows no wave formation of right cervical vestibular-evoked myogenic potential (VEMP) in response to air-conducted sounds (F), while the findings of fundus photos (C), audiometry (D), bithermal caloric tests (E), and ocular VEMP induced by ACS (G) are normal. In (A), upward deflection indicates rightward, upward, and clockwise torsional eye motion. In (B), the axes in red were calculated from the data of Della Santina et al.93; the axis in blue was constructed from the data of Blanks et al.94 (LH, horizontal position of the left eye; LV, vertical position of the left eye; LT, torsional position of the left eye; RH, horizontal position of the right eye; RV, vertical position of the right eye; RT, torsional position of the right eye; RHC, right horizontal semicircular canal; RAC, right anterior semicircular canal; RPC, right posterior semicircular canal semicircular canal).
abnormalities observed in superior and inferior vestibular neuritis (Table 2).

**Course**

Severe vertigo and static vestibular imbalance markedly improve over a few days in most patients with a gradual resolution over the following weeks. Improvement of the initial symptoms occurs by virtue of central compensation for the vestibular tone imbalance rather than by recovery of the function in the affected ear. One of the central compensation signs is the reduction of spontaneous nystagmus, which usually takes 3 weeks to subside. Neural recordings in animals recovering from unilateral vestibular deafferentation show that central compensation depends on restoration of normal spontaneous activity in the ipsilesional vestibular nucleus, thereby rebalancing the activity between the left and right vestibular nuclei. Later, when the vestibular function is restored in the damaged ear, mild spontaneous nystagmus may beat in the opposite direction (recovery nystagmus). Although the static symptoms invariably resolve, albeit often not totally, the dynamic symptoms may last longer or persist. In a previous study, the static vestibular imbalances (spontaneous nystagmus, ipsilesional SVV tilt, and ocular torsion) had mostly resolved by 3 months after symptoms onset, whereas the signs of dynamic vestibular imbalances (head impulse test, head-shaking nystagmus, vibration-induced nystagmus, and caloric test) can still be observed in more than 30% of the patients at 1 year from symptom onset. Tests of the VOR function using a rotary chair yielded similar results. When tested 3 to 5 days after the onset of vestibular neuritis, the responses to low acceleration whole-body rotation in the lesion side was decreased to 50%, and in the healthy side to 75% of normal controls. Of note, the VOR responses to rotations in the healthy side were also decreased, which may be ascribed to central compensation to improve the response asymmetry. The VOR responses to the rotation in the affected side increase with time. This improvement may result from either central compensation or periphery recovery, observed as improvement in the affected ear within a few weeks, which suggests that the otoconia may be loosened due to inflammation of the labyrinth. The second most important complication is phobic postural vertigo, which entails persistent dizziness and unsteadiness associated with fear of falling without any real falls or vestibular dysfunction that can explain the symptoms.

**Differential Diagnosis**

Acute unilateral peripheral vestibulopathy may also be caused by vascular compromise of the peripheral vestibular labyrinth. Diagnosis of isolated labyrinthine infarction remains a diagnostic challenge because current imaging techniques cannot detect it. However, isolated labyrinthine infarction is exceedingly rare, and usually accompanies cochlear damage and resultant hearing loss. Rarely, isolated labyrinthine infarction may progress to involve the brainstem or cerebellar territory supplied by the anterior inferior cerebellar artery (AICA). Vestibular pseudoneuritis has also been described in infarctions involving the vestibular nucleus or inferior cerebellum. Occasionally, serial evaluation is necessary because even the diffusion-weighted MRI may fail to detect a small infarction involving the brainstem or cerebellum during the acute phase. Plaques of multiple sclerosis or lacunar infarctions involving the root entry zone of the eighth nerve may mimic vestibular neuritis. From a clinical point of view, the first question to be answered in patients with acute vertigo and nystagmus is whether the symptoms are caused by vestibular neuritis or by central vestibular pseudoneuritis. It is not always easy to differentiate isolated vascular vertigo from acute peripheral vestibulopathy at the bedside. However, a rather simple neuro-otological examination including a normal horizontal head impulse test, direction-changing nystagmus, and skew deviation (HINTS) can reliably detect central vertigo with a high sensitivity and specificity. Even these bedside tests are more sensitive for stroke than early MRIs, while maintaining a high specificity. Indeed, initial diffusion-weighted MRI may be false-negative in 12 to 20% of the stroke patients within the first 48 hours. Because a mild degree of skew deviation is hard to detect in the presence of spontaneous nystagmus, and gaze-evoked nystagmus may be absent in cerebellar stroke, a bedside head impulse test appears to be the best tool for differentiating isolated vascular vertigo from acute vestibular neuritis. However, because a bedside head impulse test may be negative in patients with covert corrective saccades and may be inconclusive in patients with nystagmus, recording of the head impulse test using a video-based equipment may be helpful in patients with equivocal results. Of course, a positive head impulse test does not necessarily eliminate the possibility of central lesions. Because recurrence is rare in vestibular neuritis, an alternative diagnosis should be considered whenever patients report more than one episode.

**Treatment**

Treatments of vestibular neuritis generally include supportive care during the acute phase, steroids, and vestibular rehabilitation.
Symptomatic care with vestibular suppressants should be applied only during the first several days when the patients suffer from severe nausea/vomiting and vertigo because these medications may delay the central compensation. Based on the shared theory of viral etiology in vestibular neuritis and Bell’s palsy, antiviral agents and steroids have been tried. However, the efficacy of corticosteroids is controversial. A recent Cochrane review concluded that there is currently insufficient evidence to support the use of corticosteroids in patients with idiopathic acute vestibular dysfunction. This review found four trials (total of 149 participants) that compared oral corticosteroids against placebo. Although there was a significant effect of corticosteroids on complete caloric recovery at 1 month, the review demonstrated no significant effect of corticosteroids on complete caloric recovery at 12 months, the extent of caloric recovery after 1 or 12 months, symptomatic recovery from vertigo at 24 hours, or the Dizziness Handicap Inventory score at 1, 3, 6, and 12 months. Further studies are also required regarding effects on subjective improvements and quality of life. The administration of valacyclovir alone or its administration in combination with glucocorticoids revealed no effect either.

A prospective study demonstrated that specific vestibular exercises significantly improve vestibulospinal compensation in patients with acute vestibular neuritis. There is moderate evidence that vestibular rehabilitation therapy is effective during the acute period of vestibular neuritis. Equilibrium training significantly lessens the time required for vestibulospinal compensation and postural regulation to develop. Voluntary eye movements and fixations are exercised to improve impaired visual fixation. Furthermore, active head movements as well as balance tasks, goal-directed movements, and walking should be encouraged to realign the vestibular reflex, and to improve vestibulospinal postural regulation and goal-directed motor function. Patients should exercise at least for 30 minutes three times a day. A recent report showed that the visual compensation device such as Nintendo Wii Balance Board (Nintendo, Inc., Redwood, WA) also aids in vestibular compensation, even with steroids.

**Conclusion**

Careful history taking and a focused neurologic examination are usually enough for diagnosis of vestibular neuritis. With the aids of a head impulse test and cervical and ocular VEMPs, each subtype of vestibular neuritis can be securely diagnosed as superior, inferior, or total vestibular neuritis. Even though very rare, inferior vestibular neuritis should be considered in patients with acute spontaneous vertigo and torsional downbeat nystagmus. Imaging should be considered whenever there is any finding inconsistent with vestibular neuritis because it is a diagnosis of exclusion. Management during the acute phase is primarily supportive while long-term treatment should be designed to improve vestibular compensation.

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