Introduction
Vestibular migraine is the frequent and rarely diagnosed caused of episodic vertigo. Therefore, its prevalence is not clearly known, but is estimated to be 4.3–29.3%.[1] It was previously named migraine-related vertigo, migraine-associated vestibulopathy and migrainous vertigo as well, and was defined as vestibular migraine by the members of Barany Society and the International Headache Society (IHS) in 2012. In 2013, the diagnostic criteria of the disease were published in the appendix of the International Classification of Headache Disorders 3 (ICHD-3) beta.[2,3]

Case Report
A 46-year-old woman had vertigo attacks of several seconds, triggered by movement for 5 months. She had increased and more frequent dizziness and in the last 2 months, which was prolonged for as long as 1 hour. Some of the attacks were accompanied by throbbing headache of half of the head with photophobia and phonophobia, which lasted for as long as 3 hours. Due to these complaints, she was previously evaluated in ear-nose-throat and neurology clinics of an external medical center, no pathology was determined in the examination and the laboratory findings, and her treatment was begun with betahistine dihydrochloride. Despite long-term therapy, she presented again to our clinic since her attacks persisted, and nausea-vomiting had accompanied her complaints for 15 days, and dizziness had become continuous for 2 days. Her history included non-aura migraine in three members of her family with an onset at the second decade. Her neurological examination included nystagmus with continuous, minimal rotational component gaze induced left direction.
Cranial MRI, MRI venography, carotid-vertebral doppler ultrasonography, electroencephalography and blood analysis findings were all normal. No hearing loss was determined in the audiogram tests performed during the attack and the non-attack periods. The patient was considered to have VM and oral administration of valproic acid 500 mg was begun. 80% of the symptoms regressed at the 2nd hour of medication, and the nystagmus disappeared. She had no attack under valproic acid therapy during her one-month follow-up, and the agent also had a positive effect on her emotional condition.

Discussion

Presence of dizziness and headache is a frequent cause of admissions to neurology clinics. VM is one of these causes. Its incidence among the general population is about 1% life-long and 0.9% annually. It is 1.5–5 fold more common among women than men. Genetic heritance is not clear; however chromosomes 11q and 5q35 have been investigated recently for migraine-related vertigo. [4,5] Despite numerous studies, the pathophysiology of VM is not clear. The main hypothesis on the subject includes the pathophysiology of migraine and the role of reciprocal connections between vestibular nuclei in the brain and structures regulating the trigeminal nociceptive inputs.[1,8]

In vestibular migraine, the onset of the symptoms is observed 5–10 years after the original onset of the migraine.[6,7] Episodic attacks characteristically include spontaneous, positional or movement-induced vertigo, dizziness, feeling of imbalance, and migrainous headache (may be aura or non-aura). Dizziness and headache may be observed at different times or at the same time. Horizontal or vertical nystagmus, gaze-evoked (single direction or both directions) nystagmus, saccadik pursuit and central positional nystagmus may be observed in the attacks.[8] In a study analyzing the contribution of nystagmus to the diagnosis of VM, the types of nystagmus observed in patients were analyzed, and it was concluded that every type of nystagmus may be observed in patients with VM, and that nystagmus may be induced positionally during the evaluation. [9] The laboratory findings of our case were normal, and the diagnosis of VM was made upon clinic symptomatology and history of migraine. Gaze-evoked nystagmus was observed on the neurological examination in both directions during her attack. Neurological examination was normal during the non-attack period. This is important for the diagnosis of VM.

Diagnosis is vestibular migraine is made on the basis of clinical symptomatology. Therefore, the attacks of the patients should briefly be questioned for the differential diagnosis. The differential diagnosis includes basilar migraine, Meniere’s disease, benign paroxysmal vertigo, transient ischemic attack. Vertigo attacks may be misjudged as migraine aura; however, aura may be distinguished by its relation to headache and its duration.[10] Vertigo may accompany basilar migraine; however, according to the ICDH-2 criteria, migrainous headache that is subsequent to a minimum of two posterior circulation findings that take 5–160 minutes may be present for basilar migraine. Furthermore, it has been reported that cases with basilar migraine have more severe vertigo attacks compared to vestibular migraine.[11] While pulsatile, progressive hearing loss with vertigo attacks, accompanied by tinnitus and low-frequency hearing loss in the early period may indicate Meniere’s disease, it is differentiated from VM this way. In addition to the similarity in the symptomatology between Meniere’s disease and VM, Meniere’s disease has been reported to be more common among patients with a history of migraine.[12] Another pathology in the differential diagnosis is that the vertigo attacks take seconds in benign paroxysmal vertigo, and spontaneously end typically within weeks or a month, whereas in VM attacks that repeat frequently within a year takes long and no spontaneous improvement is expected. During the acute attacks of vertigo, the analysis of the positional nystagmus usually permits differentiation of positional VM from benign paroxysmal vertigo.[13] TIA is considered more frequently in later ages. Ischemic attack is different to VM, with other accompanying neurological findings, risk factors, and a pathological doppler ultrasonography.[1]

There is no consensus guideline for the treatment in VM, and the recommendations are based on randomized controlled studies, case reports, retrospective cohort studies and open label trials. In studies on tryptan use in acute attacks, almotryptan 12.5 mg orally was found to be highly effective in relieving
the vertiginous symptoms, sumatryptan (variable doses) was found to be effective, and zolmitryptan 2.5 mg oral was found to be slightly effective.[14,15] While antiemetic treatments (e.g., dimenhydrinate and benzodiazepine) are believed to be effective in acute attacks, the effect of methylprednisolone (1000 mg/day, 1–3 min.) in long-term use in serious episodes has also been reported.[16] Prophylactic therapy includes acetazolamide, cinnarizine, flunarizine, lamotrigine, pizotifen, propranolol, rizatriptan, topiramate and valproic acid; drug selection may be made separately in each patient.[17,18] In the study of Çeliker et al., valproic acid was found to be effective on the vestibular symptoms of patients with migraine, whereas in another study comparing venlafaxine and flunarizine, the efficacy of valproate on the vestibular symptoms was found to be lower compared to other agents.[19,20] Our case had presented in the attack period with dizziness only, and did not benefit from intravenous dimenhydrinate therapy. However, a rapid response was observed to valproic acid 500 mg oral, which was recommended during prophylactic therapy.

In conclusion, vestibular migraine should be considered in neurological examinations, history should be obtained in detail in patients presenting with dizziness, and diagnosed cases undergo attack therapy, and prophylactic therapy should also be recommended. The selection should be patient-specific in prophylactic therapy.

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References