Oscillatory interaction between amygdala and hippocampus coordinates behavioral modulation based on reward expectation

Terada, Satoshi; Takahashi, Susumu; Sakurai, Yoshio

Terada, Satoshi ...[et al]. Oscillatory interaction between amygdala and hippocampus coordinates behavioral modulation based on reward expectation. Frontiers in Behavioral Neuroscience 2013, 7: 177.

http://hdl.handle.net/2433/179544

© 2013 Bernheim, Halfon, Boutrel. This is an open-access article distributed under the terms of the Creative Commons Attribution License (CC BY). The use, distribution or reproduction in other forums is permitted, provided the original author(s) or licensor are credited and that the original publication in this journal is cited, in accordance with accepted academic practice. No use, distribution or reproduction is permitted which does not comply with these terms.
Oscillatory interaction between amygdala and hippocampus coordinates behavioral modulation based on reward expectation

Satoshi Terada1‡*, Susumu Takahashi2‡ and Yoshio Sakurai1‡

1 Department of Psychology, Graduate School of Letters, Kyoto University, Kyoto, Japan
2 Laboratory of Neural Circuitry, Graduate School of Brain Science, Doshisha University, Kizugawa, Japan

**Correspondence:**
Satoshi Terada, Department of Psychology, Graduate School of Letters, Kyoto University, Yoshida-hon-machi, Sakyo-ku, Kyoto 606-8501, Japan
e-mail: steradamm@gmail.com

‡These authors have contributed equally to this work.

INTRODUCTION

The aim of this study is to examine how the amygdala and hippocampus interact for behavioral performance modulated by different Reward-expectations (REs). We simultaneously recorded neuronal spikes and local field potential from the basolateral amygdala and hippocampal CA1 while rats were performing a light-side discrimination task with different expectations of a high or low probability of reward delivery. Here, we report the following results. First, the rats actually modulated their behavioral performance on their expectations of a high or low probability of reward. Second, we found more neurons related to RE in the amygdala and more neurons related to task performance in the hippocampus. Third, a prominent increase in the coherence of high-frequency oscillations (HFOs) (90–150 Hz) between the amygdala and the hippocampus was present during high RE. Fourth, coherent HFOs during inter-trial intervals and theta coherence during trials had significant correlations with the behavioral goal-selection time. Finally, cross-frequency couplings of LFPs within and across the amygdala and hippocampus occurred during ITI. These results suggest that the amygdala and hippocampus have different functional roles in the present task with different REs, and the distinctive band of coherence between the amygdala and the hippocampus contributes to behavioral modulation on the basis of REs. We propose that the amygdala influences firing rates and the strength of synchronization of hippocampal neurons through coherent oscillation, which is a part of the mechanism of how reward expectations modulate goal-directed behavior.

Keywords: reward expectation, behavioral modulation, oscillation synchrony, amygdala, hippocampus

---

In the present study, we focused on oscillatory interactions between the amygdala and the hippocampus. The amygdala is a well-known reward-related brain region and is critical for RE (Blundell et al., 2001; Savage and Ramos, 2009; Morrison and Salzman, 2010). The hippocampus, on the other hand, is not critical for RE (Ramirez and Savage, 2007) but has important roles in the learning and performance of behaviors. Also, hippocampal oscillations are correlated with the improvement of task-related behavioral performance (Rutishauser et al., 2010; Ahmed and Mehta, 2012). The hippocampal functions for behavioral performance have been shown to be affected by reward (Holscher et al., 2003; Tabuchi et al., 2003; Singer and Frank, 2009). Moreover, the interaction between the amygdala and the hippocampus is thought to be involved in the emotional modulation of behavior. For example, a recent disconnection study indicated that the amygdala-hippocampus interaction is essential for context-induced cocaine-seeking behavior (Wells et al., 2011). In Pavlovian fear conditioning, these brain regions were proposed to communicate with each other via the coherent theta oscillations (Pape et al., 2005; Lesting et al., 2011). Electrical stimulation of the amygdala was shown to be able to produce alterations in the firing properties of hippocampal place cells (Kim et al.,
Despite these previous observations, few studies have directly specified the role of coherent oscillations between the amygdala and the hippocampus in modulating behaviors based on RE.

In this study, we simultaneously recorded neuronal spikes and local field potential (LFP) from the amygdala and hippocampus while rats were performing a reward-probability-biased discrimination task. We hypothesized that the amygdala represents the RE for different probabilities of reward and the hippocampus represents the appropriate behavioral performance required for the task. The activities of the amygdala and hippocampus are expected to exhibit synchrony in oscillations when the REs modulate the behavioral performance of the task.

MATERIALS AND METHODS

SUBJECTS

Eight male Wistar albino rats (Shimizu Laboratory Supplies, Kyoto, Japan), each weighing 400–480 g at the time of the experiment, and housed in a 25 × 15 × 24 cm cage, were used as experimental subjects. All rats were handled extensively, provided with a sufficient amount of lab chow 1–3 h after each daily training or recording session to maintain approximately 85% of their ad libitum weight during daily training or recording sessions, and allowed free access to water. They were exposed to light between 08:00 and 21:00 h each day. All experiments were conducted between 10:00 and 20:00 h in accordance with the guidelines presented in Guidelines for Care and Use of Laboratory Animals at Kyoto University (2007) and with the approval of the Animal Research Committee of Kyoto University.

APPARATUS

In a dim, sound-attenuated, electrically shielded box (Japan Shield Enclosure, Osaka, Japan), rats were trained in a behavioral task in a 22 × 32 × 45 cm operant chamber (Ohara Ika, Tokyo, Japan). One wall of the chamber had three 15-mm-diameter illuminated sensor holes in the horizontal direction 60 mm above the floor to detect the nose-poke behavior of rats. Access to the left and right holes was controlled using a guillotine door immediately in front of each hole. A food dispenser behind the wall delivered 25 mg food pellets to a food magazine located at the center of the wall and 10 mm above the floor. The dispenser delivered pellets with an intermittent low buzzer tone (reward tone). Another buzzer was located behind the food dispenser to present a continuous high buzzer tone (error tone) when the rats made erroneous responses. Visual stimuli were presented on the left or right wall using a light-emitting diode (LED). Auditory stimuli consisting of two pure tones (2 or 10 kHz) of approximately 70 dB SPL were presented via a loudspeaker (15 cm in diameter) set 30 cm above the top of the operant chamber. The task was controlled and the behavioral data were recorded using a personal computer (NEC, Tokyo, Japan). Each rat’s behavior was monitored using a video camera (Sony, Tokyo, Japan), and the extracellular neuronal activity from 24 channels was simultaneously amplified using a multichannel amplifier system (Nihon Kohden, Kyoto, Japan) and stored in a custom-made personal computer.

LIGHT-SIDE DISCRIMINATION TASK

Rats were trained to perform a reward-probability-biased light-side discrimination task (Figure 1A). Each trial started with the rat holding its nose in the central hole for 1000–1400 ms. Then the left or right LED was illuminated for 1000 ms (fixation phase) during which the rat continued to keep its nose in the hole. After turning off the LED illumination, the rat was required to poke its nose into the left or right hole illuminated by the LED (selection phase). The correct response was always indicated by presentation of the reward tone for 1000 ms. The probability of delivery of the reward (pellet) accompanying the reward tone was determined by the following reward probability (RP) conditions. Each trial was separated by an intertrial interval (ITI) of 5000 ms.

REWARD PROBABILITY CONDITIONS

In each session, we divided the total of 100 trials into 4 blocks, each of which consisted of 25 trials with one RP condition (Figure 1B). We prepared high (H) and low (L) RPs during which 10 and 2 kHz pure tones were constantly presented, respectively (Figure 1A). Under the H–RP condition, the probability of reward delivery was 80% for all correct trials. Only reward tones were presented in the remaining 20% of correct trials. Under
the L-RP condition, the probability of reward delivery was 20% for all correct trials and in the remaining 80% of correct trials, only reward tones were presented. A session contained two blocks with the H-RP condition and two blocks with the L-RP condition. The order of blocks with H- or L-RPs in one session was H–L–H–L, L–H–L–H, H–L–L–H, or L–H–H–L, and was randomly varied day by day. We carried out two trials in an inter-block interval (IBI), during which neither the high nor the low tone was presented and the rats obtained no reward in all trials.

PROBE TEST
After all recording sessions under the above conditions, we implemented a probe test session for each rat to confirm that the rats actually predicted the high and low probabilities of reward delivery using the presented high and low tones (Figure 1C). No reward was delivered during all trials throughout the session. The extinction phase (5 min), no tone was presented and the rats were able to start each trial with no ITI. Following the extinction phase, probe test blocks were implemented. One block and the IBI were 2 min each. The procedure of the probe test was identical to that of the extinction phase but the high or low tone was constantly presented during each block. The probe test blocks were H–L–H–L or L–H–L–H in the session.

DATA ANALYSIS OF BEHAVIOR
To investigate whether the rats modulated their behavior in accordance with their REs, selection times and correct rates under the different H- and L-RP conditions were analyzed. The selection time was defined as the time taken by the rats to select the right or left hole in the selection phase (Figure 1A). To confirm that the rats predicted the high or low probability of reward delivery on the basis of the high or low tone, the number of trials started by the rats under the different RP conditions indicated by the high and low tones in the probe test was analyzed. ANOVA was used to compare the differences between the reward probabilities (H-RP condition and L-RP condition). To exclude position bias, we added the cue direction (right and left) for the nose poke as the second variable to exclude position bias in the analysis of the selection time.

ELECTRODE CONSTRUCTION
Neuronal recording was performed with tetrodes (Wilson and McNaughton, 1993), each of which was composed of four tungsten microwires (12.5 micron in diameter; California Fine Wire, Grover Beach, CA). The four microwires were mounted in a 33 gauge stainless-steel cannula (Small Parts, Miami, FL) with 500 micron of the tip protruding. The tips were cut at right angles with sharp surgical scissors. The tip impedance was approximately 400 kΩ at 1 kHz. Two or three cannulas with tetrodes were attached in a row to construct an array of tetrodes, with a center-to-center spacing of 500 micron between the cannulas. The array of tetrodes was mounted on a microdrive assembly (McNaughton et al., 1989; Sakurai, 1993, 1994, 1996, 2002; Sakurai and Takahashi, 2006) designed to allow fine movements of the cannulas with tetrodes and stable recording of multineuronal activity for extended periods.

IMPLANTATION SURGERY
Details of surgery and data collection have been described previously (Takahashi and Sakurai, 2009a,b; Sakurai and Takahashi, 2013). After the completion of behavioral training, each rat was anesthetized with sodium pentobarbital (40 mg/kg) in preparation for the attachment of two microdrives, each of which contained two or three tetrodes, to the skull surface. After holes were drilled into the skull for electrode implantation into the basolateral amygdala (−2.28 mm from the bregma and 3.0 from the midline) and the hippocampal CA1 (−3.5 mm from the bregma and 2.5 mm from the midline), the tips of the tetrodes were implanted into the brain to a depth of approximately 4000 micron prior to the basolateral amygdala and 2000 micron prior to the hippocampal CA1. The reason for the small depth of the tetrodes targeting the amygdala was to decrease the damage to the brain and to obtain stable and long-term recordings. The craniotomy was filled with white petrolatum to a level slightly above the exit of the tetrodes from the skull surface. After the supports of the microdrives and cannulas were coated with a thin film of white petrolatum, the entire assembly was embedded in dental cement on the skull surface. A recovery period of about a week was assigned after surgery. We performed microdrive lowering of the tetrodes post-surgery to obtain stable long-term recordings as in the previous studies (Takahashi and Sakurai, 2009a,b; Sakurai and Takahashi, 2013).

DATA COLLECTION
Brain activity data were recorded and stored on a hard disk of a personal computer at a 20 kHz sampling rate while the rats were performing the behavioral task. Head stages containing 24 field-effect transistors (Toshiba, Tokyo, Japan) that had been set as source followers were used to connect a 24-channel plastic connector that had been cemented to the animal’s head with preamplifiers. The output signals of the preamplifiers, which contained differential operational amplifiers, were transmitted to the main amplifiers in which amplified analog signals were band-pass filtered at 0.5–10 kHz for spiking activities and 0.08–300 Hz for LFP activity.

SPIKE SORTING FOR DATA ANALYSIS
Details of spike sorting were reported previously (Takahashi et al., 2003a,b; Sakurai and Takahashi, 2006). Recorded spike trains were sorted to isolate individual neuronal activities by a method of independent component analysis (ICA) and k-means clustering called ICsort (Takahashi et al., 2003a,b). After spike sorting, the isolation quality was visually inspected in the 1st to 3rd principal components feature spaces.

ANALYSIS OF TASK-RELATED SPIKING ACTIVITY OF NEURONS
Significant difference in the spiking activity of all neurons were confirmed by a statistical test (confidence limit, Abeles, 1982). The procedure of calculating the upper and lower confidence limits (Sakurai et al., 2004) is as the following. Under the null hypothesis that the spiking activity arises at a constant average rate and that this firing rate is independent of the history of the neuron firing and other events (e.g., the task phases and the RP conditions), including the firing of other neurons expected to be
independent Poisson processes, the firing rate in each bin (50 ms) 
\( (n) \) is expected with the average firing rate during whole periods 
of the tasks, the bin size and the number of events. Under these 
assumptions, the probability of finding \( m \) spikes in the bin is given 
by the Poisson formula:

\[
P(m, n) = \frac{e^{-n}n^m}{m!}
\]

Then the lower confidence limit is set at one less than the smallest 
\( m \) for:

\[
\sum_{i=0}^{m} P(i, n) > 0.005
\]

and the upper confidence limit is the smallest \( m \) for

\[
\sum_{i=0}^{m} P(i, n) > 0.995
\]

The spiking activity is defined as significant task-related activity 
when more than two successive bins (100 ms) are above twice 
the band between the upper and lower confidence limits. If a 
near showed statistically significant differences in its activity 
between H- and L-RP conditions and/or between nose-pokes to 
the left and right holes, the neuron was designated as a task-
related one. Then we classified each of the task-related neurons as 
reward-expectation (RE) neurons and/or task-performing (TP) 
nurons. The former were defined as neurons that show a sta-
tistically significant difference in the spiking activity between the 
different RP conditions, and the latter were defined as neurons 
that show significantly different spiking activity between left and 
right nose-pokes on the basis of discriminative cues provided by 
LED illumination.

**ANALYSIS OF LFP**

Coherence and averaged power spectral densities (PSDs) of LFP 
were calculated by multitaper Fourier analysis by applying the 
Chronux toolbox (http://www.chronux.org) (Mitra and Bokil, 
2008; Bokil et al., 2010). To obtain fine time dynamics of the coher-
ence of LFP between the amygdala and the hippocampus, coherograms 
were computed for each LFP pair per session (van der Meer and Redish, 
2011). Coherograms estimate the coherency \( C \) between the two LFP power spectra \( X \) and \( Y \) for each frequency \( f \) as

\[
C_{XY}(f) = \frac{S_{XY}(f)}{\sqrt{S_{XX}(f)S_{YY}(f)}}
\]

with

\[
S_{XY}(f) = \frac{1}{K} \sum_{k=1}^{K} x_k(f) y_k(f),
\]

where \( x_k \) and \( y_k \) are the two LFP spectra as follows:

\[
x_k(f) = \sum_{t=1}^{T} w_t(k)x_t e^{-2\pi ft}.
\]

We used the Chronux cohergram function and mtspectrumc func-
tion with the following parameters: window size, 0.5 \( s \); time step, 
100 ms; five or ten tapers. Before the analysis, all LFP data were 
removed of their direct current offsets, slowly changing compo-
ments, and 50/60 Hz line noise by using the locdetrend function 
and rmlinese function. The frequencies of each LFP were class-
ified into theta band (5–10 Hz), gamma band (30–80 Hz), or high-
frequency band (90–150 Hz) oscillations (Fujisawa and Buzsaki, 
2011; van der Meer and Redish, 2011; Buzsaki and Silva, 2012; 
Tort et al., 2013).

To quantify the amplitude modulation by phases, we calculated the modulation index (MI) in accordance with the pro-
cedure described in Tort et al. (2008). This index can detect cross-
frequency coupling (phase-amplitude coupling; PAC) between 
two frequency ranges of interest (e.g., HFO/gamma and theta). 
To test for significant differences among cross-frequency cou-
plings within and across the regions, we compared the peak MIs 
by three-way ANOVA with the following factors: REs (H-RP 
condition and L-RP condition), task periods (ITI and the trial 
period), and pairs for PAC (PACs within the amygdala, PACs 
within the hippocampus, amygdala theta phase—hippocampal 
HFO amplitude, and hippocampal theta phase—amygdala HFO 
amplitude).

**CORRELATION BETWEEN COHERENCE AND BEHAVIORAL PERFORMANCE**

We calculated the Pearson product-moment correlation coeffi-
cient (Pearson's \( r \)) between LFP coherence in the ITI or the 
trial and the selection time of behavioral performance following 
the ITI. To avoid spurious correlation, we calculated Pearson's \( r \) 
under H- and L-RP conditions respectively. Moreover, to exclude 
correlations that arose by chance (sham correlations), we recal-
culated Pearson's \( r \) using selection times in the next trials (shifted 
trials). We had also calculated the Spearman's rank correlation 
coefficient, but the results were similar to those obtained by 
Pearson's \( r \).

**HISTOLOGY**

After the experiment was conducted, the rats were anesthetized 
depth with an overdose of sodium pentobarbital (120 mg/kg) 
before being perfused and fixed with 10% buffered formalin 
solution. After the brain was sectioned at 50 \( \mu m \) intervals, the 
locations of the electrode tips and tracks in the brains were identi-
fied with the aid of a stereotaxic atlas (Paxinos and Watson, 
2009).

**RESULTS**

**BEHAVIOR**

We used data from 58 sessions to test the behavior of rats 
based on ANOVA. The rats showed different behavioral performance 
under different RP conditions (Figure 2). Under the H-RP 
condition, all rats responded with nose-pokes significantly faster
than under the L-RP condition [Figure 2A; two-way ANOVA, $F_{(1, 5194)} = 593.67, p < 0.001$]. We have confirmed the significantly different selection times between RP conditions for each rat. Also, the rats under the H-RP condition performed the task more accurately than those under the L-RP condition [Figure 2B; ANOVA, $F_{(1, 84)} = 26.8, p < 0.001$]. In the probe test during which no reward was delivered, the rats repeated significantly more trials when the high tone indicating H-RP was presented than when the low tone indicating L-RP was presented [Figure 2C; ANOVA, $F_{(2, 15)} = 48.5, p < 0.001$]. Therefore, the rats actually modulated their behavior according to their expectation of a high probability of reward under the H-RP condition.

**NEURONS FOR REWARD EXPECTANCY AND TASK PERFORMANCE IN AMYGDALA AND HIPPOCAMPUS**

Figure 3 shows averaged and smoothed firing-rate histograms of two examples of amygdala neurons recorded during the task. Both neurons changed their activities in accordance with the different RP conditions. The neuron in the upper portion (Figure 3A) gradually increased its firing rate during the fixation phase. This increase was sustained even after the fixation phase under the H-RP condition. Under the L-RP condition, this neuron decreased its firing rate after the presentation of the discriminative LED cue. Thus, the increase in firing rate was dependent on the rat’s expectation of a high probability of reward. We classified such a neuron as an RE positive neuron. The amygdala neuron in the lower portion (Figure 3B) increased its firing rate under the L-RP condition during the task periods. We classified such a neuron as an RE negative neuron. In these examples of RE neurons, no change was observed in the firing rate for discriminative nose-pokes between the left and right holes.

Among 47 amygdala neurons recorded, 21 neurons were classified as RE neurons. The majority (17, 80.9%) of the RE neurons were RE positive neurons. Most RE neurons (18, 85.7%) also showed no significant difference in their firing rates between discriminative nose-pokes. Three amygdala neurons were classified as TP neurons, which showed differential activity between the discriminative nose-pokes. Two of the TP neurons also showed a significant firing-rate difference between RP conditions and could

![Figure 2](https://example.com/figure2.png)  
**Figure 2** | Behavioral performance in recording sessions. (A) Mean selection times under the H-RP and L-RP conditions in all sessions. Error bars show standard deviations. (B) Mean correct rates under the H-RP and L-RP conditions in all sessions. (C) Mean numbers of trials the rats performed in blocks and IBI in the probe test. Asterisks indicate significant differences of ***$p < 0.005$***.

![Figure 3](https://example.com/figure3.png)  
**Figure 3** | Examples of amygdala RE neurons. Firing rates in all correct trials of a session are cumulated and smoothed separately under the H-RP and L-RP conditions. (A) Example of RE positive neuron, which fired more frequently for the rat under the H-RP condition than the L-RP condition. The time of zero is the time when the rat poked its nose into the selected hole. The period of significantly ($p < 0.05$) higher firing rate under the H-RP condition is indicated by the red horizontal line at the top of the graph. (B) Example of RE negative neuron, which fired more frequently for the rat under the L-RP condition than the H-RP condition. Periods of significantly ($p < 0.05$) higher firing rates under the L-RP condition are indicated by the blue horizontal lines at the top of the graph.
be classified as TP and RE neurons. Therefore, almost all of the
task-related amygdala neurons were RE neurons.

**Figure 4** shows averaged and smoothed firing-rate histograms
of two examples of hippocampal neurons recorded during the
task. Each of the hippocampal neurons was a TP neuron that
showed differential firing rates between discriminative nose-
pokes to the left and right holes. The neuron in the upper portion
(Figure 4A) increased its firing rate after the onset of the right
discriminative cue and drastically increased it further in the selec-
tion period prior to nose-pokes to the right hole. In contrast, its
firing rate gradually decreased after the onset of the left LED cue.
Thus, the activity of this neuron depended on the performance
of the discriminative behavior. The neuron in the lower portion
(Figure 4B) is another example of a TP neuron. It increased its fir-
ing rate after the onset of the left LED cue and increased it further
in the selection period prior to nose-pokes to the left hole. In these
eXamples of TP neurons, firing rates between the RP conditions
did not change.

Among 23 hippocampal neurons recorded, 10 neurons were
classified as TP neurons and one neuron was classified as an RE
neuron. Six of the TP neurons also showed significantly different
firing rates between RP conditions, i.e., they exhibited selective
firing for the cue directions and also changed their firing rates
between the RP conditions. Thus, such TP neurons were also RE
neurons. The one RE, not TP, neuron recorded from the hip-
pocampus significantly increased its firing rate under the L-RP
condition and thus was an RE negative neuron. Consequently,
most of the task-related hippocampal neurons were TP neurons
and the majority of them were also RE neurons that changed their
firing rates in accordance with the differences in RP.

To examine functional dissociation between the amygdala and
the hippocampus, we used Fisher’s exact test for the 2 × 2 contin-
gency table of brain regions and task-related neurons (**Figure 5**).
The results show that the amygdala has a significantly larger
proportion of RE neurons and the hippocampus has a signifi-
cantly larger proportion of TP neurons ($x^2 = 16.6463, df = 1,
p < 0.001$).

**REWARD-EXPECTATION-MODULATED LFP COHERENCE BETWEEN
AMYGDALA AND HIPPOCAMPUS**

**Figure 6** shows an example of LFP coherences between the amyg-
dala and the hippocampus recorded during the task. In the
example, clear differences in the coherences between the different
RP conditions in the upper and lower coherograms were observed
(Figure 6A). We calculated averaged PSDs for the amygdala and
hippocampus data under both RP conditions (**Figure 6B**). The
normalized overall power in the amygdala did not differ between
the RP conditions. Theta and HFOs power in the hippocampus
showed no difference between the RP conditions, and only
the normalized gamma power in the hippocampus under the H-
RP condition was significantly lower than that under the L-RP
condition [ANOVA, $F(1, 50) = 4.226, p < 0.05$]. Therefore, the
difference in LFP coherence between the RP conditions was not
likely to result from an increase in the average PSDs.

As shown in **Figure 6C**, the HFO coherence was significantly
higher under the H-RP condition during most task periods
[ANOVA, $F(1, 6866) = 2428.3, p < 0.001$]. This result indicates
that the elevated HFO coherence of LFP was involved in a high
expectation of reward in the rats, and this involvement was
prominent during the ITI, selection period, and around and
after reward delivery. The gamma coherence was significantly
higher under the H-RP condition than under the L-RP condition.
During most of the task periods \[F_{(1, 6866)} = 84.75, p < 0.001\] (Figure 6D). This result shows that the elevated gamma coherence, similarly to the HFOs, was involved in the high expectation of reward in the rats, although the difference between the RP conditions was not as prominent as between the HFOs.

As shown in Figure 6E, the theta coherence under the H-RP condition significantly decreased in some periods of the task and was drastically ramped up during the period of reward delivery \[F_{(1, 6866)} = 2387.29, p < 0.001\]. This result indicates that, although the modulation of theta coherence under both RP conditions was similar, the theta coherence was somewhat involved in the low expectation of reward under the L-RP condition and prominently involved in getting a highly expected reward under the H-RP condition in the task.

**TRIAL-BY-TRIAL RELATIONSHIP BETWEEN LFP COHERENCE AND BEHAVIORAL PERFORMANCE**

We hypothesized that the activities of the amygdala and hippocampus would show synchrony of oscillation when the RE modulates the task performance. To test this hypothesis, we combined LFP data of all rats and examined the correlation between LFP coherence and behavioral performance, i.e., selection time (Figure 7). The correlation was examined during the ITI and trial periods.

Under the H-RP condition, coherent HFOs had a significant inverse correlation with selection time during the ITI (Figure 7A; \(r = -0.249, p < 0.001\)). During the trial periods, theta coherence showed a significant inverse correlation with selection time (Figure 7A; \(r = -0.200, p < 0.001\)). Such correlation, however, was not observed under the L-RP condition (Figure 7B). There was no correlation between LFP coherence and selection time in shifted trial periods (Figure 7C). These results show that when the rats highly expected a reward under the H-RP condition, LFP coherence had a relationship with the selection time of behavior, and the coherent HFOs during ITIs in particular had a predictive function of the subsequent behavioral performance.

To examine the possibility that the correlation between selection time and theta coherence simply reflected an increase in...
theta power associated with the movement speed during the trial periods, we calculated Pearson’s \( r \) between the theta power during the trial periods and the selection time under both RP conditions. We found that the theta power of the hippocampus showed a significant inverse correlation under both RP conditions (Figure 7D; H-RP condition: \( r = -0.15, p < 0.001 \); L-RP condition: \( r = -0.134, p < 0.001 \)). The theta power of the amygdala also showed a significant inverse correlation under the L-RP condition (Figure 7D; \( r = -0.145, p < 0.001 \)). These results indicated that although the increase in theta power was clearly associated with the movement speed, the association did not correlate with the H-RP condition. Thus, the inverse correlation between task performance and theta coherence under the H-RP condition is not likely to be explained by an increase in theta power.

**CROSS-FREQUENCY COUPLINGS OF LFPs WITHIN AND ACROSS AMYGDALA AND HIPPOCAMPUS**

A recent study (Tort et al., 2013) reported interesting data suggesting that HFO power can be modulated by the theta phase in the hippocampus and neocortex, indicating the contribution of this theta-HFO coupling to cognitive processes such as memory. We hypothesize that such theta-HFO PAC is closely involved in the correlation between LFP coherence and behavioral performance and is important for modulating behavior on the basis of RE. Moreover, Tort et al. (2008) reported that cross-structure coupling between the striatal theta phase and the amplitude of the hippocampal HFOs occurs. We also hypothesize that such coupling interaction between the amygdala and the hippocampus occurs during ITI under the H-RP condition. To test these hypotheses, we examined MI within and across the amygdala and hippocampus during the ITI and trial periods. Three-way ANOVA showed that there was no main effect for all factors: REs (H-RP condition and L-RP condition), task periods (ITI and trial period), pairs for PAC (PACs within the amygdala, PACs within the hippocampus, amygdala theta phase—hippocampal HFO amplitude, and hippocampal theta phase—amygdala HFO amplitude). The interactions between REs and task periods, and among all factors were significant \( F_{(1, 17)} = 9.734, p < 0.01; F_{(3, 51)} = 4.631, p < 0.01 \).

In both the amygdala and the hippocampus, theta-HFO PAC characteristically co-occurred during ITI (Figures 8A,B,E). Within the amygdala, MI in the theta phase and HFO power were significantly higher under the H-RP condition during ITI [Figures 8A,E; \( F_{(1, 136)} = 12.361, p < 0.001 \)]. No such difference in peak MI, however, was observed within the hippocampus (Figures 8B,E). These results suggest that theta-HFO PAC within the amygdala emerges when the rats highly expect a reward, whereas PAC within the hippocampus emerges without a high expectation of reward. Note that the range of frequencies in which PAC occurs was different for different RP conditions in the hippocampus (Figure 8B), although PAC emerged during ITI. Specifically, theta-band oscillations under the H-RP condition modulated a wider range of HFO (>90 Hz), and the modulation tended to occur at low (3–6 Hz) theta frequencies. Interestingly, these ranges are similar to those of the amygdala under the H-RP condition. This result may suggest that the hippocampal PAC is influenced by the amygdala PAC, corresponding to the elevated HFO coherence of LFP (Figure 6C).

As shown in Figures 8C,D, cross-structure coupling did occur. The PAC between the hippocampus theta phase and the amplitude of amygdala HFOs during ITI under the H-RP condition was significantly higher than that under the L-RP condition [Figures 8C; \( F_{(1, 136)} = 16.002, p < 0.001 \)] and the most prominent among the cross-structure couplings (Figure 8E; Ryan’s method, \( t = 2.865, p < 0.05 \)). In contrast, the PACs between the amygdala theta phase and the amplitude of hippocampal HFOs during ITI were not different for different RP conditions (Figures 8D,E). These results suggest that cross-structure couplings between the amygdala and the hippocampus are asymmetric and involved in the correlation between HFO coherence and behavioral performance.

**DISCUSSION**

**HIGH EXPECTATION OF FUTURE REWARD FACILITATES AMYGDALA-HIPPOCAMPU S INTERACTION VIA IMPROVED COHERENT OSCILLATIONS**

In the present study, we concluded that LFP coherence between the amygdala and the hippocampus improved under the H-RP condition when the rats highly expected the next reward (Figure 6). We confirmed that the rats indeed had a high expectation of reward under the H-RP condition by the behavior analysis (Figure 2). The coherence between the amygdala and the hippocampus also had a significant correlation with the subsequent
behavioral performance (Figure 7). These findings are consistent with the notion that interactions between different brain regions are essential for modulating behavior on the basis of RE and that LFP oscillations contribute to such interactions. The finding that coherent HFOs in particular contributed to the regional interactions suggests that the short-time-scale communication between the amygdala and the hippocampus is coordinated with the oscillatory activity during the reward-expectation-induced behavioral modulation.

A notable finding of this study is that the different band coherences between the amygdala and the hippocampus showed different relationships with behavioral performance (Figure 7A).
This finding suggests that different bands of oscillation have different roles in the amygdala-hippocampus interaction behind the adaptive behavior in accordance with reward expectancy. The theta coherence had a direct association with the selection time of behavior during trials. This band of coherence, therefore, may reflect the brain interaction for task performance. Supporting this assumption, some previous studies reported that hippocampal theta oscillation was correlated with the improvement of various task-related behavioral performances such as spatial navigation, memory-based behavior, and running speed (Buzsaki, 2005; Rutishauser et al., 2010; Ahmed and Mehta, 2012). On the other hand, the HFO coherence was predictively associated with the selection time under the H-RP condition, suggesting that this band of coherence may reflect the brain interaction for RE. By switching the different bands of frequency of their oscillatory interaction, these regions might change the type of information they communicate.

**FUNCTIONAL DISSOCIATION BETWEEN THE AMYGDALA AND THE HIPPOCAMPUS**

The types of task-related neurons depend on the brain regions in this study. We found more RE neurons in the amygdala and more task performing (TP) neurons in the hippocampus. These results suggest that the amygdala represents expectation of various reward probabilities and the hippocampus is involved in the appropriate behavioral performance required for the task. This suggestion is also supported by previous findings that the amygdala encodes the predictive representation of future reward (Schoenbaum et al., 1998; Belova et al., 2007; Savage and Ramos, 2009; Roesh et al., 2010), and the hippocampus has important roles in behavioral performance such as spatial navigation (Buzsaki, 2005) and in the improvement of task-related cognition and behavior (Rutishauser et al., 2010; Ahmed and Mehta, 2012).

A question that remains to be answered is what roles the oscillatory interaction between the amygdala and the hippocampus have in reward-expectation-induced behavioral modulation. One possibility is that the interaction serves to bind different information about reward and behavior, which are represented in the amygdala and hippocampus, respectively. Some groups previously studies reported that the interaction of theta waves between these regions contributed to the consolidation and retrieval of fear memory (Seidenbecher et al., 2003; Pape et al., 2005), and the extinction of fear was controlled by theta-rhythmic simultaneous electrical stimulation to these regions (Lesting et al., 2011). In addition to those studies, disconnection between the amygdala and the hippocampus was shown to impair reconsolidation of cocaine-related associative memory (Wells et al., 2011). In this study, the interaction between these regions might associate RP, which resulted in reward expectancy, with behavioral performance. The improved LFP coherence might correspond to the rapid retrieval or recall of such associative memory, hence improving the behavioral performance based on reward expectancy.

Another possibility is that the oscillatory interaction reflects the modulation of the hippocampal function by the amygdala. Firing properties or long-term potentiation of hippocampal cells can be altered by electrical stimulation of the amygdala (Ikegaya et al., 1994; Kim et al., 2012), and by reward information (Kobayashi et al., 1997; Holscher et al., 2003; Tabuchi et al., 2003; Hok et al., 2007; Kennedy and Shapiro, 2009; Singer and Frank, 2009; Dupret et al., 2010). Actually in the present study, many TP neurons in the hippocampus changed their firing rates for different reward probabilities. It is suggested, therefore, that the amygdala influences the firing rates and synchronization of hippocampal neuronal populations through the neuronal oscillations and that the improved LFP coherence corresponds to the strength of this modulation.

**ASYMMETRIC CROSS-STRUCTURE COUPLING BETWEEN THE AMYGDALA AND THE HIPPOCAMPUS**

We found cross-frequency couplings within and across the amygdala and hippocampus during ITI (Figure 8). The PAC within the amygdala varied considerably with the RP conditions, suggesting that cross-frequency couplings of theta-HFO in the amygdala are derived from RE in the rats. In contrast, PAC within the hippocampus emerges without expectation of reward, although the ranges of theta and HFO bands were different for different RP conditions (Figure 8B), and is involved in decision-making or behavioral choice (Tort et al., 2008). These findings suggest that cross-frequency couplings of theta-HFO in the hippocampus reflect task-performing-related cognitive processes. Therefore, the view of the functional dissociation between the amygdala and the hippocampus is also supported by the observations of cross-frequency coupling within these regions.

Moreover, the PAC between the hippocampus theta phase and the amplitude of amygdala HFOs during ITI under the H-RP condition was very distinct (Figure 8C). This asymmetric cross-structure coupling between the amygdala and the hippocampus might explain the elevated HFO coherence and the changes in the ranges of the hippocampal theta and HFO frequencies. First, the PAC within the amygdala under high expectation of reward allows the amygdala HFO to be modulated by the hippocampal theta phase. Herewith, the amygdala HFO influences the hippocampal HFO. Then, the range of the hippocampal HFO band changes and the PAC within the hippocampus becomes similar to that within the amygdala. From this viewpoint, the elevated HFO coherence can reflect the process that the activities of the amygdala affect those of the hippocampus, and our observation of the oscillatory interaction between the amygdala and the hippocampus might reflect the information flow during the behavioral modulation based on RE. Future works are needed to confirm this hypothesis. Consequently, we conclude that the activities of the amygdala and hippocampus are synchronized in oscillation, which is one of the mechanisms of how RE modulates goal-directed behavior.

**FUTURE PROSPECTS**

Although we focused on the amygdala and hippocampus in this study, other brain regions, in particular, the prefrontal cortex (PFC), and the striatum, should be considered to play some important roles in behavioral modulation on the basis of RE (Corbit et al., 2001; Ostlund and Balleine, 2007; Holmes et al., 2010). The interaction between PFC, especially the orbitofrontal cortex, and the amygdala is crucial for generating and using the predictive representation of future reward (Schoenbaum et al.,
2003; Holland and Gallagher, 2004; Schoenbaum and Roesch, 2005). In addition, the prelimbic area of the medial PFC might be involved in generating representations of reward in connection with the hippocampus (Hok et al., 2005; Burton et al., 2009), and the oscillation between PFC and the hippocampus is related to the regulation of goal-directed memory and behavior (Jones and Wilson, 2005; Benchenane et al., 2010). The striatum is projected from the amygdala and hippocampus and is reported to convert motivational signals to motor signals (Holmes et al., 2010; Corbit, L. H., Muir, J., and Balleine, B. W. (2001). The role of the nucleus accumbens and landmark navigation, episodic and semantic memory.

Buzsaki, G., and Silva, F. L. (2012). High frequency oscillations in the intact brain. Prog. Neurobiol. 98, 241–249. doi: 10.1016/j.pneurobio.2012.02.004

Corbit, L. H., Muir, J., and Balleine, B. (2001). The role of the nucleus accumbens in instrumental conditioning: evidence of a functional dissociation between accumbens core and shell. J. Neurosci. 21, 3251–3260.

Dupret, D., O’Neill, J., Pleydell-Bouverie, B., and Caiccsvari, J. (2010). The reorganization and reactivation of hippocampal maps predict spatial memory performance. Nat. Neurosci. 13, 995–1004. doi: 10.1038/nn.2599

Fujiwara, S., and Buzsaki, G. (2011). A 4 Hz oscillation adaptively synchronizes prefrontal, VTA, and hippocampal activities. Neuron 72, 153–165. doi: 10.1016/j.neuron.2011.08.018

Hok, V., Lenck-Santini, P.-P., Roux, S., Muller, R. U., and Pouzet, B. (2007). Goal-related activity in hippocampal place cells. J. Neurosci. 27, 472–482. doi: 10.1523/JNEUROSCL.2864-06.2007

Hok, V., Save, E., Lenck-Santini, P. P., and Pouzet, B. (2005). Coding for spatial goals in the prelimbic/infralimbic area of the rat frontal cortex. Proc. Natl. Acad. Sci. U.S.A. 102, 4602–4607. doi: 10.1073/pnas.0407332102

Holland, P. C., and Gallagher, M. (2004). Amygdala-frontal interactions and reward expectancy. Curr. Opin. Neurobiol. 14, 148–155. doi: 10.1016/j.conb.2004.03.007

Holmes, N. M., Marchand, A. R., and Coutureau, E. (2010). Pavlovian to instrumental transfer: a neurobehavioural perspective. Neurosci. Biobehav. Rev. 34, 1277–1295. doi: 10.1016/j.neubiorev.2010.03.007

Holscher, C., Jacob, W., and Mallot, H. A. (2003). Reward modulates neuronal activity in the hippocampus of the rat. Behav. Brain Res. 142, 181–191. doi: 10.1016/S0166-4328(02)00422-9

Ikegaya, Y., Saito, H., and Abe, K. (1994). Attenuated hippocampal long-term potentiation in basolateral amygdala-lesioned rats. Brain Res. 656, 157–164. doi: 10.1016/0006-8993(94)91377-3

Ito, R., Robbins, T. W., Pennartz, C. M., and Everitt, B. (2008). Functional interaction between the hippocampus and nucleus accumbens shell is necessary for the acquisition of appetitive spatial context conditioning. J. Neurosci. 28, 6950–6959. doi: 10.1523/JNEUROSCL.1615-08.2008

Johnson, A. W., Gallagher, M., and Holland, P. (2009). The basolateral amygdala is critical to the expression of pavlovian and instrumental outcome-specific reward devaluation effects. J. Neurosci. 29, 696–704. doi: 10.1523/JNEUROSCL.3758-08.2009

Jones, M. W., and Wilson, M. A. (2005). Theta rhythms coordinate hippocampal-prefrontal interactions in a spatial memory task. PLoS Biol. 3(4):e402. doi: 10.1371/journal.pbio.0030402

Kennedy, P. J., and Shapiro, M. L. (2009). Motivational states activate distinct hippocampal representations to guide goal-directed behaviors. Proc. Natl. Acad. Sci. U.S.A. 106, 10805–10810. doi: 10.1073/pnas.0903259106

Kim, E. J., Kim, E. S., Park, M., Cho, J., and Kim, J. (2012). Amygdala stimulation produces alterations on firing properties of hippocampal place cells. J. Neurosci. 32, 11424–11434. doi: 10.1523/JNEUROSCL.1108-12.2012

Kobayashi, T., Nishifo, H., Fukuda, M., Bures, J., and Ono, T. (1997). Task-dependent representations in rat hippocampal place neurons. J. Neurophysiol. 78, 597–613.

Lesting, J., Narayanan, R. T., Kluge, C., Sangha, S., Seidenbecher, T., Pape, H. C. (2011). Patterns of Coupled theta activity in amygdala-hippocampal-prefrontal cortical circuits during fear extinction. PLoS ONE 6:e21714. doi: 10.1371/journal.pone.0021714

McNaughton, B. L., Barnes, C. A., Meltzer, J., and Sutlward, R. J. (1989). Hippocampal granule cells are necessary for normal spatial learning but not for spatially-selective pyramidal cell discharge. Exp. Brain Res. 76, 485–496. doi: 10.1007/BF01248904

Mitra, P., and Bokil, H. (2008). Observed Brain Dynamics. New York, NY: Oxford UP.

Morrison, S. E., and Salzman, C. D. (2010). Re-valuing the amygdala. Curr. Opin. Neurobiol. 20, 221–230. doi: 10.1016/j.conb.2010.02.007

Ostlund, S. B., and Balleine, B. W. (2007). Orbitofrontal cortex mediatesoutcome in pavlovian but not instrumental conditioning. J. Neurosci. 27, 4819–4825. doi: 10.1523/JNEUROSCL.5443-06.2007

Pape, H. C., Narayanan, R. T., Smid, J., Stork, O., and Seidenbecher, T. (2005). Theta activity in neurons and networks of the amygdala related to long-term fear memory. Hippocampus 15, 874–880. doi: 10.1002/hipo.20120

Paxinos, G., and Watson, C. (2009). The rat brain, compact sixth edition. New York, NY: Academic Press.

Popescu, A. T., Popa, D., and Pare, D. (2009). Coherent gamma oscillations couple the amygdala and striatum during learning. Nat. Neurosci. 12, 801–808. doi: 10.1038/nn.2305

Ramirez, D. R., and Savage, L. M. (2007). Differential involvement of the basolateral amygdala, orbitofrontal cortex, and nucleus accumbens core in the...
acquisition and use of reward expectancies. Behav. Neurosci. 121, 896–906. doi: 10.1037/0735-7044.121.5.896

Retailleau, A., Etienne, S. E., Guthrie, M. G., and Boraud, T. (2012). Where is my reward and how do I get it. Interaction between the hippocampus and the basal ganglia during spatial learning. J. Physiol. Paris 106, 72–80. doi: 10.1016/j.jphysparis.2011.10.002

Roesh, M. R., Calu, D. J., Esber, G. R., and Schoenbaum, G. (2010). Neuronal correlates of variations in event processing during learning in basolateral amygdala. J. Neurosci. 30, 2464–2471. doi: 10.1523/JNEUROSCI.5781-09.2010

Rutishauser, U., Ross, I. B., Mamalak, A. N., and Schuman, E. M. (2010). Human memory strength is predicted by theta-frequency phase-locking of single neurons. Nature 464, 903–907. doi: 10.1038/nature08860

Sakurai, Y. (1993). Dependence of functional synaptic connections of hippocampal and neocortical neurons on types of memory. Neurosci. Lett. 158, 181–184. doi: 10.1016/0304-3940(93)90255-N

Sakurai, Y. (1994). Involvement of auditory cortical and hippocampal neurons in singing. J. Neurosci. 13, 121–134. doi: 10.1016/j.neuron.2009.11.016

10.1073/pnas.0810524105

10.1038/neuro.04.009.2009

10.1080/07357044.2013.821404

Conflict of Interest Statement: The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

Received: 14 September 2013; accepted: 07 November 2013; published online: 03 December 2013.

Citation: Terada, S., Takahashi S and Sakurai Y (2013) Oscillatory interaction between amygdala and hippocampus coordinates behavioral modulation based on reward expectation. Front. Behav. Neurosci. 7:177. doi: 10.3389/fnnbeh.2013.00177

This article was submitted to the journal Frontiers in Behavioral Neuroscience.

Copyright © 2013 Terada, Takahashi and Sakurai. This is an open-access article distributed under the terms of the Creative Commons Attribution License (CC BY). The use, distribution or reproduction in other forums is permitted, provided the original author(s) or licensor are credited and that the original publication in this journal is cited, in accordance with accepted academic practice. No use, distribution or reproduction is permitted which does not comply with these terms.