RAPID EYE MOVEMENTS DURING PARADOXICAL SLEEP IN PATIENTS WITH CEREBROVASCULAR DISEASE

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ABSTRACT - Rapid eye movements that occur during paradoxical sleep are generated from the brainstem and are modulated by cerebral hemispheres. In an attempt to establish the participation of cerebral hemispheres on rapid eye movements, we carried out a quantitative study of eye movements density in patients bearing hemispheres vascular lesions. The polysomnographic recordings of 24 patients were compared to those of 24 healthy volunteers. Density of rapid eye movements was defined as the percentage of eye movements during the respective time of paradoxical sleep. Based on the present results, we concluded that: stroke patients with hemispheric lesions displayed increased density of rapid eye movements; there was no difference on the density of rapid eye movements according to the hemispheric lesion; higher density of rapid eye movements was observed in patients with anterior hemispheric lesion.

KEY WORDS: rapid eye movements, paradoxical sleep, dreams, stroke.

Paradoxical sleep events may be classified as tonic and phasic. Tonic events are continuous and may be characterized by hippocampal theta rhythm, synchronization of cerebral electrical activity and muscle atonia. Phasic events are periodic and are characterized by rapid eye movements, bursts of sawtooth waves and by ponto-geniculo-occipital waves. Both, tonic and phasic events, result from the activation of gigantic cells in the pons. Eye movements occur as a consequence of phasic activation of oculomotor and vestibular system via pontine reticular formation.

Although the role of brainstem on rapid eye movements is beyond doubt, some authors suggest a hemispheric participation on different phases of sleep and, therefore, on paradoxical sleep, dreams and rapid eye movements. Rapid eye movements may vary as a function of genetic, constitutional
and personality factors, and, as consequence, they are considered an individual characteristic. Several studies analyzed the variation of rapid eye movements density in mood disorders. Gould et al. established the association between rapid eye movements and sleep apnea.

With regard to the relationship between sleep and cerebrovascular disease, damage of the brainstem modifies sleep architecture, mainly the organization of paradoxical sleep. Few studies investigating the relationship between encephalic vascular lesions and rapid eye movements. Laffont et al. observed a reduction of ocular movements when vascular lesions are situated in the brainstem, more specifically in the medial pontine tegmentum (gigantocellular and center-caudal pontine nuclei). Doricchi et al. examined rapid eye movements in patients bearing hemispheric vascular lesions, who presented the unilateral negligence syndrome, in order to establish the participation of attention mechanisms on rapid eye movements. Except for the lack of rapid eye movements towards the neglected side, all other sleep parameters were similar, regardless of whether the lesion was localized in the right or left hemisphere.

In order to examine the participation of brain hemispheres on rapid eye movements during paradoxical sleep, we performed a quantitative analysis of this phasic event in patients with brain lesions of vascular nature.

**METHOD**

**Subjects**

The experimental group consisted of 24 patients, whose age varied from 32 to 69 years. These patients came from the Department of Neurology of UNIFESP, with a history of stroke, that occurred more than 30 days earlier, since we were interested in well-established vascular lesions and considering that in the acute phase brain alterations are more diffuse. All patients were submitted to a cranial computerized tomography, in order to establish the localization of the lesion. Patients who presented other neurologic diseases, as well as those who used any substance that could interfere with sleep architecture, were also excluded (Table 1).

The control group was composed by 24 healthy volunteers, whose age varied from 32 to 69 years. They did not present any sleep alterations, nor did they bear any kind of neurologic disease or used any medicine that could alter the normal sleep pattern (Table 1).

**Polysomnographic recordings**

Polysomnograms were performed throughout one night, with an 8-channel Grass electroencephalograph. The recordings included brain electrical activity, eye movements, chin muscle activity, nasal and oral airflow, thoracic and abdominal respiratory movements, heart beat and leg movements. The electroencephalogram used four electrodes: central left (C3), central right (C4), left ear (A1) and right ear (A2), and placement C3-A2 and C4-A1, according to the international 10-20 system. Other electrodes placement and sleep stages classification followed the criteria established by Rechtschaffen and Kales. The following parameters were considered: total recording time, total sleeping time, duration of non-paradoxical sleep and its four stages, duration of paradoxical sleep, total time of awakening, sleep latency, paradoxical latency, REM density, apnea/hipopnea index and heart beat.

REM density is defined as the percentage of time (in minutes) in which rapid eye movements occurred during the respective period of paradoxical sleep, considering that amplitude of eye movements is higher than 25 µV. Calculation of this index was made by the following equation: total duration of rapid eye movements/total duration of paradoxical sleep x 100.

**Statistical analysis**

For statistical purposes, two groups were initially considered: control and cerebrovascular disease. After these groups were compared, cerebrovascular diseases patients were divided in bearers of left hemisphere lesions and right hemisphere lesion. For each hemisphere we contemplated lesion situated on the anterior or posterior areas. One-way ANOVA and Student's-t-test were employed in the analyses of total
recording time, total sleep time, sleep latency, awakenings during sleep, total duration of non-paradoxical sleep and its respective sleep stages, total duration of paradoxical sleep, paradoxical sleep latency. Analyses of sleep efficiency, REM density and apnea/hipopnea index were made by the Kruskal-Wallis Rank test and by Mann-Whitney U test. Correlations between density of rapid eye movements and the patient’s age, time elapsed since the stroke occurred, and sleep efficiency were made by the Sperian correlation test. The level of significance was set at p< 0.05.

RESULTS

Four healthy volunteers presented sleep apnea/hipopnea, the indices of which varied from 8.2 to 11.8/hour of sleep. In the cerebrovascular group, six patients presented respiratory alterations with indices varying from 10.1 to 59.4/hour of sleep. Statistical analysis of these parameters showed that mean index of sleep apnea/hipopnea was higher in cerebrovascular patients than in healthy volunteers (27.9±20.2/hour and 9.5±1.7/hour respectively). Cerebrovascular disease patients with sleep apnea and hypopnea displayed shorter latency to sleep than those patients who did not present respiratory alterations.

With regard to localization of lesions, the tomography revealed their presence in the left hemisphere in eight patients, four presented lesion in anterior areas, whereas in four patients the lesions were located in posterior regions. Lesions in the right hemisphere were detected in 11 patients: in four patients, these lesions were located in anterior areas, and in seven, they were situated in posterior areas. Imaging evaluation detected lesions in both hemispheres in three patients, whereas in two, there was no evidence of brain lesions.

There were no significant correlations between REM density and the patient’s or volunteer’s age, and with time elapsed from the stroke, nor could we detected a correlation between REM density and sleep efficiency in either group. There was an increase in stage 1 and a reduction of stages 2 and 3 in cerebrovascular disease patients, compared to controls. Overall, however, no changes in non-paradoxical sleep and paradoxical sleep duration were detected.

Cerebrovascular patients presented increased REM density, compared to volunteers from control group. This augmented density was observed in total, as in the first paradoxical period (Table 2). No differences were detected when volunteers were compared with patients with left or right hemisphere lesion (Table 3), however, there was a significant increase of REM density in the group with anterior lesions (Table 4).

DISCUSSION

Respiratory disorders in patients with cerebrovascular disease

The mean index of sleep apnea and hypopnea was higher in cerebrovascular patients than in healthy volunteers. Respiratory alterations during sleep, characterized by an index of apnea and hypopnea above 10/hour, were observed in six patients. In three cases in which we could determine the type of respiratory disorder, one patient presented a high index of obstructive apnea (50.9/hour).

The relationship between sleep apnea syndrome and cardiovascular alterations is well established. Askenasy and Goldhammer described the case of a patient who developed sleep apnea after having a
stroke with an ischemic infarction of the lateral medulla. In the mammalian medulla there are three clusters of respiratory neurons. For a normal respiration it is vital that these nuclei modulate the vestibulocerebellar regulation of the supraglotic, genioglossus, pharyngolarynx, diaphragm and intercostal muscle tonus. The proximity between hypnogenic and respiratory somatic centers may provide an explanation for the development of sleep apnea after an infarction of the lateral region of the medulla.

Palomaki et al. studied 177 male patients with cerebral infarction and observed that the frequency of vascular conditions is higher in snorers than non-snorers. The authors suggest that ischemic cerebral infarction occurs more frequently during sleep, at the end of the night.

Sleep architecture of patients with cerebrovascular disease

Korner and coworkers studied the sleep pattern of 19 patients with cerebrovascular disease due to thromboembolism of the medial cerebral artery diagnosed by cranial computerized tomography, 14 days after the ischemic accident. Nine of these patients presented lesion in the right hemisphere and ten in the left hemisphere. Compared to a control group, composed by 13 healthy volunteers, these patients spent more time in bed, presented lower sleep efficiency, increased time of non-paradoxical sleep, as a result of increased time of stages 1 and 4, and reduction of paradoxical sleep. Our results showed that cerebrovascular patients present an increased time of stage 1, however, without differences in paradoxical sleep.

Another study with 18 cerebrovascular patients, whose ischemia occurred at the level of the medial cerebral artery and five hours before the onset of symptoms, reported reduction of paradoxical sleep, proportional to the severity of the neurological features and depth of the lesion. On the third

Table 3. Density of rapid eye movements and localization of hemispheric lesion. Data are expressed as median and semi-interquartil interval.

<table>
<thead>
<tr>
<th></th>
<th>Total paradoxical sleep</th>
<th>Paradoxical sleep (first period)</th>
<th>Paradoxical sleep (second period)</th>
<th>Paradoxical sleep (third period)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Control (n=24)</td>
<td>4.5 (3.1)</td>
<td>4.8 (2.4)</td>
<td>6.7 (3.8)</td>
<td>5.8 (4.6)</td>
</tr>
<tr>
<td>LBH (n=7)</td>
<td>18.3 (12.3)</td>
<td>5.3 (14.7)</td>
<td>8.6 (7.5)</td>
<td>19.9 (13.8)</td>
</tr>
<tr>
<td>RBH (n=8)</td>
<td>8.0 (9.4)</td>
<td>10.8 (9.8)</td>
<td>26.3 (9.7)</td>
<td>14.5 (10.5)</td>
</tr>
</tbody>
</table>

LBH, left brain hemisphere; RBH, Right brain hemisphere.

Table 4. Density of rapid eye movements and localization of the lesion. Data are presented as median and semi-interquartil interval.

<table>
<thead>
<tr>
<th></th>
<th>Total paradoxical sleep</th>
<th>Paradoxical sleep (first period)</th>
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<td>6.7 (3.8)</td>
<td>5.8 (4.6)</td>
</tr>
<tr>
<td>Anterior (n=5)</td>
<td>19.5 (4.8)*</td>
<td>27.8 (12.4)*</td>
<td>24.2 (6.5)*</td>
<td>20.8 (15.8)</td>
</tr>
<tr>
<td>Posterior (n=10)</td>
<td>5.9 (7.8)</td>
<td>6.2 (7.0)</td>
<td>8.5 (15.6)</td>
<td>17.1 (11.6)</td>
</tr>
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*significantly different from control group; p<0.05, Kruskal-Wallis rank test, followed Mann-Whitney U test.
week following the ischemia, the authors observed normalization of paradoxical sleep. Three of our patients did not display paradoxical sleep and their lesions were located in deep regions of the brain hemisphere. It has been suggested that during the acute phase that follows ischemic cerebrovascular disease, the disappearance of paradoxical sleep may be a consequence of the suppression of eye movements and sawtooth waves, and persistency of the muscle tonus indicating that paradoxical sleep generator is still active.

**Rapid eye movements in patients with cerebrovascular disease**

The cerebrovascular patients presented augmented REM density, when we considered both total paradoxical sleep and the first paradoxical period. According King et al., patients with neurologic disorders, other than vascular disease do not present altered REM density, compared to normal controls, however, augmented REM density is reported to occur in depressive patients.

An inverse relationship between REM density and sleep depth has been suggested. Feinberg et al. observed a reduction of REM density in ten healthy volunteers after one night of sleep deprivation, and interpreted this result as a consequence of increased sleep depth. If this were true, augmented REM density in cerebrovascular patients should be due to a more superficial sleep. We did not, however, find any correlation between REM density and sleep efficiency.

Considering the localization of brain lesion, our results showed that REM density was augmented in patients whose brain lesions were located in anterior regions, compared to control subjects. This increased REM density may be due to cortical lesions that could, in turn, induce eye movements, generated in the brainstem. In cats, frontal cortex lesions suppress isolated eye movements, originated in the cortex, however, eye movements that occur in burst, generated in the brainstem, are augmented. On the contrary, occipital lesions reduce the bursts of eye movements. Jeannerod et al. suggested a facilitation role for the occipital lobe and an inhibitory role for the frontal lobe on rapid eye movements. Appenzeller and Fischer also proposed a facilitation role for the visual cortex and an inhibitory influence of the frontal cortex on REM density. Similarly, two patients with atrophy of the frontal brain presented high REM density, leading King et al. to suggest the frontal cortex plays an inhibitory role on eye movements.

Doricchi and coworkers studied rapid eye movements in patients with left unilateral negligence syndrome, due to lesions on the right hemisphere, and found a suppression of eye movements toward the neglected side. They concluded that attention mechanisms, located in parietal lobes, influence rapid eye movements during sleep.

**The rapid eye movements, the cortical areas and dreams**

A more relevant participation of posterior brain areas on rapid eye movements was suggested by studies that employed neurophysiologic methods. Indirect evidence for the involvement of posterior regions on paradoxical sleep was obtained with analysis of the ability to recall dreams following encephalic lesions. Murri and coworkers studied a group of 53 patients with focal lesions of either vascular or neoplastic nature. The patients with posterior lesions showed more difficulty to recall dreams, associated with a performance deficit in visual-perceptive tests, than patients with anterior lesions. These results were recently confirmed. Doricchi and Violani hypothesized that inability to recall dreams is a consequence, not only of the loss of memory and verbal impairment, but also of a deficit of cognitive dream decoding, since the lesions impair cognitive processes related to speech, which is essential for processing internal information. Murri et al. showed, more specifically, that patients with posterior lesions, on temporal, occipital and parietal lobes, mainly in the right hemisphere, presented more difficulty to recall dreams.

Our results did not indicate greater influence of either hemisphere on REM density, since patients did not differ from normal volunteers in this respect. Several authors have reported the higher specialization of the right hemisphere as regards paradoxical sleep, specially concerning its
main role in the genesis of dreams. Neurophysiological studies, employing analysis of amplitude of each brain hemisphere electrical activity during sleep, showed greater participation of the right hemisphere. Similar results were reported with tests of hemisphere function after awakening from paradoxical sleep. Tests of tactile recognition after each sleep phase demonstrated the major importance of the right hemisphere during paradoxical sleep. More recently, the same authors emphasized that, although the right hemisphere plays a major role in spatial tests, there is no consistent evidence that it is more active during paradoxical sleep. Reinsel and Antrobus did not observe hemisphere lateralization, using specific tests for each hemisphere, after awakening from either non-paradoxical or paradoxical sleep. Similarly, Armitage et al. did not find support in the literature for the hypothesis of greater influence of the right hemisphere on paradoxical sleep. Hobson et al. related that during paradoxical sleep occurs an activation of the pontine brain stem and of limbic and paralimbic cortical structures involved in mediating emotion and a corresponding deactivation of dorsolateral prefrontal cortical structures involved in the executive and mnemonic aspects of cognition.

CONCLUSION
Our results, together with previous reports, suggest that cortical and subcortical structures, particularly in frontal regions of both hemispheres, exert an inhibitory influence on rapid eye movements generated from the brainstem. If dreams were related to eye movements, we could hypothesize that both phenomena are triggered by structures located in the brainstem and modulated by brain hemisphere. This hypothesis does not exclude the psychological content of dreams, that could be generated from cortical structures responsible for emotion.

REFERENCES


23. Feinberg I, Floyd TC, March JD. Effects of sleep loss on delta (0.3-3 Hz) EEG and eye movement density: new observations and hypotheses. Electroenceph Clin Neurophysiol 1987; 67:217-221.


