THE RELATION BETWEEN EEG PREFRONTAL ASYMMETRY AND SUBJECTIVE FEELINGS OF MOOD FOLLOWING 24 HOURS OF SLEEP DEPRIVATION

Camila Ferreira¹, Andréa Deslandes¹, Helena Moraes¹, Maurício Cagyu², Luiz Fernando Basile³, Roberto Piedade⁴, Pedro Ribeiro⁵

ABSTRACT - Several studies have investigated the relationship between asymmetrical EEG activity over the frontal cortex and mood. This study aimed at investigating the association between state fluctuations in frontal alpha EEG asymmetry and state changes followed by 24 h of sleep deprivation (SD). Our results show that sleep deprivation caused a significant alteration in the asymmetry values. Activation shifted from the left hemisphere, before SD, to the right hemisphere, after SD, in all frontal electrode pairs. In addition, according to the self-rating scale of SD-related mood effects, subjects became significantly less alert and active, and sleepier. According to these results, increased right prefrontal activation might be potentially associated with the negative mood states typically seen after sleep deprivation, although the causal relationship is still uncertain. However, more studies will be necessary to establish the viability of EEG asymmetry and the cerebral lateralization hypothesis to explain the SD-related affective changes.

KEY WORDS: EEG prefrontal asymmetry, sleep deprivation, affective disorders.

Relação entre assimetria pré-frontal no EEG e sensações subjetivas de humor após 24 horas de privação de sono

RESUMO - Diversos estudos têm investigado a relação entre a atividade assimétrica do EEG no córtex frontal e mudanças no humor. Adotando tal abordagem, o presente estudo teve como objetivo investigar a associação entre os estados de oscilação na assimetria frontal de alfa e mudanças no estado emocional ou motivacional após 24h de privação de sono. Os resultados mostram que 24h de privação de sono ocasionaram alterações significativas nos valores de assimetria. Ativação cerebral mudou do hemisfério esquerdo, antes da privação de sono, para o hemisfério direito, após a privação de sono, em todos os pares de eletrodos frontais. Além disso, de acordo com a escala relacionada aos efeitos subjetivos do humor após privação de sono, os sujeitos mostraram-se significativamente menos alerta e ativos e mais sonolentos. É possível que as duas variáveis estejam associadas, embora a relação causal seja ainda incerta. Estudos serão ainda necessários para que se possa estabelecer a viabilidade da assimetria pré-frontal e da hipótese de lateralização cerebral na elucidação de mudanças emocionais relacionadas à privação ou falta de sono.

PALAVRAS-CHAVE: assimetria pré-frontal, privação de sono, desordens afetivas.

Over the past years, clinical and laboratory observations suggest that frontal EEG asymmetry reflects individual differences in the regulation of an elicited emotion, a biological marker of affective style1-6. According to Petruzzello et al.6 EEG frontal asymmetry reflects a diathesis that, in concurrence with an emotion elicitor stimulus, will result in a change in positive or negative affect appropriate with the emotion-eliciting stimulus. Even though such evidence does not involve sleep deprivation (SD), there have been many studies examining EEG activity as a consequence of total or partial sleep deprivation which reported decreased alpha power and concomitant reduced alertness and increased irritability7-9.

However, up to now, no investigations have observed the relationship between motivational or mood changes after SD and prefrontal asymmetry. But, given the acknowledged SD-related negative

¹Laboratório de Mapeamento Cerebral e Integração Sensório-Motora, Instituto de Psiquiatria (IPUB), Universidade Federal do Rio de Janeiro, Brasil (UFRJ); ²COPE, UFRJ; ³Laboratório de Neurociências (LIM-27) Faculdade de Medicina da Universidade de São Paulo SP, Brasil; ⁴Coordenador Laboratório de Mapeamento Cerebral e Integração Sensório-Motora, IPUB, UFRJ; ⁵Escola de Educação Física e Desportos (EEFD) - Departamento de Biociências e Atividade Física, Laboratório de Mapeamento Cerebral e Integração Sensório-Motora, IPUB-UFRJ, Professor Pesquisador Universidade Castelo Branco (PRGHH-UCB).

Received 9 September 2005, received in final form 20 February 2006. Accepted 2 March 2006.

Dra. Camila Ferreira - Rua Delfina 47 / 104 - 20511-270 Rio de Janeiro RJ - Brasil. E-mail: camilaferreira@libero.it.
mood/affective changes and according to the cerebral lateralization hypothesis, we may well observe if these variables are correlated, helping to understand how the brain adjusts itself when confronted with sleep loss or sleep deprivation.

Therefore, the primary purpose of this study was to observe alterations in EEG prefrontal asymmetry after sleep deprivation; and if SD-related affective changes are associated with such electrophysiological adjustments. We predicted that more activation in the anterior portions of the right hemisphere, relative to the anterior portions of the left hemisphere should occur as a result of sleep deprivation, and that such modifications are related to enhanced negative affect.

**METHOD**

**Subjects** – The sample consisted of 11 individuals, 4 males and 7 females, with ages varying between 21 and 40 years (30.8±6.0 years), weighing between 55 and 92 kg (70.5±17.2 kg). All subjects were healthy, right-handed, non-smokers, free of cognitive deficits and were not making use of oral contraceptives or any psychoactive or psychotropic substance at the time of the test. To ensure that subjects did not present any impairment of their physical and mental health, a questionnaire was applied. The questionnaire aimed at identifying food intake, body temperature, fatigue, and drugs use, among others. Subjects were not allowed to consume any alcoholic beverages or caffeine. The Edinburgh inventory was used to assess laterality and exclude left-handed individuals from the experiment. All subjects signed a consent form, where the experimental condition was thoroughly described. The Psychiatric Institute's Ethics Committee approved the experiment.

**Study design and procedures** – The procedures were standardized in two different times; pre- and post-sleep deprivation, following the subsequent routine: subjects arrived at the laboratory by 8:00 p.m., performed a baseline eight-minute EEG recording (4 minutes with eyes closed and 4 minutes with eyes open) and answered to an 11-item sleep deprivation-related mood changes questionnaire (better described below). An abbreviated form was favored because of the several estimations that were made along which objected a rapidly conclusion, allowing a close time correspondence with EEG measures. It is relevant to remind that the questionnaire used here attempted to reflect mood changes related only to the sleep deprivation effects, reproducing pleasant (active, energetic, alert, satisfied) and unpleasant states (sleepy, drowsy, tired, lazy, depressed). Nevertheless, the same elicitor will produce an array of different emotions across subjects, even in response to elicitors that are specifically chosen to target particular emotions. During all night long, volunteers played games, watched videos and carried out recreational activities, and were also fed each three or four hours. Two experimenters were required to ensure continuous wakefulness of the subjects monitored sleep deprivation continuously. Subjects abstained from smoking or drinking xanthine-containing beverages (coffee, tea, cola or soft drinks). No concurrent treatment was allowed during the study. At 7:00 a.m. (Time 2 - after sleep deprivation) of the subsequent morning, all subjects repeated the same identical EEG recording and answered on more time the mood-related changes questionnaire.

**Data acquisition** – The study design respected the International Pharmacology—EEG group guidelines. During the task, all lights remained turned off to minimize visual stimuli interferences, besides the video monitor. Individuals sat comfortably in a large supported chair, in order to also minimize also muscular artifacts inside this sound-light-attenuated room, while EEG was collected from 20 monopolar derivations for eight minutes (eyes closed, alert but resting). Data were collected with eyes closed in order to observe the cortex electrical activity without any external stimuli, minimizing possible visual artifacts. Electrodes were positioned according to the International 10/20 System (referred to linked earlobes with ground at FPZ). All electrode impedances were kept below 5 kΩ. The signal was amplified with a gain of 22,000, analogically filtered between 0.01 Hz (high-pass) and 100 Hz (low-pass), and sampled at 240 Hz using a BrainTech-3000® (EMSA-Medical Instruments, Rio de Janeiro, Brazil) EEG acquisition system. The EEG was recorded by means of the software ERP Acquisition (Delphi 5.0®, Borland-Inprise), developed at the Brain Mapping and Sensorimotor Integration Lab, employing the following digital filters: notch (60 Hz), high-pass of 0.3 Hz and low-pass of 30 Hz. Visual inspection was employed for detection and elimination of artifacts. Eye-movement (EOG) artifact was monitored with a bipolar electrode montage using two 9-mm-diameter electrodes attached superior to and on the external canthus of the right eye.

**Data analysis: Alpha asymmetry** – EEG containing artifact was marked and excluded from each EEG trial prior to further analysis of the data. All artifact-freedata extracted from the EEG’s total recordings were subjected to a Matlab 5.3® (The Mathworks Inc., Massachusetts, USA) routine to perform spectral power analysis, producing estimates of absolute spectral power (µV2) and asymmetry for the alpha frequency band. A Fast Fourier Transformation (FFT) was applied, analyzing the repetitive data during regular time intervals. Therefore, through the FFT, it is possible to define how much energy (power) exists in a given frequency band. Afterwards, a natural log transformation was applied to power density values to normalize the distribution. Power spectral analysis had to be carried out, in order to extract the asymmetry values. Subjects had a minimum of 25 artifact-free epochs for each experimental time. After the data acquisition and storage, all statistics were computed to extract the asymmetry values for the alpha frequency band in the frontal areas of the brain cortex (Fp1-Fp2, F3-F4, and F7-F8). Although the software provides us with power spectral data from all electrode sites in the four frequency bands, we have focused our analysis on the three anterior
electrode pairs, because previous researches\cite{2,3,15} showed relations between brain electrophysiology at these sites and measures of affective states. In addition, power in the alpha band (typically in the 8-13 Hz) is inversely related to activation and has been the measure most consistently obtained in studies\cite{14} of EEG asymmetry. Alpha power has been found to be more reliably related to task performance compared with power in other frequency bands, when the tasks that are compared are carefully matched on psychometric properties and since it is inversely related to activation, blocking of or decreases in alpha are seen\cite{16} when underlying cortical systems engage in active processing. Moreover, alpha asymmetry has been found\cite{15} to be relatively stable over a three-week interval leading\cite{1} Davidson suggested that this measurement reflects a trait-like tendency to respond differentially to positive (i.e. approach-related) and negative (i.e. withdrawal-related) stimuli. Therefore, we expected that the SD-related affective changes would differ on measures of alpha power asymmetry, especially in anterior regions. The alpha power asymmetry may be considered a gradient of power that exists between the two homologous electrodes in the pair, with the slope of the gradient being towards the electrode with the greatest amount of power in this frequency band. Computational indices of power asymmetry provide a left-right comparison in the alpha frequency band, as well as homologous bipolar pairs and across the specified multivariate electrode arrays (for example, left hemisphere versus right hemisphere power asymmetry). The basic mathematical formula\cite{1} (5.3) The Mathworks Inc., Massachusetts, USA) for asymmetry calculations is % Asymmetry = \((L-R)/L+R \times 100\). Where L refers to the left homologous electrode in the pair of compare electrodes, and R is the right homologous electrode.

Self-rating scale of mood effects – This is an 11-item four-point self-rating scale utilized\cite{17} in sleep deprivation-paradigms. Motivational or affective states in this study should be related to sleep deprivation or sleep loss. The most frequent emotions considered in asymmetry studies, such as happy or sad wouldn’t make sense in this paradigm. The subjective effects evaluated should be related to the emotion elicitor\cite{14}, although the same elicited produces an array of unintended emotions across subjects. For example, when investigating hemispheric activation and exercise-related affective states, Petruzzello\cite{16} and Landers and Petruzzello et al.\cite{15} assessed pleasant emotions related to the elicitor, exercise, such as, reduced anxiety or increased energetic arousal. In the present study, individuals reported their subjective state in terms of the eleven clusters of the following effects: alert, satisfied, nervous, sleepy, active, depressed, talkative, headache, upset stomach, lazy, irritable.

Statistical analysis – Data were expressed as mean ± s.d. The statistical software SPSS for Windows was used for all data analysis. Asymmetry values were submitted to a nonparametric Wilcoxon analysis, in order to observe variations in the hemispheres activation pre and post 24h of sleep deprivation. A paired t-test was performed to compare the questionnaire scores in the two experimental times (pre- and post-sleep deprivation). Sleep deprivation effects on asymmetry-related brain activation were examined comparing EEG data collected pre sleep deprivation with the first post sleep deprivation measures. Significance levels were set at \(p<0.05\).

RESULTS

Electrophysiological results – Twenty-four hours of sleep deprivation caused an activation shift from the left hemisphere to the right hemisphere in all frontal and prefrontal electrode pairs analyzed in this study. Statistical analysis comparing all pre and post sleep deprivation asymmetry values showed a significant increase in hemispheric asymmetry. According to the mathematical formula from the software utilized in this study (Matlab 5.3 The Mathworks Inc., Massachusetts, USA) the preponderance of negative asymmetry values observed after sleep deprivation demonstrates that the activation in the right side of the frontal cortex was greater, therefore, creating the distinctive electrophysiological patterns observed: Asymmetry = \((L-R)/(L+R \times 100\). Statistical analysis demonstrated a significant difference in hemispheric asymmetry before and after SD in the three electrode pairs representing the frontal cortex: Fp1 Fp2 p= 0.026, F3 F4 p= 0.004 and F7 F8 p=0.021. The energy shift in the alpha frequency band from the left hemisphere to the right hemisphere after 24 h of SD resulted in greater right-side activation (negative asymmetry values), which could be related to

![Fig 1. Twenty-four hours of sleep deprivation (SD) caused a shift in hemispheric activation resulting in greater right side activation, which is confirmed by the asymmetry negative values (not specifically shown here) observed after SD. Statistical analysis p reduced the following p values 0.026, 0.004 and 0.021, representing electrode pairs Fp1-Fp2, F3-F4 and F7-F8, respectively. Pre SD=before sleep deprivation; Post SD=after sleep deprivation.](image-url)
the negative emotional state often seen after sleep loss\textsuperscript{9-12}. Electrophysiological results from the three prefrontal electrode pairs are shown in Figure 1. Figure 2 has illustrative purposes only and represents sites of greater and lower activation in the frontal cortex after sleep deprivation. It is perceptible that the electrodes from the right frontal hemisphere experienced higher activation following SD.

\textit{Subjective evaluation of mood} – To observe the effects of sleep deprivation on mood or affective sta-
tes, a t-test was applied. The statistical analysis examined the following mood/affective changes related to sleep deprivation: alertness, satisfied, sleepy, active, depressive, talkactive, headache, upset stomach, lazy and irritable. After 24 h of SD subjects became significantly less alert (p=0.01) and active (p=0.04) and more sleepy (p=0.001). According to Davidson et al.14, the subjective effects evaluated should be related to the emotion elicitor, which in this case is the deprivation or lost of sleep, and the emotion elicitor-negative affect or state should be associated with the withdrawal system. The significant effects of sleep deprivation on affective states are demonstrated in Figure 3.

DISCUSSION

Total or partial sleep deprivation causes significant performance impairment throughout different cognitive and psychomotor tasks and is also related to reductions in alertness, sustained and selective attention levels7,9-11. Besides cognitive deterioration, SD is also followed8,10 by changes in mood or emotional states, often resulting in negative affective states. Therefore, the primary purpose of this study was to observe the cerebral lateralization hypothesis during an investigational test. In addition, we tried to evaluate if post SD related negative affective/mood changes are associated with increased right prefrontal hemisphere activation. According to Petruzello et al.9, an emotion elicitor stimulus of sufficient intensity, will result in a change in positive or negative affect, which is associated with EEG frontal asymmetry. Following Davidson’s5,14 approach-withdrawal model and taking SD as an emotion elicitor, we predicted that post SD-related negative affective changes would be associated with an EEG state frontal asymmetry specifically, a greater right frontal hemisphere activation in the alpha frequency band. In this study, the alpha band (typically in the 8-13 Hz) has been chosen as an electrophysiological marker because, as quoted before, it is relatively stable14,15 and inversely related to activation. Hence, it has been the measure most consistently obtained in studies of EEG asymmetry, leading14,5 Davidson to suggest that this measurement reflects a trait-like tendency to respond differentially to positive (i.e., approach-related) and negative (i.e., withdrawal-related) stimuli. Furthermore, EEG power spectra studies have generally reported a decrease in alpha power (increased activation) after SD, which could account for the negative mood variations frequently observed12,13 following sleep loss or sleep deprivation. Since influential previous investigations demonstrated a relationship between brain electrophysiology in prefrontal area and measures of affective states14,20, we have also focused our analysis on the three anterior electrode pairs, Fp1-Fp2, F3-F4, F7-F8.

The results demonstrated that 24h of SD caused significant changes in EEG asymmetry, producing an activation shift from the left hemisphere to the right hemisphere in all anterior electrodes pairs, which is represented by the negative asymmetry values in the post SD time. SD-related mood variations extracted from the self-report mood scale in the two experimental times reported a significant increase in sleepiness and reduction in active and alertness levels. The association between the contextual negative emotional states observed after sleep deprivation and the increased right frontal activation in the alpha band might be related to Davidson’s withdrawal hypothesis20 and could partially explain the negative effects often seen after sleep loss. According to Davidson, the right prefrontal cortex is a biological substrate of the withdrawal behavior, inhibition and negative emotional states, mediating also attention and alertness (toward the stimulus). Our results demonstrate a likely association between negative emotional states, particularly elicited by sleep deprivation, and increased right prefrontal activation, although it is not entirely clear if prefrontal asymmetry plays a causal role in the outcome measures. In other words, it is still unsure if the shift in hemispheric activation seen after sleep deprivation produced the negative motivational or mood effects or if it is the contrary. Hence, the small sample presented here could have influenced our results. It is, therefore, essential to continue other investigations using a larger number of individuals. It is relevant to remind that the present study did not attempt to examine if EEG asymmetry measured during resting conditions (trait frontal asymmetry) could predict such affective responses to the emotion elicitor (sleep deprivation). Instead, it has followed a different research approach where EEG asymmetries (state frontal EEG asymmetry) may be thought of as those that are responsive to specific environmental conditions.

In conclusion, it appears from the present findings that changes in brain activation, characterized by increased right prefrontal activation, might at least partially explain the negative mood states typically seen after sleep deprivation, and therefore, the theory of individual differences in response to this specific emotion elicitor may be hypothesized. In ad-
dition to replicating the present findings, future work also needs to examine SD-related affective responses of subjects with: 1) previously higher levels of negative affect, 2) suffering from mood or anxiety disorders and 3) resting EEG asymmetry as a predictor of emotional responses to sleep deprivation or sleep loss. Measuring the differences in individual response to SD-mood effects might be interesting and useful, given the involvement of almost all psychopathologies in the sleep regulation.

REFERENCES