

Original Research Article

Locus cœruleus noradrenergic neurons phase-lock to prefrontal and hippocampal infra-slow rhythms that synchronize to behavioral events

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Abstract

The locus cœruleus (LC) is the primary source of noradrenergic projections to the forebrain, and, in prefrontal cortex, is implicated in decision-making and executive function. LC neurons phase-lock to cortical infra-slow wave oscillations during sleep. Such infra-slow rhythms are rarely reported in awake states, despite their interest, since they correspond to the time scale of behavior. Thus, we investigated LC neuronal synchrony with infra-slow rhythms in awake rats performing an attentional set-shifting task. Local field potential (LFP) oscillation cycles in prefrontal cortex and hippocampus on the order of 0.4 Hz phase-locked to task events at crucial maze locations. Indeed, successive cycles of the infra-slow rhythms showed different wavelengths, as if they are periodic oscillations that can reset phase relative to salient events.

Simultaneously recorded infra-slow rhythms in prefrontal cortex and hippocampus could show different cycle durations as well suggesting independent control. Most LC neurons (including optogenetically identified noradrenergic neurons) recorded here were phase-locked to these infra-slow rhythms, as were hippocampal and prefrontal units recorded on the LFP probes. The infra-slow oscillations also phase-modulated gamma amplitude, linking these rhythms at the time scale of behavior to those coordinating neuronal synchrony. This would provide a potential mechanism where noradrenaline, released by LC neurons in concert with the infra-slow rhythm, would facilitate synchronization or reset of these brain networks, underlying behavioral adaptation.

1 Introduction

1 The brain coordinates activity among interconnected regions via coherent oscillatory cycles of
2 excitation and inhibition (Womelsdorf, et al., 2007). This can facilitate communication among
3 selected subsets of neurons, groups of neurons, and brain regions. Sensory stimuli or behavioral
4 events can reset the phase of these oscillations (Canovier, 2016; Voloh and Womelsdorf, 2016),
5 linking activity of multiple neurons to process information in concert. However, the principal
6 brain rhythms studied in behaving animals are at the time scale of cell neurophysiological
7 processes, which are much faster (on the order of tens and hundreds of milliseconds) than real life
8 behavioral events, which typically occur at second and supra-second time scales. The brain has
9 several mechanisms linking these two time scales, some of which involve the hippocampus
10 (reviewed in Banquet, et al., 2021) and associated networks, including the prefrontal cortex and
11 striatum.

12 Little is known about brain rhythms that operate in this crucial behavioral time scale during
13 awake behavior. The brain is indeed capable of generating rhythms on the order of 0.1-1.0 Hz,
14 although these have been principally characterized during sleep (Steriade, 1993). Furthermore,
15 during sleep or under anesthesia, rat noradrenergic locus cœruleus (LC) and prefrontal cortical
16 (Pfc) neurons are phase-locked to infra-slow rhythms (Lestienne, et al., 1997; Eschenko, et al.,
17 2012; Totah, et al., 2018). LC stimulation exerts powerful influence on neurophysiological
18 activity in Pfc and hippocampus (Hip; Berridge and Foote, 1991). LC actions in prefrontal cortex
19 are implicated in vigilance, decision-making, and executive function, while in Hip they are
20 associated with learning and processing contextual information (e.g., Wagatsuma, et al., 2018;

21 Sara, 2009 for review). Since oscillations can coordinate activity in brain networks, we reasoned
22 that there might also be rhythmicity on this behavioral time scale in awake animals, and
23 investigated this possibility in rats performing a task engaging Pfc, Hip and LC (Oberto, et al.,
24 2022; Xiang, et al., 2019). Such coordinated activity could provide a possible link between
25 neuromodulation and oscillatory coordination of brain areas on the time scale of behavior.

26 **2 Materials and Methods**

27 All experiments were carried out in accordance with local (Comité d'éthique en matière
28 d'expérimentation animale no. 59), institutional (Scientific Committee of the animal facilities of
29 the Collège de France) and international (US National Institutes of Health guidelines; Declaration
30 of Helsinki) standards, legal regulations (Certificat no. B751756), and European/national
31 requirements (European Directive 2010/63/EU; French Ministère de l'Enseignement Supérieur et
32 de la Recherche 2016061613071167) regarding the use and care of animals. The data analyzed
33 here were recorded in experiments described by Xiang, et al. (2019) and further details can be
34 found there.

35 **2.1 Animals**

36 Nine male Long-Evans rats (Janvier Labs, Le Genest-Saint Isle France; weight, 280–400 g) were
37 maintained on a 12 h:12 h light-dark cycle (lights on at 7 A.M.). The rats were handled on each
38 workday. To motivate animals for behavioral training on the T maze, water was partially
39 restricted except for a 10–30 min period daily to maintain body weight at 85% of normal values
40 according to age. Rats were rehydrated during weekends. Food was restricted to 14 g of rat chow
41 daily (the normal daily requirement) to prevent the animals from becoming overweight.
42 Recording properties and behavioral correlates of the LC neurons of four of the rats were
43 reported in Xiang, et al. (2019), but no LFP data are presented there. Recordings of LC neurons
44 during task performance was not possible in the remaining five rats included here.

45 **2.2 The automated T maze with return arms**

46 The behavioral task took place in an elevated automated T-maze (see Fig. 3A) consisting of a
47 start area, a central arm, two reward arms and two return arms, which connected the reward arms

48 to the start area. Small wells at the end of each reward arm delivered liquid reward (30 μ l of
49 0.25% saccharin solution in water) via solenoid valves controlled by a CED Power1401 system
50 (Cambridge Electronic Design, Cambridge, UK) with a custom-written script. As the rats crossed
51 a central photo-detector, visual cues (VCs) were displayed in pseudo-random sequence on video
52 monitors positioned behind, and parallel to the two reward arms. This is the “VC”, or “central
53 arm PD” event. The VCs were either lit or dim uniform fields. The rat then selected the left or
54 right arm and crossed the Reward arm (Rew) photodetector (PD), triggering reward release from
55 an audible solenoid valve. Crossing the photodetector in the middle of the return arm of the T-
56 maze (Return arm PD; Ret) triggered the visual cue to be turned off. Photodetectors detected task
57 events and triggered cues and rewards via the CED Spike2 script.

58 **2.3 Viral vector preparation, injection and immunohistochemistry**

59 The Canine Adenoviral vector (CAV2-PRS-ChR2-mCherry) was produced at the University of
60 Bristol. Details about it and injections procedures appear in Xiang, et al. (2019). Also see Xiang,
61 et al. (2019) for histological and immunohistochemical procedures and neuron characterization.

62 **2.4 Electrode and optrode implants**

63 Following VC task pre-training, at least one day before surgery, rats were returned to ad libitum
64 water and food. Surgical procedures and electrode construction are the same as in Xiang, et al.
65 (2019). Moveable tungsten microelectrodes (insulated with epoxylite®, impedance = 2-4 M Ω ,
66 FHC Inc, USA) were used for LC recordings. A single microelectrode, or two or three such
67 electrodes glued together was implanted at AP -3.8-4 mm relative to lambda, and ML 1.1-1.2
68 mm, with a 15° rostral tilt. A stainless steel wire (Teflon coated, diameter=178 μ m, A-M
69 systems Inc) implanted in the midbrain area about 1-2 mm anterior to the LC electrode tip served
70 as a fixed LC reference electrode, permitting differential recording. The rat with the virus
71 injection (R328) was implanted with an optrode composed of a tungsten microelectrode
72 (insulated with epoxylite, impedance = 2-4 M Ω , FHC Inc, USA) glued to a 200 μ m optic fiber
73 implant with a ferrule (0.37 numerical aperture, hard polymer clad, silica core, multimode,
74 Thorlabs), with tip distances 1 mm apart (the electrode was deeper). The optic fiber implant and
75 optic fiber cables were constructed at the NeuroFabLab (CPN, Ste. Anne Hospital, Paris). Two
76 screws (diameter = 1 mm, Phymep, Paris) with wire leads were placed in the skull above the

77 cerebellum to serve as ground. LC electrodes were progressively lowered under
78 electrophysiological control until characteristic LC spikes were identified (located ~ 5-6 mm
79 below the cerebellar surface, see Xiang, et al., 2019 for details). For the virus-injected rat, LC
80 spikes could also be identified by responses to laser stimulations (described below). Following
81 implantation, the microelectrode was fixed to a micro-drive allowing for adjustments along the
82 dorsal-ventral axis. The headstage was fixed to the skull with dental cement, and surrounded by
83 wire mesh stabilized with dental cement for protection and shielding. After the surgery, animals
84 were returned to their home cages for at least one-week recovery with ad libitum water and food
85 and regular observation.

86 **2.5 Electrophysiological recordings**

87 Rats were then returned to dietary restriction. The movable electrodes were gradually advanced
88 until a well-discriminated LC unit was encountered and then all channels were recorded
89 simultaneously while the rat performed in the T-maze. If no cells could be discriminated, the
90 electrodes were advanced and there was at least a 2 h delay before the next recording session.

91 For daily online monitoring of LC spikes, pre-amplified signals were filtered between 300-3000
92 Hz for verification on the computer screen (Lynx-8, Neuralynx, Bozeman, MT, USA) and also
93 transmitted to an audio monitor (audio analyzer, FHC). For recordings, brain signals were pre-
94 amplified at unity gain (Preamp32, Noted Bt, Pecs, Hungary) and then led through a flexible
95 cable to amplifiers (x500, Lynx-8, Neuralynx) and filters (0.1-9 kHz, Lynx-8, Neuralynx). Brain
96 signals were digitized at ~20 kHz using CED Power1401 converter and Spike2 data acquisition
97 software. The LC unit activity was identified as in Xiang, et al. (2019) by: 1) spike waveform
98 durations ≥ 0.6 ms; 2) low average firing rate (1-2 Hz) during quiet immobility; 3) brief
99 responses to unexpected acoustic stimuli followed by prolonged (around 1 s) inhibition; 4) for the
100 virus-injected rat (R328), LC units were verified by responses to laser stimulation. A laser driver
101 (Laserglow Technologies, Canada, wavelength 473 nm) was controlled by signals from a
102 stimulator (Grass Technologies, USA, Model SD9). Light intensity from the tip of optic fiber was
103 measured by a power meter (Thorlabs, Germany, Model PM100D). If unit firing was entrained to
104 the pulses with an increased rate (to at least twice the baseline firing rate) averaged over all the
105 stimulations, they were considered to be noradrenergic LC units.

106 A light emitting diode (LED) was mounted on the cable that was plugged into the headstage. This
107 was detected by a video camera mounted above the T-maze and transmitted to the data
108 acquisition system at a sampling rate of ~30 Hz for the purpose of position tracking.

109 **2.6 Signal processing, spike sorting and data analyses**

110 For off-line spike detection of LC activity in three of the rats, the wide-band signals were
111 converted and digitally high-pass filtered (nonlinear median-based filter). Waveforms with
112 amplitudes passing a threshold were extracted, and then subjected to principal component
113 analysis (PCA). All of these processes were performed with NDManager (Hazan, et al., 2006).
114 Spikes were sorted with a semi-automatic cluster cutting procedure combining KlustaKwik (KD
115 Harris, <http://klustakwik.sourceforge.net>) and Klusters (Hazan, et al., 2006). Spikes with
116 durations less than 0.6 ms were rejected. In one rat (R311), the LC signal was filtered from 300-
117 3000 Hz during recording, and the spike sorting was performed with Spike2 software (which
118 employs a waveform template-matching algorithm). Most data analyses were performed using
119 Matlab (R2010a) with the statistical toolbox FMAToolbox (developed by M. Zugaro,
120 <http://fmatoolbox.sourceforge.net>) and scripts developed in the laboratory as well as some
121 statistical analyses performed with Microsoft© Excel©. Phase was computed with the 'Phase'
122 function of the FMAToolbox, employing the Hilbert transform. To characterize periods with
123 infra-slow rhythms, a criterion for salient phase-locking to task events was established as when
124 the SEM range of LFP phase was less than $0.75 * \pi$ radians (cf., Figure 4, middle column). This is
125 termed "regular phase-locking". Sessions tallied for phase-locking of LC neurons to infra-slow
126 rhythms were included only if they had at least 1000 LC neuron spikes.

127 **3 Results**

128 **3.1 LC neuron phase-locking to prefrontal and hippocampal infra-slow rhythms**

129 Infra-slow rhythms were readily apparent in visual inspections of hippocampal (Hip) and
130 prefrontal cortical (Pfc) local field potentials (LFPs) (Fig. 1A). These were rendered more salient
131 by filtering the signal in a 0.1-1.0 Hz window (Fig. 1B). We applied an amplitude threshold to
132 examine data from those periods when the infra-slow rhythm amplitude was elevated (Fig. 1C).
133 We applied an amplitude threshold to examine data from those periods when the infra-slow

134 rhythm amplitude was elevated (Fig. 1C). This was intended to limit analyses to those periods
135 when the infra-slow oscillation was sufficiently robust, thus avoiding possibly spurious
136 computations of phase angle from low amplitude oscillations. In each of five rats, LC neurons
137 were phase-locked to Pfc (e.g., Figs. 1D and 2), as well as Hip infra-slow LFP rhythms. The
138 incidence of phase-locking of the LC neurons in sessions with regular phase-locking of the infra-
139 slow rhythms to task events was 20 out 23 for Hip LFP and 15/20 for Pfc LFP (Rayleigh test,
140 $p<0.05$; for histology, see Fig. 2 of Xiang, et al., 2019). In the animal where noradrenergic LC
141 neurons were identified optogenetically (see Methods), all were phase-locked to the infra-slow
142 rhythms ($n=8$ for both Pfc and Hip; Rayleigh test, $p<0.05$).

143 The modal preferred infra-slow phase among these neurons was $0.35*\pi$ radians for Hip infra-
144 slow and $0.15*\pi$ radians for Pfc infra-slow ($p<0.05$, Rayleigh test; not shown).

145 **3.2 Prefrontal and hippocampal infra-slow rhythms are synchronized to maze events**

146 The infra-slow rhythms were phase-locked to task-relevant positions on the maze (Fig. 3B; Supp.
147 Fig. 1B). To quantify this phase-locking, the mean (\pm SEM) phase of the rhythm was plotted in
148 peri-event time color plots (see Fig. 4 and Supp. Fig. 1A for examples) over all trials in 57
149 sessions from eight rats (including the five with LC recordings). Infra-slow rhythms were phase-
150 locked to the reward arm photodetector crossing (Rwd) in 51 of the recording sessions for Hip,
151 and 46 sessions for Pfc (see Table 1). The other maze events had fewer incidences of regular
152 phase-locking (Pfc return arm photodetector crossing, or Rtn: 11; Pfc central arm visual cue onset
153 PD, or VC: 18; Hip Rtn: 18; Hip VC: 20). The mean phases at the respective PD crossings (in
154 those cases when $SEM \leq 0.75*\pi$ radians there) were $0.70*\pi$ and $0.24*\pi$ radians for Pfc and Hip
155 Rtn, $0.25*\pi$ and $0.22*\pi$ radians for Pfc and Hip Rwd, and $0.19*\pi$ and $0.01*\pi$ radians for Pfc and
156 Hip VC. The root-mean-square differences between Pfc and Hip mean phase (calculated pairwise
157 by session) at the respective PD crossings were $0.14*\pi$, $0.13*\pi$ and $0.12*\pi$ rad. The regular
158 phase-locking could last from less than one to over 2.5 successive rhythmic cycles (Supp. Figs. 1,
159 and 2, Table 1) and could continue from one event to the next (Fig. 4, Supp. Fig. 1). For PL Rwd
160 and Hip Rwd, 30 and 37 sessions had durations of regular phase-locking lasting one or more
161 cycles, respectively. These permitted quantification of the temporal duration of the cycles, which
162 ranged from 2.0 to 2.6 s, the equivalent of 0.4 to 0.5 Hz. In the six cases of Rwd PD phase-

163 locking which had a second complete cycle, the mean of the first was 2.3 s, while the second was
164 lower, 2.0 s (pairwise t-test, $p=0.0009$, $df=5$). Thus, these are not regular periodic oscillations,
165 but, rather, this is consistent with phase-locking to task events. Pfc and Hip infra-slow rhythms
166 sometimes resembled one another (e.g., Fig. 2). To compare them, sessions were classified as
167 having Pfc and Hip regular phase-locking in the following ranges of cycles (see Table 1). In 17 of
168 the 57 sessions, these numbers of cycles were different between Pfc and Hip for VC, Rwd and/or
169 Rtn (e.g., Supp. Fig. 2). This indicates that it is unlikely that Pfc and Hip infra-slow rhythms are
170 related by volume conduction, and suggests that they could be independently generated.

171 The infra-slow rhythms were regularly phase-locked to two (in 24 sessions), or even all three (in
172 8 sessions) different task events. Thus, they were not linked to any specific task-related behavior.
173 To test whether infra-slow rhythms were triggered by rapid head movements, regression analysis
174 compared the onset of regular phase-locking and times of peak acceleration, or deceleration
175 around the Rwd PD crossings, and were not significant ($r^2=0.034$, $p=0.49$ and $r^2=0.0056$, $p=0.80$
176 respectively; $df=15$; see Supp. Fig. 3). In Xiang et al. (2019), we showed that LC neurons fire
177 more during accelerations. Indeed, the periods with the greatest increase in LC activity were not
178 those most frequent for the start of regular phase-locking (i.e., reset) of the infra-slow rhythm;
179 rather phase-locking occurred most frequently to Rwd PD crossing (see above), where no
180 consistent accelerations occurred (see Supp. Figs. 3 and 4). These results indicate it is unlikely
181 that Pfc and Hip infra-slow rhythms are due to a biomechanical artifact, for example, from
182 locomotion or head rocking.

183 **3.3 Coordination of neuronal activity across time scales**

184 In the four sessions where Hip and Pfc neurons could be discriminated from the LFP electrodes,
185 most were also modulated by infra-slow rhythms (Pfc LFP modulated 6/12 Pfc units and 8/11
186 Hip units; Hip LFP modulated 8/12 Pfc units and 8/11 Hip units; Rayleigh test $p<0.05$). The LC
187 neurons could have relatively consistent phases with respect to the two infra-slow rhythms (not
188 shown). LC neurons could be phase-locked to oscillations in the delta frequency range (1-4 Hz)
189 in Pfc ($n=15/37$) and Hip (11/37) as well as theta (5-10 Hz; 7/37 and 5/37 respectively) for Pfc
190 and Hip (see Supp. Fig. 5). While phase-locking of LC neurons to gamma (40-80 Hz) was rare

191 (n=2 for both structures' LFPs), the infra-slow rhythm did modulate the amplitude of gamma
192 oscillations at 35-45 Hz in hippocampus and prefrontal cortex (Fig. 5).

193 **4 Discussion**

194 LFP oscillation cycles on the order of 0.4 Hz in prefrontal cortex and hippocampus were phase-
195 locked to task events at crucial points on the maze. Successive cycles had different cycle lengths,
196 indicating that, if they are indeed periodic oscillations, their phase can reset to salient events.
197 Simultaneous recordings in prefrontal cortex and hippocampus could have different cycle lengths
198 as well, while still phase-locking to task events. This would seem to exclude any single structure
199 from entraining these independent rhythms simultaneously. This also would rule out a
200 contribution of volume conduction. The intriguing issue of the origin of these rhythms merits
201 further investigation. Most of the LC neurons were phase-locked to these infra-slow prefrontal
202 cortical and hippocampal LFPs, including all of the optogenetically identified noradrenergic
203 neurons. Hippocampal and prefrontal units were also phase-locked to the infra-slow oscillations.
204 While the number of LC neurons recorded may appear low, this is typical for the rare chronic
205 recording studies of this structure in behaving rats, likely because its diminutive dimensions and
206 deep location render accurate electrode placements challenging.

207 This is consistent with previous work showing neuronal activity adapting to the time scale of
208 behavioral events. For example, in behavioral tasks with delays, several brain structures show
209 "time cell" activity: neurons with sequential "tiling" activity lasting on the order of several
210 seconds. These periods can expand or contract depending upon the duration of task-imposed
211 intervals (MacDonald, et al., 2011). We speculate that this infra-slow rhythm may originate in the
212 hippocampal-prefrontal system since neuro-physiological activity there tracks time intervals on
213 the order of several seconds based upon regularities in temporal structure of behavioral or
214 environmental events.

215 Steriade, et al. (1993) observed infra-slow (0.3-1.0 Hz) rhythms in neocortical activity in
216 anesthetized and naturally sleeping cats. Eschenko et al (2012) showed that LC neuronal activity
217 in sleeping rats is synchronized with the sleep infra-slow wave cycle (1 Hz) and is out of phase
218 with Pfc neuronal activity. Similarly, in rats under ketamine anesthesia, there is a negative
219 correlation between activity of LC NE neurons and prefrontal neurons, when neuron activation

220 oscillates at ~1 Hz (Sara and Hervé-Minvielle 1995; Lestienne, et al. 1997). Furthermore, when
221 the latter authors pooled their LC recording data, they were significantly phase-locked to cortical
222 LFP delta oscillations. While these infra-slow cycles of UP-DOWN state transitions are not
223 generally observed in awake animals, this does demonstrate that LC can fire rhythmically, and
224 that these structures can coordinate their activity at this time scale. Furthermore, in rats under
225 urethane anesthesia, Totah, et al. (2018) found that the firing rate of locus coeruleus neurons
226 oscillates at 0.4-0.5 Hz. And, in head-fixed awake mice, cortical noradrenergic axons exhibited
227 rhythmic Ca^{2+} activity at 0.5–0.6 Hz (Oe, et al., 2020). Thus, the LC could also be associated
228 with the Pfc-Hip in the origin, maintenance and communication of behaviorally relevant infra-
229 slow rhythms in the brain. Further work is required to elucidate the respective roles of these
230 structures in these processes.

231 In the awake state, there is evidence for infra-slow neural processing although this was not
232 observed as rhythms per se. Molter, et al. (2012) observed a 0.7 Hz modulation of the power of
233 theta rhythm recorded in rat Hip. This 0.7 Hz modulated Hip neuronal activity during sleep, as
234 well as during behavior in a maze, a running wheel, and an open field. Positions on a figure-8
235 maze corresponded to specific phases of this modulatory rhythm, similar to the infra-slow rhythm
236 recorded here. (Their filter settings excluded 0.7 Hz rhythms and thus this could not be directly
237 measured in that work.) In Molter et al. (2012), the 0.7 Hz modulation of the power of the theta
238 infra-slow modulation was locked at π radians to junction points in the maze (their Figure 7B),
239 where accelerations might be expected. However, they found no overall correlation between
240 phase and acceleration. Halgren et al. (2018) observed a rhythm at less than 3 Hz generated in the
241 superficial layers of the cerebral cortex in awake humans. The phase of this rhythm reset to
242 infrequent tones in their oddball task, similar to the reset of the infra-slow rhythm here in relation
243 to salient task events.

244 Villette, et al. (2015) used calcium imaging to observe CA1 pyramidal cells in head fixed mice
245 moving in the dark on a non-motorized treadmill. They found that different neurons fired
246 sequentially in cycles at the same time scale as the infra-slow oscillations observed here.
247 Furthermore, the cycles could occur singly, or consecutively in groups of two or three. The
248 authors interpreted this as representing an intrinsic metric for representing distance walked. This
249 resembles time cell activity (Pastalkova et al., 2008; MacDonald et al., 2011) evoked above,

250 where the length of the cycle extends to the time scale of the ongoing task (Kraus, et al., 2013;
251 Ravassard, et al., 2013). The 2 to 5 s durations of the cycles in the Villette, et al. (2015) study
252 may represent a default value since their task had no temporal structure. This is on the order of
253 the time scale of the infra-slow rhythm recorded here, and the variable numbers of cycles they
254 observed might flexibly adapt to the positions of task-relevant events to lead to the results found
255 here.

256 The present observations of phase-locking of LC neurons to infra-slow rhythms in hippocampus
257 could ostensibly be due to independent synchrony of the infra-slow rhythms and the LC neurons
258 to task events. However, the LC neurons showed phase preferences in the infra-slow rhythms in
259 data pooled over multiple task events. We did not observe any simple relation between infra-slow
260 rhythms and motor events (e.g., as we showed for LC neurons with acceleration or deceleration
261 by Xiang, et al., 2019), since regular phase-locking could start before (Supp. Fig. 1) or after the
262 same task events in different sessions (not shown), and continue over periods including a variety
263 of behaviors.

264 The phase-locking of LC neurons to infra-slow rhythms in Hip and Pfc, as well as to oscillations
265 in the delta, theta and gamma frequency bands could reveal coordinated neuronal processing
266 within a unified temporal framework (cf., Totah, et al., 2018a). The scale of this corresponded to
267 the temporal and spatial regularities characterizing the current behavioral patterns. Cross-
268 frequency coupling could serve as a mechanism to link processing at different time scales. This
269 could facilitate both ‘Communication through coherence’ (CTC, Bosman et al., 2012; Fries,
270 2005) and ‘Binding by synchrony’ (Eckhorn, et al., 1990; Engel, et al., 1999; Buehlmann and
271 Deco, 2010). Thus, infra-slow rhythms would serve as a scaffold to link the time scales of
272 dynamics of neuronal processes to those of behavior and cognitive processes. Noradrenaline,
273 released by LC neurons in concert with the infra-slow rhythm, would participate in synchronizing
274 or resetting those brain networks underlying behavioral adaptation to these events (Bouret and
275 Sara, 2005; Sara and Bouret, 2012).

276 **5 Competing interests**

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

279 **6 Author contributions**

280 S.I.W. and L.X. designed the experiments; S.J.S. and L.X. developed and implemented the LC
281 optogenetics and recordings; L.X. and H.Y.G. performed the experiments; L.X., S.I.W., R.T.,
282 A.H. and S.J.S. designed the analyses; L.X., R.T., and A.H. performed the analyses; S.J.S.,
283 S.I.W., and L.X. wrote the paper. All authors approved of the final version of the manuscript.

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291 **9 Contribution to the field statement**

292 Periodic oscillations of excitability coordinate neuronal activity within and between brain
293 structures for perception, cognition and goal-directed behavior, processes implicating
294 noradrenergic activity in forebrain circuits. To better understand the link between the time scales
295 of behavior (on the order of seconds) and underlying neuronal processing (on the order of
296 milliseconds), we recorded phase-locking of neurons in the noradrenergic locus coeruleus to
297 brain oscillations in rats performing in a maze. Most neurons synchronized with hippocampal and
298 prefrontal cortical infra-slow (~0.4 Hz) rhythms. The infra-slow rhythms phase-locked to
299 principal events in the maze, and thus were not strictly periodic. They modulated the amplitude of
300 gamma rhythms, known to coordinate neuron activity, and thus could provide a scaffold linking
301 behavior to neuronal activity.

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304 **10 References**

305 Berridge, C.W., and Foote, S.L. (1991). Effects of locus coeruleus activation on
306 electroencephalographic activity in neocortex and hippocampus. *J. Neurosci.* 11(10), 3135-45.
307 doi: 10.1523/JNEUROSCI.11-10-03135.1991

308 Bosman, C.A., Schoffelen, J.-M., Brunet, N., Oostenveld, R., Bastos, A.M., Womelsdorf, T., et
309 al. (2012). Attentional stimulus selection through selective synchronization between monkey
310 visual areas. *Neuron*. 75, 875–888. doi: 10.1016/j.neuron.2012.06.037.

311 Bouret, S., and Sara, S.J. (2005). Network reset: a simplified overarching theory of locus
312 coeruleus noradrenaline function. *Trends Neurosci.* 28(11), 574-82. doi:
313 10.1016/j.tins.2005.09.002

314 Buehlmann, A., and Deco, G. (2010). Optimal information transfer in the cortex through
315 synchronization. *PLoS Comput. Biol.* 6. doi: 10.1371/journal.pcbi.1000934

316 Canavier, C.C. (2015). Phase-resetting as a tool of information transmission. *Curr. Opin.*
317 *Neurobiol.* 31, 206-213. doi:10.1016/j.conb.2014.12.003

318 Eckhorn, R., Reitboeck, H.J., Arndt, M., and Dicke, P. (1990). Feature linking via
319 synchronization among distributed assemblies: Simulations of results from cat visual cortex.
320 *Neural Comput.* 2, 293–307. <https://doi.org/10.1162/neco.1990.2.3.293>

321 Engel, A.K., Fries, P., König, P., Brecht, M., and Singer, W. (1999). Temporal binding, binocular
322 rivalry, and consciousness. *Conscious. Cogn.* 8, 128–51.

323 Eschenko, O., Magri, C., Panzeri, S., and Sara, S.J. (2012). Noradrenergic neurons of the locus
324 coeruleus are phase-locked to cortical Up-Down states during sleep. *Cereb. Cortex* 22(2), 426-43.
325 doi:10.1093/cercor/bhr121

326 Fries, P.A. (2005). A mechanism for cognitive dynamics: neuronal communication through
327 neuronal coherence. *Trends Cogn. Sci.* 9, 474–80. doi: 10.1016/j.tics.2005.08.011.

328 Halgren, M., Fabó, D., Ulbert, I., Madsen, J.R., Erőss, L., Doyle, W.K., et al. (2018). Superficial
329 slow rhythms integrate cortical processing in humans. *Sci. Rep.* 8(1), 2055. doi: 10.1038/s41598-
330 018-20662-0.

331 Hazan, L., Zugaro, M., and Buzsáki, G. (2006). Klusters, NeuroScope, NDManager: A free
332 software suite for neurophysiological data processing and visualization. *J. Neurosci. Meth.* 155,
333 207–216. doi: 10.1016/j.jneumeth.2006.01.017.

334 Hickey, L., Li, Y., Fyson, S.J., Watson, T.C., Perrins, R., Hewinson, J., et al. (2014).
335 Optoactivation of locus ceruleus neurons evokes bidirectional changes in thermal nociception in
336 rats. *J. Neurosci.* 34, 4148–4160. DOI: 10.1523/JNEUROSCI.4835-13.2014

337 Hwang, D.Y., Carlezon, W.A., Isacson, O., and Kim, K.S. (2001). A high-efficiency synthetic
338 promoter that drives transgene expression selectively in noradrenergic neurons. *Human Gene
339 Ther.* 12, 1731–1740. doi: 10.1089/104303401750476230.

340 Kraus, B.J., Robinson, R.J., 2nd, White, J.A., Eichenbaum, H., and Hasselmo, M.E. (2013).
341 Hippocampal “time cells”: time versus path integration. *Neuron* 78, 1090–1101. doi:
342 10.1016/j.neuron.2013.04.015.

343 Lestienne, R., Hervé, A., Robinson, D., Brios, L., and Sara, S.J. (1997). Slow oscillations as a
344 probe of the dynamics of the locus coeruleus-frontal cortex interaction in anesthetized rats. *J.
345 Physiology (Paris)* 91, 273-284. doi: 10.1016/s0928-4257(97)82407-2.

346 Li, Y., Hickey, L., Perrins, R., Werlen, E., Patel, A.A., Hirschberg, S., et al. (2016). Retrograde
347 optogenetic characterization of the pontospinal module of the locus coeruleus with a canine
348 adenoviral vector. *Brain Res.* 1641(Pt. B), 274–290. doi: 10.1523/JNEUROSCI.4835-13.2014

349 MacDonald, C., Lepage, K., Eden, U., and Eichenbaum, H. (2011). Hippocampal “time cells”
350 bridge the gap in memory for discontiguous events. *Neuron* 71(4), 737–749. doi:
351 10.1016/j.neuron.2011.07.012.

352 Molter, C., O'Neill, J., Yamaguchi, Y., Hirase, H., and Leinekugel, X. (2012). Rhythmic
353 modulation of theta oscillations supports encoding of spatial and behavioral information in the rat
354 hippocampus. *Neuron* 75, 889–903. doi: 10.1016/j.neuron.2012.06.036

355 Oberto, V.J., Boucly, C.J., Gao, H., Todorova, R., Zugaro, M.B., and Wiener, S.I. (2022).
356 Distributed cell assemblies spanning prefrontal cortex and striatum. *Curr Biol.* 32(1), 1-13.e6.
357 doi: 10.1016/j.cub.2021.10.007

358 Oe, Y., Wang, X., Patriarchi, T., Konno, A., Ozawa, K., Yahagi, K., et al. (2020). Distinct
359 temporal integration of noradrenaline signaling by astrocytic second messengers during vigilance.
360 *Nat. Comm.* 11(1), 471. doi: 10.1038/s41467-020-14378-x

361 Pastalkova, E., Itskov, V., Amarasingham, A., and Buzsáki, G. (2008). Internally generated cell
362 assembly sequences in the rat hippocampus. *Science* 321(5894), 1322-7. doi:
363 10.1126/science.1159775.

364 Ravassard, P., Kees, A., Willers, B., Ho, D., Aharoni, D., Cushman, J., et al. (2013).
365 Multisensory control of hippocampal spatiotemporal selectivity. *Science* 340, 1342–1346.
366 doi: 10.1126/science.1232655

367 Sara, S.J. (2009). The locus coeruleus and noradrenergic modulation of cognition. *Nat Rev
368 Neurosci.* 10(3), 211-23. doi: 10.1038/nrn2573.

369 Sara, S.J., and Bouret, S. (2012). Orienting and reorienting: the locus coeruleus mediates
370 cognition through arousal. *Neuron* 76(1), 130-41. doi: 10.1016/j.neuron.2012.09.011.

371 Sara, S.J., and Hervé-Minvielle, A. (1995). Inhibitory influence of frontal cortex on locus
372 coeruleus neurons. *Proc. Nat. Acad. Sci. (USA)* 92, 6032-6036. doi: 10.1073/pnas.92.13.6032.

373 Steriade, M., Contreras, D., Curro Dossi, R., and Nunez, A. (1993). The infra-slow (<1 Hz)
374 oscillation in reticular thalamic and thalamocortical neurons: scenario of sleep rhythm generation
375 in interacting thalamic and neocortical networks. *J. Neurosci.* 13, 3284-3299. doi:
376 10.1523/JNEUROSCI.13-08-03284.1993.

377 Totah, N.K., Neves, R.M., Panzeri, S., Logothetis, N.K., and Eschenko, O. (2018). The locus
378 coeruleus is a complex and differentiated neuromodulatory system. *Neuron*. 99(5), 1055-1068.e6.
379 doi: 10.1016/j.neuron.2018.07.037.

380 Totah, N.K.B., Logothetis, N.K., and Eschenko, O. (2019). Noradrenergic ensemble-based
381 modulation of cognition over multiple timescales. *Brain Res.* 1709, 50-66. doi:
382 10.1016/j.brainres.2018.12.031.

383 Villette, V., Malvache, A., Tressard, T., Dupuy, N., and Cossart, R. (2015). Internally recurring
384 hippocampal sequences as a population template of spatiotemporal information. *Neuron*. 88(2),
385 357-66. doi: 10.1016/j.neuron.2015.09.052

386 Voloh, B., and Womelsdorf, T. (2016). A role of phase-resetting in coordinating large scale
387 neural networks during attention and goal-directed behavior. *Front. Syst. Neurosci.* 10, 18.
388 doi: 10.3389/fnsys.2016.00018

389 Wagatsuma, A., Okuyama, T., Sun, C., Smith, L.M., Abe, K., and Tonegawa, S. (2018). Locus
390 coeruleus input to hippocampal CA3 drives single-trial learning of a novel context. *Proc. Natl.*
391 *Acad. Sci. (USA)* 115(2), E310-E316. doi: 10.1073/pnas.1714082115.

392 Womelsdorf, T., Schoffelen, J.M., Oostenveld, R., Singe, W., Desimone, R., Engel, A.K., et al.
393 (2007). Modulation of neuronal interactions through neuronal synchronization. *Science* 316,
394 1609–1612. doi: 10.1126/science.1139597

395 Xiang, L., Harel, A., Gao, H., Pickering, A.E., Sara, S.J., and Wiener, S.I. (2019). Behavioral
396 correlates of activity of optogenetically identified locus coeruleus noradrenergic neurons in rats
397 performing T-maze tasks. *Sci. Rep.* 9(1), 1361. doi: 10.1038/s41598-018-37227-w.

398 **11 Figure Legends**

399 Figure 1. Calculation of LC spike phase relative to Hip or Pfc LFP. A) Unfiltered signal of a
400 hippocampal recording, with theta oscillations dominating. B) The signal from A band-pass
401 filtered at 0.1-1.0 Hz. Red dots indicate LC neuron action potentials in all panels. C) The
402 amplitude of the signal in B was z-scored. Low amplitude oscillations were excluded from

403 analyses according to a selected criterion of $z \leq 0$ (excluded zones are demarcated by the dotted
404 rectangles). D) Phase of the filtered signal in B. Note that the LC spikes generally occur at phases
405 between 0 and $\pi/2$ radians in this example. The discontinuities near 138.5 and 143 s correspond
406 to excluded data, where phase could not be computed reliably.

407 Figure 2. Spike phase-locking to infra-slow rhythms from two example LC neurons (A and B).
408 Radius values are spike counts. Red arrows represent resultant vectors.

409 Figure 3. A) The automated behavioral task. When the trained rat crosses the central arm
410 photodetector (VC onset PD), this triggers one of the two cue screens behind the reward arms to
411 be lit in pseudo-random sequence. Crossing the appropriate reward delivery PD triggers a drop of
412 sweetened water to arrive at the corresponding reward site. Crossing the VC OFF PD's on the
413 return arms triggers the lit screen to be turned off. These three photodetector events are used to
414 synchronize activity in other Figures. B) Distribution of mean phase (left) and p-values of phase-
415 locking (right; Rayleigh test) for Pfc infra-slow oscillations in pooled data from multiple sessions
416 (top), and in an example session (bottom).

417 Figure 4. An example of simultaneous recordings of Pfc and Hip LFP infra-slow oscillations
418 phase-locked to principal maze events, the PD crossings (at time zero). Each row of the color
419 plots corresponds to a single trial. The phase of the infra-slow LFP is color-coded. Black rings
420 correspond to the PD crossing prior to (left) or after (right) the event at zero for each plot. Note
421 that the time scales vary among the events, in order to display prior and subsequent PD's. The
422 dark traces in the 2nd and 4th rows show mean phase while the lighter shaded upper and lower
423 lines are \pm SEM. In the middle column, the blue vertical bars and blue double-headed arrow
424 illustrate the calculation of the range of regular phase-locking (defined here as the period with the
425 criterion of SEM range $< 0.75 \pi$ radians; pink double-headed arrows). Here, desynchronization
426 (zones with wider SEM ranges) and discontinuities in the mean phase result from inter-trial
427 variability in speed and distance from the synchronization point. (PD - photodetector crossing).
428 This is from the same session as the recording in Fig. 3B.

429 Figure 5. Example of infra-slow modulation of gamma rhythm LFP in Pfc (top) and Hip
430 (bottom). The black trace schematically illustrates the phase of the infra-slow rhythms. Gamma

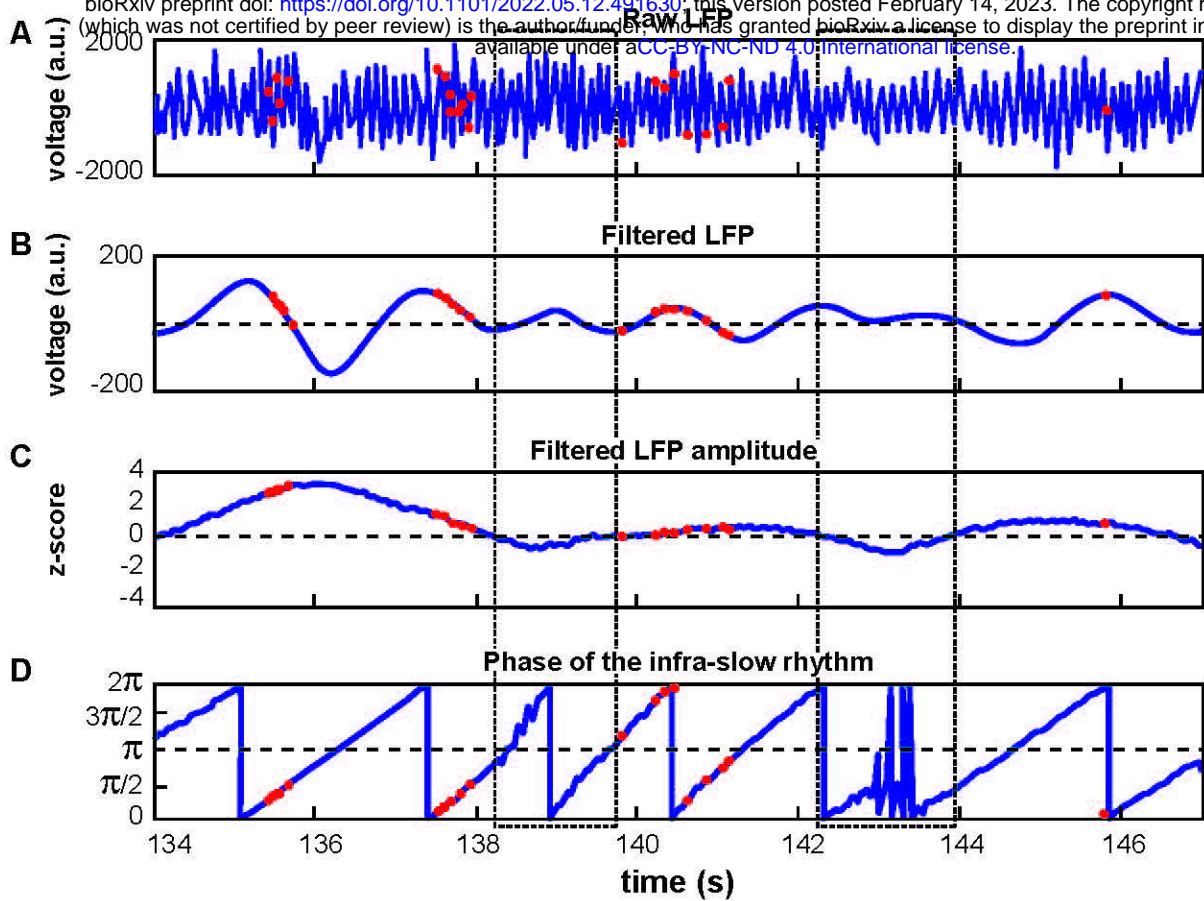
431 power was elevated between the phases of zero and pi radians of the infra-slow rhythm, while
432 this was not apparent at other frequencies. a.u. – arbitrary units.

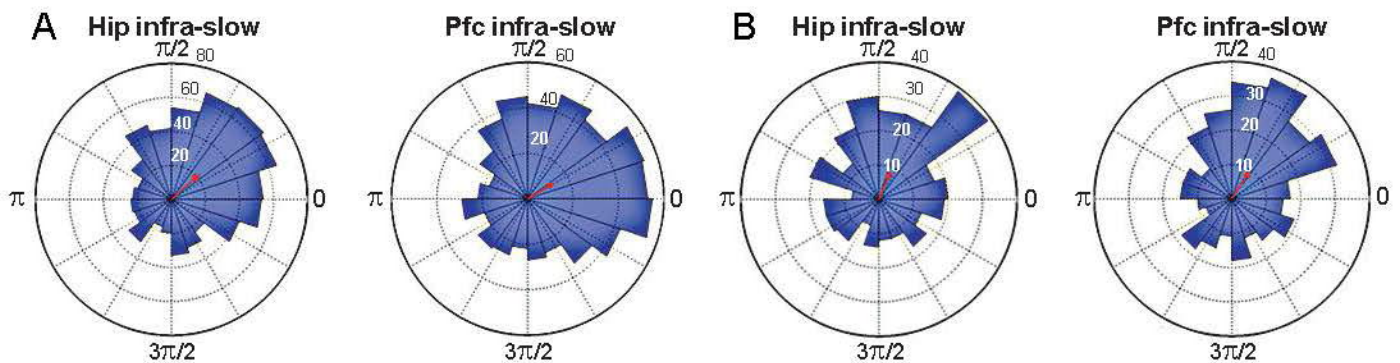
433 **12 Table**

	Pfc Rtn	Pfc Rwd	Pfc VC	Hip Rtn	Hip Rwd	Hip VC
<1 cycle (n)	4	16	10	13	14	9
1 to 1.49 cycles (n)	6	20	6	3	28	8
1.5 to 1.99 cycles (n)	1	6	2	2	7	3
2 to 2.49 cycles (n)		4			1	
2.5 to 3 cycles (n)					1	
Mean cycle period (s)	2.48	2.22	2.05	2.62	2.45	2.32
Mean frequency (Hz)	0.40	0.45	0.49	0.38	0.41	0.43

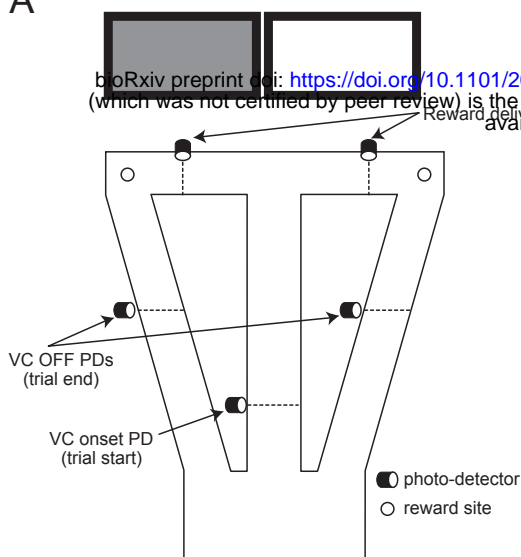
434

435 Table 1. Characterization of periods in sessions with regular phase-locking (SEM \leq 0.75* π
436 radians) of infra-slow LFP oscillations in prefrontal cortex (Pfc) and hippocampus (Hip) to task
437 events. In the six cases of two or more cycles, only data from the first cycle were counted for
438 mean cycle period and frequency. Cycles are only counted in the period from the previous trial
439 event to the next one, even though successive cycles could extend before or after (cf., Fig. 4,
440 Supp. Fig. 1).





A



B

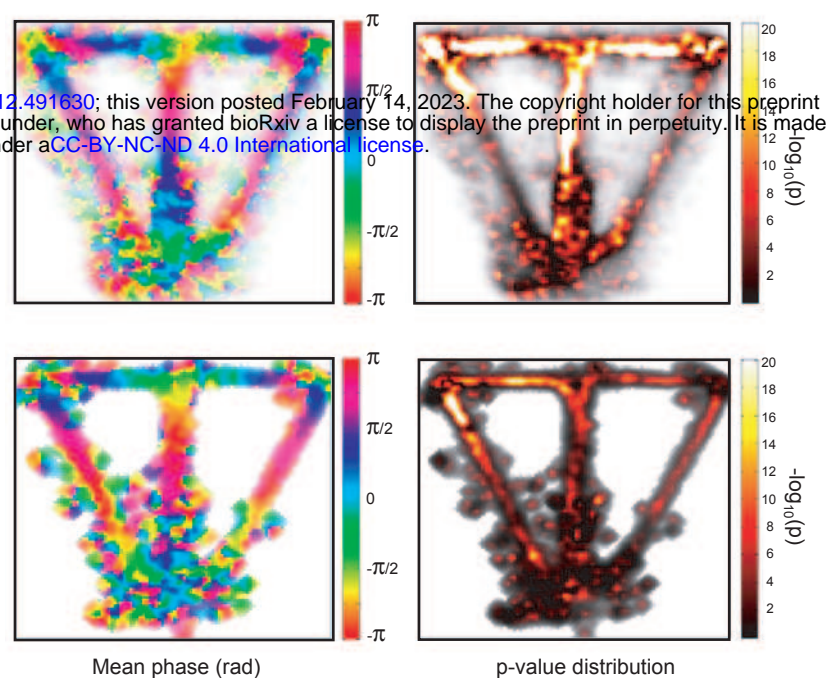


Figure 3. A) The automated behavioral task. When the trained rat crosses the central arm photodetector (VC onset PD), this triggers one of the two cue screens behind the reward arms to be lit in pseudo-random sequence. Crossing the appropriate reward delivery PD triggers a drop of sweetened water to arrive at the corresponding reward site. Crossing the VC OFF PD's on the return arms triggers the lit screen to be turned off. These three photodetector events are used to synchronize activity in other Figures. B) Distribution of mean phase (left) and p-values of phase-locking (right; Rayleigh test) for Pfc infra-slow oscillations in pooled data from multiple sessions (top), and in an example session (bottom).

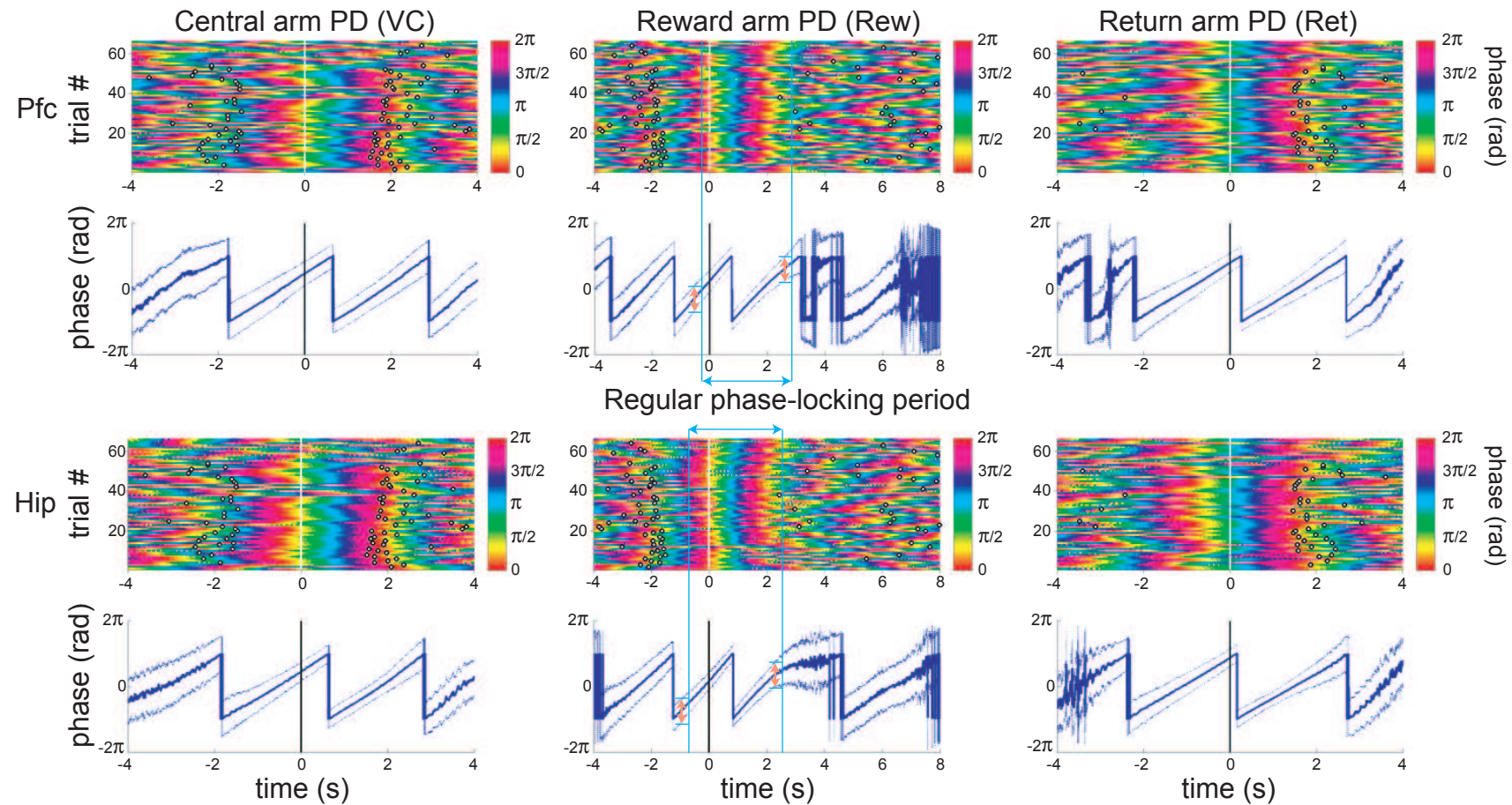


Figure 4. An example of simultaneous recordings of Pfc and Hip LFP infra-slow oscillations phase-locked to principal maze events, the PD crossings (at time zero). Each row of the color plots corresponds to a single trial. The phase of the infra-slow LFP is color-coded. Black rings correspond to the PD crossing prior to (left) or after (right) the event at zero for each plot. Note that the time scales vary among the events, in order to display prior and subsequent PD's. The traces in the 2nd and 4th rows show mean phase and dashed lines are \pm SEM. In the middle column, the blue vertical bars and blue double-headed arrow illustrate the calculation of the range of regular phase-locking (defined here as the period with the criterion of SEM range $< 0.75 \times \pi$ radians; pink double-headed arrows). Here, desynchronization (zones with wider SEM ranges) and discontinuities in the mean phase result from inter-trial variability in speed and distance from the synchronization point. (PD - photodetector crossing). This is from the same session as the recording in Fig. 3B.

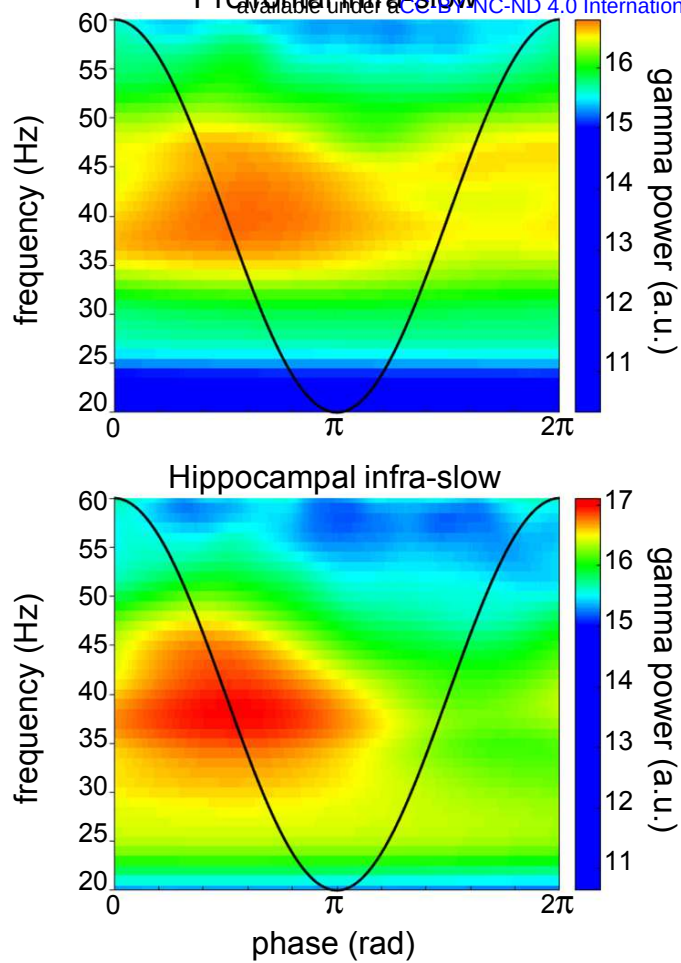


Figure 5. Example of infra-slow modulation of gamma rhythm LFP in Pfc (top) and Hip (bottom). The black trace schematically illustrates the phase of the infra-slow rhythms. Gamma power was elevated between the phases of zero and pi radians of the infra-slow rhythm, while this was not apparent at other frequencies. a.u. – arbitrary units.