**Delta EEG Activity in Left Orbitofrontal Cortex in Rats Related to Food Reward and Craving**

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Abstract: The orbitofrontal cortex (OFC) is particularly important for the neural representation of reward value. Previous studies indicated that electroencephalogram (EEG) activity in the OFC was involved in drug administration and withdrawal. The present study investigated EEG activity in the OFC in rats during the development of food reward and craving. Two environments were used separately for control and food-related EEG recordings. In the food-related environment rats were first trained to eat chocolate peanuts; then they either had no access to this food, but could see and smell it (craving trials), or had free access to this food (reward trials). The EEG in the left OFC was recorded during these trials. We showed that, in the food-related environment the EEG activity peaking in the delta band (2-4 Hz) was significantly correlated with the stimulus, increasing during food reward and decreasing during food craving when compared with that in the control environment. Our data suggests that EEG activity in the OFC can be altered by food reward; moreover, delta rhythm in this region could be used as an index monitoring changed signal underlying this reward.

Key words: Orbitofrontal cortex; EEG; Reward; Craving; Delta band

The orbitofrontal cortex (OFC) receives efferent projections of the nucleus accumbens (Ray & Price, 1993; Koob & Bloom, 1988) and the ventral tegmental area (Oades & Halliday, 1987). The mesolimbic dopamine
(DA) pathway, which projects from the ventral tegmental area to the nucleus accumbens, is important for mediating the perception of reward (Berridge, 1996; Rolls, 2004). Previous studies have suggested that the left OFC rather than the right is associated with a strong reward bias (Pizzagalli et al, 2005).

Electroencephalogram (EEG) is a sensitive measure to study the global activity of the brain. Evidence shows that EEG activity is associated with many important functions, such as learning and memory (Basar et al, 2001; Basar et al, 1999). EEG activity in the OFC is altered during the development of drug dependence and tolerance (Sosina, 1993; Wikler & Altschul, 1950; Grasing & Szeto, 1993; De & Caballero, 1989), such as with morphine (Liu et al, 2005; Sun et al, 2006). However, data related to EEG activity in the OFC during the development of reward and craving of general substances, such as food, is limited.

Previous studies have indicated that eating behavior is mainly motivated by the reward value of the food, which was available in large amounts, rather than by feelings of hunger or maintaining homeostasis (Berridge, 2004). In addition, food craving was driven by the presence of stimuli, which also predicted the rewarding effect of food (Rolls, 1997).

In this study, we recorded EEG activity in the left OFC during the development of food reward and craving to further our understanding of the changes in EEG activity in this region in the process of general substance reward.

1 Methods

1.1 Animals

Experiments were performed on ten male Sprague-Dawley rats (Animal House Center, Kunming Medical College, PR China) weighing 300–350g. They were housed in individual cages under constant temperature and humidity conditions on a 12-h light/12-h dark cycle. The amount of rodent chow was adapted in order to maintain nearly constant body weight. Water was freely available all the time. All experimental and animal care procedures were carried out in accordance with the guidelines for the National Care and Use of Animals approved by the National Animal Research Authority.

1.2 Surgery

As previously described (Sun et al, 2006), surgery was performed under pentobarbital sodium (40mg/kg; Shanghai Chemical Factory, PR China) anesthesia. After a midline scalp incision, a burr hole was drilled in the skull over the left OFC. A Teflon-insulated recording electrode was implanted in the left hemisphere OFC according to the following stereotaxic coordinates: (relative to bregma) anteroposterior (AP), 3.7 mm; mediolateral (ML), 2.4 mm; dorsoventral (DV), 4.6 mm (Paxinos & Watson, 1998). Two stainless steel wash screws implanted in the bone above the right cerebral regions served as both reference (AP, 3mm; ML, 2mm) and ground (AP, −5mm; ML, 2mm) electrodes. Two or more additional support screws were positioned, and the entire ensemble was secured to the skull with dental acrylic. All electrodes were attached to male pins that were secured in a rectangular three-by-one pin array and secured with dental acrylic. In addition, a general penicillin antibiotic was injected (50,000 units per rat; i.m.; Harbin Pharmaceutical Group, PR China) immediately after the surgery. Rats were allowed at least one week to recover from surgery. The locations of the recording electrode were confirmed histologically after the EEG experiment. Data were excluded from any rats in which the recording locations were misplaced.

1.3 Apparatus for EEG recording

With the socket on the head of the rat, the electrodes were first connected by a cable to an amplifier (bandpass: 0.5–100 Hz, with no 50 Hz filter). The cable was suspended to allow the rat free movement. The amplified EEG signals were digitized with an AD (analog to digital) board (biphase, 1000 Hz), and then displayed instantly and saved by a computer.

EEG recording was performed in two chambers (60 cm×30cm×50cm). Chamber A was painted white with a textured floor, which served as the control environment. Chamber B was painted with dark stripes (5 cm width), and the floor was smooth, which served as the stimulus-related environment.

1.4 Experimental procedure

The experimental procedure consisted of three phases: habituation, training and testing (Fig. 1). EEG recording was performed during the testing phase.

Habituation: rats were placed into chamber A for 30 min before being removed. After 15 min, they were placed in chamber B for 30 min. This procedural sequence represented a trial, and took place on each of the five habituation days. In this phase no food was offered in both chambers.

Training: following habituation, the same procedure was conducted except that the rats stayed in the chambers for 5 min only. In addition, rats were given free
access to chocolate peanuts in chamber B, as well as a perforated bottle containing the same chocolate peanuts. This training phase continued over 6 days, with one trial taking place in the morning and the other in the evening. In all, food was always offered in chamber B but not in chamber A.

Fig. 1  Schematic illustration of the experimental procedure

The experimental procedure included three phases: habituation, training and testing. “R”, the rats were given free access to chocolate peanuts; “C”, the rats were presented with only a perforated bottle containing chocolate peanuts allowing no access to food; “a”, the trial was performed in the morning; “p”, the trial was performed in the evening. EEG recording was performed during the testing phase.

Testing: the same procedure was applied in the testing phase, with an exception of including craving trials. One craving trial was followed by two reward trials. This phase comprised 18 trials in total, including 6 craving trials and 12 reward trials. The reward trials were the same as trials in the training phase. The only difference between the reward trials and the craving trials was that in the craving trials, the rats were presented with only a perforated bottle containing chocolate peanuts allowing no access to the food. In summary, there was no food in chamber A, but in chamber B food was offered during reward trials and not offered during craving trials.

1.5  Data analysis and statistics

As previously described (Liu et al, 2005), EEG activity was examined by off-line analysis and spectral analysis was performed using Matlab 6.5 software. After computer-assisted rejection of EEG segments with artifacts, the relative power across various frequency bands were characterized for (1) delta: 2–4Hz, (2) theta: 4–8 Hz, (3) alpha: 8–12 Hz, (4) beta: 12–20 Hz and (5) gamma: 20–100Hz.

Based on our experimental design, for each rat, the relative EEG power in each frequency band was averaged for 12 reward trials and for 6 craving trials in chamber A and chamber B. Pearson correlations were used to assess the relationship between changes in relative EEG power and food-related stimuli. Differences were assessed with analysis of variance (ANOVA) with repeated measures, using the SPSS 13 statistical software. Stimulus (craving and reward) and chamber (A and B) were two within-group factors. Data were expressed as mean ± SEM. A level of \( P<0.05 \) was considered significant.

2  Results

As shown in Fig. 2, during the testing phase the two food-related stimuli, craving and reward, were used for animals (\( n=9 \)) in chamber B. Pearson correlation analysis revealed no correlation between the relative EEG power and the stimulus, in any frequency band, in chamber A (\( r=-0.08, 0.02, 0.09, 0.2 \) and \(-0.07 \) for delta, theta, alpha, beta and gamma bands, respectively; \( P>0.05 \) for all). In contrast, in chamber B, there was a positive correlation between the relative power and the stimulus in the delta

Fig. 2  Food craving and reward during the testing phase induced changes in relative EEG power in the delta frequency band in chamber A and B

a: Pearson correlation analysis showed that the relative power in the delta band showed a significant stimulus-dependent change in chamber B but not in chamber A; b: The mean relative power in the delta band decreased during food craving and increased during food reward in chamber B, compared with that in chamber A. *\( P<0.05 \); ** \( P<0.001 \).
frequency band ($r=0.75$, $P<0.001$), but not in the other frequency bands ($r=-0.25$, $-0.34$, $0.28$ and $-0.07$ for theta, alpha, beta and gamma bands, respectively; $P>0.05$ for all).

ANOVA with repeated measures on the delta frequency band showed a significant main effect of stimulus [$F(4, 8)=75.98$; $P<0.001$] and chamber [$F(4, 8)=8.29$; $P<0.05$] and interaction effect of stimulus $\times$ chamber [$F(16, 32)=69.41$; $P<0.001$]. Further analysis with a paired Student's $t$-test revealed that the relative power in the delta band decreased during food craving and increased during food reward in chamber B, when compared with that in chamber A.

3 Discussion

In this study, the EEG activity in the left OFC was investigated in rats during the development of food reward and craving in a certain environment. In this environment, a significant correlation between the relative EEG power and the food-related stimulus was observed in the delta frequency band only. Moreover, this frequency band showed increased power during food reward and decreased power during food craving when compared with that in the control environment.

Our finding was that the activity in the delta band showed significant correlation with the food-related stimulus in a certain environment, reflected by relative power increased during food reward and decreased during food craving when compared to that in the control environment. Firstly, our data are in accordance with studies on cocaine in humans. For example, a reduction in delta power in the frontal cortex during touching and seeing cocaine cues has been observed, suggestive of craving, while an increase was found during guided imagery of cocaine, indicative of analogical reward (Reid et al, 2003). Cocaine also produced an increase in delta coherence over the prefrontal cortex, indicating the involvement of delta activity in the rewarding properties of cocaine (Reid et al, 2006). In addition, a decrease in delta power during withdrawal in cocaine-preferring polysubstance abusers suggested craving (Roemer et al, 1995). Secondly, during food deprivation, a decrease in delta power has been found in humans (Sulimov, 1995).

Indeed, the delta band has been linked with reward motivation by an evolutionary interpretation based on brain oscillations (Knyazev & Slobodskaya, 2003). For example, the delta activity increased after administration of testosterone (Schutter & Van Honk, 2004), which has been related to reward motivation in humans and animals (Van Honk et al, 2004). On the other hand, the OFC has an important relationship with the mesolimbic DA reward system, mediating the perception of reward (Berridge, 1996; Rolls, 2004). Previous studies have shown that the OFC dysfunction is associated with disturbances in reward motivation (Rolls, 2004). Therefore, in the present study, EEG activity peaking in the delta band in the OFC was probably associated with the reward and craving bias towards food.

In our previous studies, EEG activity in the OFC in monkeys decreased after morphine injection, and this decrease lasted for a longer period in the gamma frequency band (Liu et al, 2005). Moreover, in the OFC in rats, gamma activity showed decreases and increases during the morphine administration and withdrawal period, respectively (Sun et al, 2006). In the present study, EEG activity, especially peaking in the delta band, could be affected by food reward and craving, which is in agreement with our previous studies. On the basis of these findings, we confirm that the OFC plays an important role in the reward motivation either for morphine or for food, which could be monitored by EEG spectrum.

Although the EEG recording was well-controlled by disassociating the effects of body and jaw movements while the subjects were moving or eating, chewing or sniffing artifacts may still have contaminated our EEG signals. However, previous studies and our own data suggest that the effects of these additional factors on the present results should be minor. The rate of chewing pellets in adult male rats varies between 4 and 6 Hz (Wejs, 1975). This frequency range was again tested, and was found to increase with age in infant rats, and by 18 or 21 days of age, reached the rate of 4 – 5 Hz (Westneat & Hall, 1992). Besides, sniffing and other rhythmic movements are always associated with the theta band of the EEG (Kepecs et al, 2006). Taking all this into account, if these movement artifacts affected our EEG signals, the activity in the theta band would be expected to show some correlation with food stimulus. However, this frequency band revealed no significant alterations in this study.

In addition, we only recorded the left OFC, but several studies have shown the possibility that activity localized in the OFC may represent more generalized EEG activity (Zuo et al, 2007). Therefore, whether our findings in this region could be found in other brain regions, as well as in more global activity, are interesting subjects for future investigation.
In conclusion, EEG activity in the left OFC in rats seems to be altered by food reward, and delta EEG activity in this region could be an index that monitors changed signals underlying this reward.

References:


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