Absence of nystagmus during REM sleep in patients with vestibular neuritis

I Eisensehr, S Noachtar, M Strupp, H v Lindeiner, T Brandt, U Büttner

Abstract
Saccades, including fast phases of nystagmus, disappear during drowsiness and non-rapid eye movement (NREM) sleep, but are present during the alert state and REM sleep. The purpose of this study was to determine whether spontaneous nystagmus is present in patients with vestibular neuritis during REM sleep.

Eight patients with spontaneous nystagmus due to vestibular neuritis and eight control patients without any nystagmus underwent at least one night of polysomnography. Fast phases of nystagmus were analyzed. The number of right and left horizontal saccades were counted, first during 3–5 minute samples of the awake state before sleep onset, then during the first REM episode and the last REM episode of nocturnal sleep, and finally during the alert state in the morning after nocturnal sleep.

All patients with vestibular neuritis showed significantly more saccades (fast phases) towards the side contralateral to their vestibular lesion in the awake state before and after the polysomnography. This reflects their spontaneous nystagmus. By contrast, during REM sleep the patients with vestibular neuritis showed no preponderance in saccade direction. The eye movement pattern in REM was the same for patients and controls.

In conclusion, peripheral vestibular imbalance producing nystagmus in vestibular neuritis in the awake state is not active at the brain stem level during REM sleep.

Keywords: vestibular neuritis; spontaneous nystagmus; REM sleep; saccades

It is assumed that the same pontine neurons burst with saccades during the alert state, REM sleep, and the fast phases of nystagmus, which are saccades. There is a close relation between the presence of saccades and nystagmus. As eye movements during REM sleep are considered to be saccades, nystagmus could also be expected to occur during REM sleep.

This is the first prospective study on peripheral vestibular lesion and REM sleep which investigated whether spontaneous nystagmus during the awake state in patients with acute vestibular neuritis also occurs during REM sleep.

Methods
STUDY POPULATION
Informed consent was obtained from all participants in the study. Eight patients with vestibular neuritis and eight controls had a complete physical examination and underwent at least one night of polysomnography (PSG). Polysomnography of the patients with vestibular neuritis was performed from days 1 to 6 after the onset of symptoms of vestibular neuritis (mean 69 hours, range 24–144 hours). For inclusion in the study, nystagmus (slow phase velocity of at least 7°/s in the dark) before and after the PSG night had to be present in patients with vestibular neuritis. The diagnosis of vestibular neuritis was based on criteria described in detail elsewhere.

The patients without vestibular neuritis had been referred to the sleep laboratory because of suspected sleep disorders. They were included in the study as controls if they had no history of vertigo and no spontaneous nystagmus during the awake state as verified with Frenzl’s glasses.

SLEEP STUDIES
The recording and scoring of the PSG were performed as described elsewhere. An electro-oculogram (EOG) was obtained from electrodes placed lateral to both eyes and to the infraorbital and supraorbital regions; EOG calibration was performed at the beginning and the end of the PSG and included eye excursion of 35° from the midline to the right and left, vertical views with and without Frenzl’s glasses on, and provocation of vertical and horizontal optokinetic nystagmus. A time constant of 10 seconds was chosen for the EOG. Velocities of the slow phases of horizontal eye movements can be calculated by summarising the amplitudes of the fast phases during a certain time period. Slow phase velocities were exclusively determined for patients with vestibular neuritis during the awake state, when long periods of obvious nystagmus in one direction were present. Right and left (ipsilateral and contralateral) horizontal saccades were counted separately during 3–5 minute samples of the awake state before sleep onset, the awake state after the nocturnal sleep period, and the first and the
last continuous REM episodes of the nocturnal sleep. Horizontal eye movements were scored only if they had an excursion of 2° or more and a velocity of at least 50°/second. Any preponderance in saccade direction was considered potential nystagmus.

DATA PRESENTATION AND STATISTICS
Data were analyzed using the SPSS statistical package for Windows 9.0. They are presented as mean (SD) if not stated otherwise. The data were tested for normal distribution with the Kolmogorow-Smirnov test. Data points between and within the two study groups were compared using the χ2 test and Mann-Whitney U test for paired and independent samples. In the two groups (controls/vestibular neuritis), the numbers of right/ipsilateral and left/contralateral horizontal saccades were compared within awake and REM sections to detect directional preponderance of horizontal saccades and nystagmus. A p value<0.05 was considered significant.

Results
Patients with vestibular neuritis were slightly older than controls (vestibular neuritis 65.1 (14.7) years, controls 55.4 (19.8) years) and were more often women (n=7) than controls (n=4), (p>0.05). Vestibular neuritis was located on the right in five patients and on the left in three patients. Controls were diagnosed as having periodic limb movement disorder (n=2), narcolepsy (n=2), obstructive sleep apnoea (n=1), and sleep state misperception (n=3). These disorders are not reported to be associated with ocular-motor disorders. As we wanted to score rapid eye movement directions and were not interested in analyzing sleep macrostructure, we thought that our controls were adequate. All patients with vestibular neuritis had spontaneous nystagmus, which was best seen with Frenzl’s glasses on, and to a lesser extent during fixation before and after the night of recording. The slow phase velocity with the eyes closed in patients with vestibular neuritis had decreased slightly from the evening before to the morning after the PSG (evening 11.05 (6.80)°/s; morning 8.39 (1.32)°/s; p<0.05).

The videotapes showed that all patients and controls lay on their back without turning their heads extensively to the right or left side during the analyzed awake and REM periods. All patients with vestibular neuritis showed significantly more saccades to the side contralateral (ipsilateral) to their vestibular lesion in the awake state (eyes open and closed), indicating spontaneous nystagmus (p=0.012) (table 1).

During REM sleep, none of the patients with vestibular neuritis showed either any nystagmus or any significant preponderance in saccade direction. Similarly, no tonic deviation was found in patients with vestibular neuritis during REM sleep (analyzing eye in head position with EOG and synchronised videotape). No difference in saccade direction was found in control patients during the awake state or REM sleep. Occasionally, brief phases (about 2 seconds long) of nystagmoid jerks (one to four jerks a second) or slow horizontal eye movements with velocities between 10 to 20°/s (fig 1A) with changing directions and without any directional preponderance were found in both patients and controls during REM sleep. Saccadic eye movements were not found in non-REM (NREM) sleep.

Discussion
This is the first prospective study comparing eye movements in REM sleep of well defined patients with vestibular neuritis with those of controls. We found that the mechanisms generating REMs during desynchronised sleep are not influenced by mechanisms producing nystagmus in vestibular neuritis. Moreover, the nystagmus associated with vestibular neuritis was abolished during NREM sleep. Studies investigating eye movements during REM sleep of patients with nystagmus in the awake state due to various CNS lesions report contradictory findings.10–13 Gordon et al,10 who examined six patients in a vegetative state, and Appenzeller et al,11 who studied three patients with alcoholic and one patient with post-traumatic encephalopathy, described brief phases of nystagmoid jerks during REM sleep. These jerks differed from the continuous nystagmus seen during wakefulness (no description of nystagmus direction). However, brief phases of irregular nystagmoid jerks during REM sleep have also been described in healthy subjects,13 and we also found non-systematic brief phases of nystagmoid jerks in vestibular neuritis and controls. These may be related to a gaze holding deficit on lateral gaze, which might correspond to a neuronal integrator disorder during REM sleep. By contrast, two studies reported that the nystagmus generated by CNS lesions in the awake state disappears during NREM and REM sleep.12 13 Another study found a nearly total suppression of rapid eye movements directed away from the side of the lesion during REM sleep in six patients with unilateral attentional neglect,14 although phases of nystagmus were not described in the study.

In cats, unilateral labyrinthectomy and unilateral section of the eighth nerve resulted in spontaneous nystagmus during wakefulness for

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Table 1 Saccades during wakefulness and REM sleep of patients with vestibular neuritis and control patients

<table>
<thead>
<tr>
<th></th>
<th>Patients with vestibular neuritis</th>
<th>Control patients</th>
<th>p Value Mann-Whitney U test</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Initial wakefulness</strong></td>
<td>Contralateral/left: 1.1 (0.54)</td>
<td>0.42 (0.19)</td>
<td>0.012</td>
</tr>
<tr>
<td></td>
<td>Ipsilateral/right: 0.2 (0.21)</td>
<td>0.38 (0.18)</td>
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<tr>
<td><strong>Last wakefulness</strong></td>
<td>Contralateral/left: 0.79 (0.34)</td>
<td>0.38 (0.23)</td>
<td>0.012</td>
</tr>
<tr>
<td></td>
<td>Ipsilateral/right: 0.16 (0.10)</td>
<td>0.38 (0.24)</td>
<td></td>
</tr>
<tr>
<td><strong>Initial REM</strong></td>
<td>Contralateral/left: 0.21 (0.24)</td>
<td>0.13 (0.10)</td>
<td>0.236</td>
</tr>
<tr>
<td></td>
<td>Ipsilateral/right: 0.23 (0.21)</td>
<td>0.13 (0.11)</td>
<td></td>
</tr>
<tr>
<td><strong>Last REM</strong></td>
<td>Contralateral/left: 0.17 (0.15)</td>
<td>0.14 (0.07)</td>
<td>0.123</td>
</tr>
<tr>
<td></td>
<td>Ipsilateral/right: 0.23 (0.17)</td>
<td>0.13 (0.05)</td>
<td></td>
</tr>
</tbody>
</table>

Contralateral and ipsilateral refers to the side of the vestibular lesion in patients with vestibular neuritis; right and left refers to control patients. Values are given as mean (SD). During REM sleep, none of the patients with vestibular neuritis showed any significant preponderance in saccade direction (Mann-Whitney U test).
several days, which was largely depressed or completely abolished during NREM and REM sleep.

Studies on the influence of vestibular stimulation on eye movements during sleep in normal subjects have given contradictory results. Reding and Fernandez, using perrotatory stimulation at 10 minute intervals in a sample of five children, found that nystagmic jerks (direction not specified) were significantly associated with the occurrence of rapid eye movements in REM sleep. However, brief phases of nystagmic jerks during REM sleep have also been described in healthy subjects under normal conditions. Other studies in adult humans demonstrated that neither postrotatory nor perrotatory vestibular stimulation elicited nystagmus during any stage of sleep. However, slow compensatory conjugate deviation of the eyes occurred with each oscillation during REM sleep, whereas rapid eye movements were absent in the study of Tauber et al; this suggests some vestibular influence on eye movements during REM sleep. We exclusively evaluated the number and direction of horizontal saccades during REM sleep. Therefore we cannot exclude vestibular influence on other characteristics of rapid eye movements during REM sleep (number of clusters versus single rapid eye movements, velocities etc). However, nystagmus seen during wakefulness in our patients with vestibular neuritis could definitely not be seen during REM sleep. In analogy to the de-erefferentation of the motor output during REM sleep, it might be hypothesised that there is a deerefferentation of the peripheral vestibular input to the REM sleep generating areas. This deerefferentation might originate from REM sleep generating structures in the paramedian pontine reticular formation and inhibit peripheral input to the medial vestibular nucleus, which plays an executive part in rapid eye movements during REM sleep.

Figure 1 (A) Thirty second sample of the awake state before sleep onset of the polysomnogram for a patient with right vestibular neuritis and one control patient. The control patient shows several blinking artifacts in the channel for the vertical eye movements. To get highly artefact free awake phases the patients and controls were told to relax as much as possible during the initial and final wake phases. This might have been easier for controls than for patients with vestibular neuritis, who mostly complained about nausea when they were awake. Therefore, the low chin muscle tone in the control indicates good relaxation by contrast with the patients with vestibular neuritis. (B) Thirty second sample of the first REM period of the polysomnogram for a patient with right vestibular neuritis and one control patient. The drift of slow eye movements to the right, which preceded a series of rapid eye movements in vestibular neuritis, was not a frequent finding. Such an occasional drift occurred in various directions and was also seen in controls. A1=left ear; A2=right ear; LBE=left below eye; RAE=right above eye; ECG=electrocardiogram.

C4–A1  C3–A2  O2–Cz  O1–Cz  T4–Cz  T3–Cz  LBE–A1  RAE–A2  EMG chin  ECG

Vertical eye movement  Horizontal eye movement

5 s

Right vestibular neuritis

Control

0% a

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