OBJECTIVE: To evaluate the presence of anti-compensatory saccades (AcS) using the video head impulse test (vHIT) in the healthy inner ear in patients with vestibular neuritis (VN) during the acute and subacute stages of VN.

METHODS: We retrospectively considered a chart review of 2420 patients evaluated for acute vestibular syndrome from 2016 to 2020 in the Cassino (Italy) clinic. Nine hundred fifty-four patients with acute onset of vestibular syndrome who received an instrumental otoneurological assessment within 24 hours from the onset of the symptoms, evaluated by simultaneously using a combination of vHIT, ocular vestibular-evoked myogenic potential (VEMP), and cervical VEMP, were included in the study.

RESULTS: Thirty-two patients with superior VN that showed an altered horizontal canal function when tested with vHIT and quick AcS on the healthy side were enrolled. We found that all patients with VN, evaluated in the first 24 hours from the onset of the symptoms, showed AcS when their head was abruptly and passively turned toward the healthy inner ear side. At follow-up within 8 weeks from the first evaluation, 29 out of 32 patients did not show increased AcS.

CONCLUSIONS: Our findings support the hypothesis that the AcS on the healthy side are a clinical sign of vestibular canal hypofunction or paresis in patients affected by the acute stage of VN.

KEYWORDS: Vestibular, vestibular neuritis, anti-compensatory saccades, head impulse test, video head impulse test

INTRODUCTION

The video head impulse test (vHIT) constitutes an important clinical tool to evaluate horizontal and vertical vestibulo-ocular reflex (VOR).1-3

Individuals undergoing this test are instructed to maintain their eyes on a target placed in front of them, in solidarity with the earth, during short, passive, unpredictable, horizontal head turns. This is now called the head impulse test paradigm (HIMP).

The decreased canal function does not allow the patient to maintain fixation on the target during the vHIT, and to respond in the best way to the fixation task, the patient needs a series of corrective eye saccadic movements during or at the end of the head rotation. The first 100 ms of the eye movement does not depend on other sources of vision control, and the earliest part of the eye movement responds only to a vestibular stimulus. The head impulse test or HIMP assesses the functional state of each semicircular canal.

Over the past few years, a new test paradigm has been proposed to evaluate the vestibular function.2,10 During abrupt, unpredictable head turns, the subjects are asked to look at a target that moves with their head rather than the fixed target used in HIMP. This new paradigm is called SHIMP, suppression head impulses. In fact, at the end of each head turn, a corrective
stage (the so-called “SHIMPs” saccade) appears in normal subjects, whereas it does not appear in patients in the acute stage of peripheral diseases such as neuritis, or iatrogenic vestibular diseases such as intratympanic gentamicin or neurectomy.7 The SHIMPs paradigm is not affected by covert saccades and could give more information on slow-phase velocity of the VOR, compared to HIMPs. Recently we have demonstrated that the VOR gain value of the affected side in patients with vestibular neuritis (VN), might be different at the time of the attack (within 72 hours) rather than in the subacute or chronic stage (from 72 hours to 6 weeks after the onset of symptoms), as well as the related impact on patients’ quality of life.11 Furthermore, we found that when the operator turns the patient’s head toward the unaffected side, a series of quick anti-compensatory saccades (AcS) is generated. This group of eye movements, at a first glance, seems related to the quick phase of spontaneous nystagmus. The latter could be reduced at the time of the attack of an acute vestibular syndrome but not completely suppressed by fixation. To confirm this, it should be emphasized how the vHIT test is typically performed in a well-lit room.12

Whereas the study of VOR gain through the vHIT is entirely focused on the affected side, our clinical hypothesis is that in the very early stage of the vestibular neuritis, the healthy side can also yield important clinical signs to guide the physician in the diagnosis of acute VN. Therefore, this study aims to retrospectively investigate the presence of AcS in the healthy side in patients with VN during the acute and subacute phases.

MATERIALS AND METHODS

Study Design
The aim of this study was to explore the presence of anti-compensatory saccades (AcS) in the healthy side during the acute and subacute phase in VN patients. Ethical standards and guidelines on human experimentation were observed during all procedures of this work. Guidelines for reporting observational studies (Strengthening the Reporting of Observational Studies in Epidemiology) were followed. All participants gave their written informed consent for participation in the study. This study was approved by the IRB of the ENT MSA Academy Center.

Setting
Medical records of patients who were admitted to the ENT medical Center from 2016 to 2020 with a diagnosis of VN evaluated within 24 hours from the onset (T0) and after 8 weeks (T1) were included.

Participants
The inclusion criteria were diagnosis of VN evaluated within 24 hours from the onset of the symptoms), and patients who had not received therapy for VN. We excluded patients with a diagnosis of Meniere’s disease, benign paroxysmal positional vertigo bilateral, vestibular migraine, the presence of somatic or psychiatric disorders, and neurological diseases. The vestibular evaluation included the vHIT, a self-assessment dizziness handicap inventory (DHI), bone-conducted vibration and air-conducted sound cervical and ocular vestibular-evoked myogenic potentials (VEMPs). The VN patients were diagnosed having acute vestibular syndrome (AVS) characterized by rotatory vertigo, postural imbalance, and neurovegetative symptoms such as nausea and vomiting; abnormal results of the horizontal semicircular canal function; an asymmetry ratio (AR) greater than 40% of n10 to oVEMPs, an AR greater than 30% of p13-n23 to cVEMPs, absence of neurological signs, and the presence of HINTS peripheral pattern or the presence of horizontal spontaneous nystagmus without evidence of a central vestibular lesion; and absence of cerebral lesions at MRI of the brain evaluation performed 72 hours after the AVS onset.9

Dizziness Handicap Inventory
The quality of life was assessed by the self-assessment inventory DHI. With 25 questions, the DHI evaluates self-perceived activity limitation and restriction resulting from dizziness.10

Video Head Impulse Test
Vestibulo-ocular reflex was evaluated with a video system (OtosuiteV®, GN Otometrics, Denmark)

The functional canal state was assessed by measuring the corrective eye movement during an unpredictable head movement using vHIT.7 The patients were instructed to keep staring at an “earth-fixed” fixation target on the wall in front of them during the head turn and to return their eyes to that target as quickly as possible if they lose the target. To ensure that the pupil image was not affected by reflections, room lightening conditions were adapted. Twenty brief, quick, unpredictable horizontal head turns to each side were applied by the clinician. The head impulse for each session was “turn and stop.”

For each head turn, the eye and head velocity were recorded. The affected side was identified by VOR gain < 0.76 with 100% sensitivity and specificity. The VOR gain was represented by the ratio of the area under the curve (AUC) (0.81-1.0, P < .0001).6 AcS was defined as saccades in the direction of the head movement with a peak velocity of 50°/s. AcS start was at 10°/s, the difference between head impulse and AcS start was latency. The rate of occurrence of AcS was the proportion of impulses with AcS.11

Vestibular-Evoked Myogenic Potentials
The oVEMP n10 is a small (5-10 µV) negative (excitatory), crossed, VEMP of the stretched inferior oblique eye muscles13,14 recorded by surface EMG electrodes on the skin beneath the eyes, in response to stimulation by bone-conducted vibration (BCV) delivered to the midline of the forehead at the hairline (Fz) and to air-conducted stimuli (ACS), 500 Hz short-tone burst. Based on evidence of utriculo-ocular projections15 and neural evidence of the preferential activation by BCV and ACS of irregular otolithic afferent neurons,16-21 n10 to oVEMP mainly reflects the utricular macula activity for these stimuli.

In contrast, BCV and ACS p13-n23 to cVEMP is a positive (inhibitory) uncrossed potential recorded by surface EMG electrodes over the tensed sternocleidomastoid muscle and reflects mainly the saccular macula function.6,13,22

Statistical Analysis
Data were analyzed using the Statistical Package for the Social Sciences version 23 (IBM Corp., Armonk, NY, USA). The independent
samples t-test was used to analyze the improvement in VOR gain, and the Mann–Whitney U-test was applied to compare DHI data within subjects. P-values < .05 were considered significant.

RESULTS
Thirty-two VN patients (13 females and 29 males), mean age 59.19 ± 16.23, met the inclusion criteria and were included in the study (participants’ demographic characteristics are reported in Table 1). All patients showed an altered ipsilesional VOR gain at T0 (0.40 ± 0.16) and a physiological contralesional VOR gain (0.91 ± 0.10). The enrolled patients had an AR greater than 40% to oVEMP in response to BCV and ACS stimulus, with the n10 component being markedly reduced beneath the eye opposite the affected side. In contrast, the ACS and BCV AR p13-n23 to cVEMP was symmetrical. Significant differences were found in the within-subjects analysis at T0 and T1 in DHI scores (P = .000). Affected hVOR gain values showed significant differences at T1 with respect to T0 (P = .02), whereas no differences were found in the contralesional VOR gain values (P = .11).

All patients showed AcS on the healthy side at T0. Twenty-nine out of 32 patients evaluated at T1 did not show increased AcS.

DISCUSSION
This study aimed to explore the presence of AcS in the healthy side in patients with VN during the acute and subacute phases. We found that in all patients evaluated during the acute phase, AcS were present during the rotation toward the healthy side. The presence of AcS can be related to the VOR destabilization process that occurs during the acute stages of VN. In particular, this can also be emphasized by passive head rotations toward the healthy side at the time of the attack and seems to disappear in all T1 patients (Figure 1A-C). Essentially, AcS can be regarded as a sign of altered VOR function, similar to what happens on the affected side or the desired realization of the reflex, “eyes on target,” at the end of the head turning. These rapid eye movements always temporally follow the movements of the head, the anti-compensatory saccades, but unlike the rotation toward the affected side, they have the same direction. Other authors have speculated that these quick eye movements are an indicator of peripheral injury.

For this reason, considering our large sample of subjects evaluated in the first 24 hours—practically in the time that we like to define as “the time of the attack”—it is possible to relate this phenomenon to the acute stage of the peripheral nerve injury or a sign of the very early stages of the pathophysiological insult that underlies VN. However, the reason why AcS are present only in “early” vHIT evaluation is still under debate, and our aim is certainly to shed new light on the matter.

Indeed, related features of kinematics share both rapid phases and saccades, which are rapid eye movements. Furthermore, the genesis of horizontal eye movements recognizes 2 distinct neural circuits. The first of these circuits is undoubtedly generated by the trineuronal arch for the VOR slow phase.

The second is a separate neural circuit which is located in the brainstem, responsible for the genesis of the quick phase represented by a double network: (a) burst neurons and (b) pause neurons. The latter circuit, activated by vestibular inputs, generates both vestibular nystagmus rapid phase during horizontal angular acceleration and voluntary saccades triggered by descending axons from the superior colliculus. This is a first possibility for the genesis of saccades, the other is represented by contribution or trigger from neck afferents. There is also evidence that this rapid process of the neural circuit may also be triggered by these neck inputs.

In our sample, we found a very low VOR gain within 24 hours from the onset of the symptoms, suggesting that the acute insult to the superior component of the vestibular nerve had disabled the horizontal slow-phase mechanism (Fig. 1B).

In any case, during the vHIT toward the unaffected side, similar to what happens when the head is turned toward the affected side, the quick phase mechanism is not compromised, and right in the first hours of illness, AcS are generated similar to the quick phase of the spontaneous nystagmus, directed away from the affected side. Essentially, these quick eye movements seem to be due to the presence of spontaneous nystagmus that does not seem to be fully suppressed by fixation. Indeed, the vHIT test is usually conducted in a normally lit room. We recognize some limitations in the present study.

First, there are limitations to the interpretation of our findings, because this was a retrospective study. Second, the timing of follow-up is heterogeneous, and future studies could better identify the timing of re-evaluation to verify how long-lasting this clinical sign is, to better understand the early stage of the vestibular compensation process.

CONCLUSION
Our findings suggest that the presence of anti-compensatory overt saccades, measured using the vHIT while turning the head to the
Healthy side in a patient affected by VN, is a new instrumental clinical sign of acute VOR impairment, and a marker for recognizing the test execution timing.

**Ethics Committee Approval:** This study was approved by IRB of MSA ENT Academy Center.

**Informed Consent:** All patients gave their written consent.

**Peer Review:** Externally peer-reviewed.

**Author Contributions:** Concept – L.M.; Design - M.T., L.M.; Data Collection and/or Processing - L.M., M.T.; Analysis and/or Interpretation - L.M., M.T.; Writing - M.T., L.M.

**Conflict of Interest:** The authors have no conflict of interest to declare.

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