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Characterizing brain stage-dependent pupil dynamics based on lateral hypothalamic activity

Kengo Takahashi 10^{1,2,3}, Filip Sobczak 10¹, Patricia Pais-Roldán 10⁴, Xin Yu 10^{5,*}

¹High-Field Magnetic Resonance Department, Max Planck Institute for Biological Cybernetics, 72076 Tübingen, Germany,

²Graduate Training Centre of Neuroscience, International Max Planck Research School (IMPRS), University of Tübingen, 72076 Tübingen, Germany, ³Cognitive and Systems Neuroscience Group, Swammerdam Institute for Life Sciences, Faculty of Science, University of Amsterdam, 1098XH Amsterdam,

The Netherlands,

⁴Medical Imaging Physics, Institute of Neuroscience and Medicine (INM-4), Forschungszentrum Jülich, 52425 Jülich, Germany,

⁵Athinoula A. Martinos Center for Biomedical Imaging, Massachusetts General Hospital, Harvard Medical School, Charlestown, MA 02129, United States

*Corresponding author: 149 13th. Street, Charlestown, MA 02129, United States. Email: xyu9@mgh.harvard.edu

Pupil dynamics presents varied correlation features with brain activity under different vigilant levels. The modulation of brain dynamic stages can arise from the lateral hypothalamus (LH), where diverse neuronal cell types contribute to arousal regulation in opposite directions via the anterior cingulate cortex (ACC). However, the relationship of the LH and pupil dynamics has seldom been investigated. Here, we performed local field potential (LFP) recordings at the LH and ACC, and whole-brain fMRI with simultaneous fiber photometry Ca²⁺ recording in the ACC, to evaluate their correlation with brain state-dependent pupil dynamics. Both LFP and functional magnetic resonance imaging (fMRI) data showed various correlations to pupil dynamics across trials that span negative, null, and positive correlation values, demonstrating brain state-dependent coupling features. Our results indicate that the correlation of pupil dynamics with ACC LFP and whole-brain fMRI signals depends on LH activity, suggesting a role of the latter in brain dynamic stage regulation.

Key words: lateral hypothalamus; anterior cingulate cortex; multimodal resting-state fMRI; brain dynamic stages; pupillometry.

Introduction

Brain state fluctuation, as a fundamental feature of animal brain physiology, is related to arousal regulation during wakefulness, sleep, or under anesthesia (Saper et al. 2001; Brown et al. 2010; McGinley et al. 2015). A transition of brain dynamic stages can arise between two radically different sub-states such as REM and NREM sleep but also between more complex sub-states such as those emerging under anesthesia (Brown et al. 2010; Barttfeld et al. 2015; McCormick et al. 2020; Pais-Roldan et al. 2020). The brain dynamic stages can be assessed using eye-tracking (e.g. pupillometry) as an external readout in combination with direct brain functional measurements such as electrophysiological recordings or functional magnetic resonance imaging (fMRI) (McGinley et al. 2015; Chang et al. 2016; Yuzgec et al. 2018; Pais-Roldan et al. 2020). For instance, the slow cortical oscillations of the electrophysiological signals are negatively synchronized with pupil size fluctuations during low arousal states in unconscious animals (Yuzgec et al. 2018). Besides measuring cortical brain state changes, studies in subcortical areas have revealed the involvement of the lateral hypothalamus (LH) and multiple brainstem nuclei including the locus coeruleus (LC) and periaqueductal gray in the regulation of brain states (Saper et al. 2001; Carter et al. 2010; Weber et al. 2015; Liu et al. 2017; Szabadi 2018). In particular, LC-based noradrenergic regulation of the anterior cingulate cortex (ACC) activity is in tight relationship with the pupil size regulation (Carter et al. 2010; Costa and Rudebeck 2016; Reimer et al. 2016; Liu et al. 2017), presenting a crucial link between the ACC activity and different sleep states and wakefulness

(Gompf et al. 2010; Rudebeck et al. 2014). It should be noted that the spontaneous change of pupil size and associated brain dynamic stages are not solely dependent on the noradrenergic pathways; instead, they are likely orchestrated by multiple subcortical structures (Costa and Rudebeck 2016; Joshi et al. 2016; DiNuzzo et al. 2019). Interestingly, neural spike rates recorded in the ACC present both positive and negative correlations with pupil size change across trials (Joshi et al. 2016), suggesting more complex subcortical regulation of brain state-dependent pupil dynamics.

The LH is an important hub for brain state regulation (Saper et al. 2001) where different neural cell types co-exist, showing varied activities depending on the underlying arousal state (Adamantidis et al. 2007; Kelz et al. 2008; Hassani et al. 2009; Konadhode et al. 2013). Also, electrical activation of the LH directly causes pupil dilation (Ranson and Magoun 1933). The LH activity has been reported to modulate LC-mediated arousal state and pupil dynamics (Hagan et al. 1999; de Lecea 2015; Herrera et al. 2016). There is anatomical evidence showing direct neural projections of arousal-related nuclei from the LH to the ACC (Jin et al. 2016). However, the functional interaction between the LH and ACC has not been fully investigated to elucidate their impact on brain dynamic stages and pupil dynamics.

Here, we measured brain dynamic stages with pupillometry, electrophysiology in the LH and ACC, and whole-brain fMRI with simultaneous fiber photometry Ca²⁺ recording in the ACC. We first identified a biphasic distribution of the correlation between pupil size and power of local field potential (LFP) in both LH and ACC across different trials of recordings. Based on opposite signs of the correlation (positive vs. negative), we discovered two distinct brain dynamic stages and performed cross-frequency coupling analyses to identify brain state-specific characteristics of LFP signals for the two stages. Specifically, the LFP delta (1–4 Hz) phase coupled stronger to amplitudes of higher frequency bands at the ACC in trials where LH-LFP and pupil were negatively correlated, suggesting the trials with negative signs as a lower arousal state (Mukamel et al. 2011; Nair et al. 2016; Pal et al. 2017). Meanwhile, similar to the LFP-based pupil dynamic correlation, the ACC neuronal Ca²⁺ also showed an opposite correlation with pupil dynamics across different trials. Interestingly, in trials with the positive correlation between LH fMRI and pupil dynamics, the ACC Ca²⁺ signal showed a primarily negative correlation with pupil dynamics. This observation suggested that similar brain dynamic stages could be identified in electrophysiology and fMRI measurements based on LH activity. This work revealed unique LH activity patterns with both LFP and fMRI according to its different coupling features with pupil dynamics, which were associated with different brain dynamic stages.

Materials and methods Subjects

All experiments were performed in accordance with the European Communities Council Directive (2010/63EU) and the German Animal Protection Law and approved by the Animal Protection Committee of Tübingen (Regierungspräsidium Tübingen, Germany). Nine Sprague Dawley rats [Strain: Crl:CD (SD), Sex: Male, Age: 3–6 months, Charles River Laboratories, Sulzfeld, Germany] were used for the multimodal electrophysiology experiments and nine Sprague Dawley rats for the multimodal fMRI experiments. All experiments were performed under alpha-chloralose anesthesia. The multimodal fMRI data had been previously published (Pais-Roldan et al. 2020). Since our study was exploratory research, the sample size was determined based on similar experiments in previous studies (Hutchison et al. 2010; Paasonen et al. 2018).

Surgical procedures for electrophysiology

Prior to the acquisition of the electrophysiology data, rats were anesthetized with 5% isoflurane and orally intubated with a 14-gage cannula. They were ventilated with a mixture of 2% isoflurane, 30% oxygen, and 70% air at 60 ± 1 breaths per minute by a ventilator (CW-SAR-830/AP) during surgery. The body temperature was maintained at 37-38°C by a heating pad. In order to monitor blood pressure and infuse anesthesia intravenously, cannulation of the femoral artery and vein were performed using polyethylene tubes (PE-50). Afterwards, rats were placed pronely on a stereotactic device, and two craniotomies were performed using a drill over the ACC and LH. A tungsten electrode (the resistance at 1 $M\Omega$; FHC Inc) was placed at the ACC (M/L = +0.5 mm, A/P = +1.2 mm, D/V = -1.8 mm) and LH (M/L = +1.5 mm, A/P = -3.2 mm, D/V = -7.8 mm) respectively. A bolus of ~ 80 mg/kg alpha-chloralose anesthesia was intravenously administrated into the femoral vein and inhalation of isoflurane was stopped. Afterwards, a mixture of alpha-chloralose (~25 mg/kg/h) and pancuronium (~2 mg/kg/h), a paralyzer agent, was continuously infused through an infusion pump during the whole experiment. During the recording, eye drops were not used to ensure a reliable detection of the pupil since the use of eye drops could have interfered with physiological responses of the pupil as well as quality of video images. Once the experiment was

done, the location of the electrodes was confirmed by MR imaging (Supplementary Fig. 13).

Data acquisition of electrophysiology signals

The tungsten electrodes positioned in the ACC and LH respectively were connected to our custom-made pre-amplifier ($25 \times$ gain and 0.1 Hz high pass), and the acquired analog signals were amplified and filtered by MCP-Plus 8 ($20 \times$ gain and 10 kHz low pass; Alpha Omega LTD). The total amplification was therefore 500. A sliver wire was placed between the skull and skin above the cerebellar region as a reference. The analog signals were converted to digital signals by the CED Power1401-3A (Cambridge Electronic Design), and the digital signals were recorded by Spike2 (CED) software at 25 kHz of sampling rate. To ensure synchronization between electrophysiology and pupil size recordings, the start time to record electrophysiology signals was set to be consistent with the start time of pupil size detection by a MacroRecoder software.

Data acquisition of pupillometry and extraction of pupil diameter

The pupil size was recorded by a customized video camera (dimensions: 30 mm \times 21 mm \times 15 mm, focal length of the lens: 10 mm; RS-OV7949–1818, Conrad) in a dark room during the experiments. Infrared LED light (880 nm of wavelength) was added in order to acquire a high contrast of the pupil image. The videos were recorded with 30 frames/s, 8 bits per pixel in RGB24, and 628 \times 586 pixels during the electrophysiology experiments and 29.97 frames/s, 8 bits per pixel in RGB24, and 240 \times 352 pixels during the fMRI experiments. The acquired video was processed by the DeepLabCut toolbox (Mathis et al. 2018; Nath et al. 2019) in order to extract the pupil diameter from each video frame. Four points were labeled at the edge of pupil, and the pupil diameter was calculated based on these four points with the following formula.

$$d = \frac{\sqrt{(x_2 - x_1)^2 + (y_2 - y_1)^2} + \sqrt{(x_4 - x_3)^2 + (y_4 - y_3)^2}}{2}$$

The extracted pupil diameter was first smoothed with a 10 point moving average and then down-sampled to 1 point per 1 s window for fMRI synchronization.

Data acquisition of MR scans

Prior to the acquisition of MR scans, the same preparatory surgical procedures for electrophysiology such as femoral cannulation were performed. Anesthesia was also provided as described above. All MR scans were performed on a 14.1 T/26 cm magnet (Magnex, Oxford) with an Avance III console (Bruker, Ettlingen), as previously described (Pais-Roldan et al. 2020). A custom-made elliptic trans-receiver surface coil (\sim 2 × 2.7 cm) was used for signal transmission and reception. For the acquisition of functional data, a 3D gradient echo echo-planar imaging (GE-EPI) sequence covering the whole brain was used with the following sequence parameters: 1 s repetition time (TR), 12.5 ms echo time (TE), 14° flip angle, $48 \times 48 \times 32$ matrix size, and $400 \times 400 \times 600 \ \mu$ m resolution. The total scan time was 15 min 25 s (925 TRs) for each EPI sequence. For the registration of MR images to a template, an anatomical RARE image was acquired with the following parameters: 4 s TR, 9 ms TE, 128 × 128 matrix size, 32 slices, 150 μ m in-plane resolution, 600 μ m slice thickness, 8× RARE factor.

Data acquisition of Ca²⁺ signals

The GCaMP (AAV5.Syn.GCaMP6f.WPRE.SV40) fluorescence signal was acquired from the ACC (M/L=+0.5 mm, A/P=+1.2 mm, D/V=-1.8 mm) via an optical fiber (FT200-EMT; NA=0.39; 200 μ m diameter, Thorlabs), connected to a light path setup, as previously described (Pais-Roldan et al. 2020). The signals were recorded at 5 kHz sampling rate using an analog input to digital converter in Biopac 150 system (Biopac, Goleta, CA, USA).

Analysis of electrophysiology signals

The acquired data were processed in Matlab. In order to extract the power of LFP signals from every second, spectral analysis using wavelet decomposition was performed with 2 s sliding windows between 0.5 Hz and 100 Hz. The extracted LFP powers were categorized into different frequency bands based on a previous animal sleep study (Yuzgec et al. 2018) as follows: delta (1–4 Hz), theta (4–7 Hz), alpha (7–14 Hz), beta (15–30 Hz), low gamma (30– 60 Hz), and high gamma (60–100 Hz). The extracted power time series was first smoothed with a 11 point moving average. The correlation between the power of each frequency band and pupil diameter changes was assessed after synchronizing both signals, using the Matlab corrcoef function.

Coherence

Coherence analysis is a standard method to evaluate functional connection of different brain regions in the frequency domain in electrophysiology studies (Nunez et al. 1997). Prior to evaluating the coherence, the raw data was downsampled from 25,000 to 500 sampling points per second. In the coherence analysis between the ACC and LH electrophysiology recordings sites, the Matlab mscohere function was used to calculate the magnitude-squared coherence with a 2 s Hanning window and an overlap of 1 s. This function has been used to compute functional connectivity measures in electrophysiology studies (Chapeton et al. 2019).

Phase locking value analysis

The phase locking value (PLV) measures synchronization of the phase between two different brain regions to investigate functional connectivity (Lachaux et al. 1999). Prior to calculating the PLV, the raw data was downsampled from 25,000 to 5,000 sampling points per second. The phase of each LFP was extracted using the Morlet wavelet and the fast Fourier transform from LH and ACC LFP signals, respectively. Then, the PLV was calculated with the following formula:

$$PLV = \left| \frac{\sum_{t=1}^{n} e^{i(\phi_{LH(t)} - \phi_{ACC(t)})}}{n} \right|$$

where n represents the number of sampling points, ϕ represents phase angles with the same length of sampling points, and e^{i} represents the exponential integral.

Phase-amplitude coupling

The phase amplitude coupling, using the modulation index (MI), is a well-used method to evaluate modulation of brain functions; it estimates how the phase of the low-frequency oscillation (e.g. delta band) couples with the amplitude of the high frequency oscillation (e.g. gamma band) (Tort et al. 2010; Mukamel et al. 2011). Prior to calculating the MI, the amplitude of the upper envelope of the LFP signal in each frequency band was extracted using the Matlab envelope function, and the phase angles of

each frequency band were extracted using the Matlab hilbert and angle functions. Then, the MI was calculated using the Kullback-Leibler (KL) distance between the amplitude and phase distributions as described in Tort et al. (2010) (Tort et al. 2010) using the "modulation_index" Matlab function (https://github. com/pierremegevand/modulation_index).

Analysis of fMRI signals

The EPI data from each animal were registered to an anatomical scan (RARE) of the corresponding animal. To spatially match data across animals, one RARE scan was used as a template and other anatomical scans were registered to it, while saving the transformation matrices. Finally, the saved transformation matrices were used to register all functional data together. A bandpass filter was applied to the fMRI signals between 0.01 and 0.1 Hz. These fMRI signals were correlated with pupil diameter changes and the power of 1–4 Hz Ca²⁺ signals using the Matlab corr function. With the purpose of comparing the whole brain fMRI map with previous publications (Schneider et al. 2016; Pais-Roldan et al. 2020), the voxel-wise correlation t-value was calculated based on the following formula:

$$t = \frac{r\sqrt{n-2}}{\sqrt{1-r^2}}$$

where r represents the correlation coefficient and n represents the number of sampling points from each run. Once the whole brain t-value map was generated, it was registered on the anatomical image using the AFNI software package (Cox 1996). In addition, these anatomical scans were overlapped with a rat brain atlas (Paxinos and Watson 2007) to identify the location of brain regions.

Analysis of Ca²⁺ signals

The acquired optical fiber Ca²⁺ signals (GCaMP) were processed by Matlab. The Ca²⁺ signal was first synchronized with fMRI signals with a customized Matlab code. In order to extract the power of Ca²⁺ signals, spectral analysis using wavelet decomposition was performed with 2 s sliding windows between 0.5 Hz and 100 Hz every second. Then, 1–4 Hz of Ca²⁺ powers, equivalent to the delta band of LFP, were extracted. The power of 1–4 Hz Ca²⁺ signals was correlated with pupil diameter changes and fMRI signals using the Matlab corrcoef function.

Analysis to determine the thresholds for the positive, no, and negative correlation ACC and LH electrophysiology versus pupil

The thresholds for positive and negative correlations were statistically determined by taking the 95% confidence interval from correlations between randomized pupil signals and LFP delta power with 10,000 times of permutation using the Matlab randperm and corrcoef functions (Supplementary Fig. 3). Trials above the upper confidence limit (correlation coefficients >0.065) were defined as positive correlations and those below the lower confidence limit (correlation coefficients < -0.065) as negative correlations. Trials remaining within this 95% confidence interval were considered as uncorrelated. Using the randomized pupil signal, distribution of correlation coefficients with LFP delta power across all trials was examined to assess whether this distribution of correlations with randomized data deviates from the true data. The result shows that distributions between the true and randomized results differ (Fig. 1e, Supplementary Fig. 3), indicating that correlation between pupil and LFP signals are not random.



Fig. 1. a) The experimental setup for simultaneous electrophysiology and pupil size recordings. Images of pupillary response (constricted and dilated pupil) are shown. b) Power spectrum of LFP signals in the ACC with pupil diameter fluctuations (pink trace) in a representative rat during 15 min of resting state recording, c) Correlation coefficients between pupil dynamics and power of all LFP bands in the ACC (circular markers) and in the LH (cross markers) from all trials (n = 89, 9 animals). Each marker represents the positive (green), no (gray), or negative (red) correlation on each trial. Thresholds of positive (0.046) or negative (-0.046) correlations are shown as blue dashed lines. d) Examples of positive (green shade) and negative (red shade) correlation between pupil dynamics (pink trace) and LFP delta (1-4 Hz) band power fluctuations (blue trace) in the LH and ACC during 15 min of resting state (all plots have the same scale). e) Distribution of correlation coefficients between LFP delta power fluctuation and pupil dynamics in the LH (left; [+ (green)]: N = 46, 52%, [0 (gray)]: N = 4, 4%, [- (red)]: N = 39, 44%) and ACC (right; [+ (green)]: N = 12, 13%, [0 (gray)]: N = 7, 8%, [- (red)]: N = 70, 79%). Each pie chart represents ratios of the positive, no, and negative correlations. Vertical blue solid lines represent the mean of correlation coefficients across all trials in the LH and ACC, respectively. Pink curves and boxplots represent the distribution of trials for LH and ACC, respectively. The vertical black dashed line represents zero. f) Violin plots for the distribution of correlation coefficients between pupil dynamics and LFP delta power fluctuation in LH and ACC. g) Categorization of correlation coefficients in the ACC between LFP delta power fluctuation and pupil dynamics when LH delta power fluctuation has a negative (LFP-group 1, orange) or positive (LFP-group 2, blue) correlation with pupil dynamics. In LFP-group 1, ACC delta power fluctuation and pupil size fluctuation show a negative correlation (correlation coefficient < -0.046) in all trials. In LFP-group 2, correlations of ACC delta power fluctuation and pupil dynamics diversify to a great extent. Note: [+]: Correlation coefficient > 0.046, [0]: -0.046 < correlation coefficient < 0.046, [-]: Correlation coefficient < -0.046; ***P < 0.001, **P < 0.01, *P < 0.05

Pupil versus ACC Ca²⁺

The thresholds for positive and negative correlations were statistically determined by taking the 95% confidence interval from correlations between randomized pupil signals and 1–4 Hz Ca²⁺ power with 10,000 times of permutation using the Matlab randperm and corrcoef functions (Supplementary Fig. 9). Trials above the upper confidence limit (0.065) were defined as positive correlations and below the lower confidence limit (-0.065) were defined as negative correlations. Otherwise, trials within this 95% confidence interval were considered as no correlation. Using the randomized pupil signal, distribution of correlation coefficients with 1-4 Hz Ca²⁺ power across all trials was examined to see if this distribution of correlations with randomized data deviates from the true data. The result shows that distributions between the true and randomized results differ (Fig. 3d, Supplementary Fig. 9), indicating that correlation between pupil and Ca²⁺ signals are not random.

fMRI versus pupil

The two different brain dynamic stages (positive or no correlation between BOLD fMRI and pupil dynamics) were statistically determined by taking the 95% confidence interval from correlations between the randomized pupil signal and fMRI signal in the LH with 10,000 times of permutation using the Matlab randperm and corr functions. The result shows that the distribution of correlations with randomized data differs from the true data (Supplementary Fig. 12), indicating that correlations between pupil and fMRI signals are not random. Trials above the upper confidence limit (1.96) were defined as positive correlations and those between the upper and lower confidence limit (-1.96 < CI < 1.96) were defined as no correlation. The LH region of interest was located in a single slice centered at the same position as electrophysiology LH recordings (33 manually selected voxels at A/P = -3.2 mm). Tvalues for correlation were calculated for the whole brain analysis as described above. In addition, the correlation between pupil and each fMRI voxel was corrected by the false discovery rate using the "FDR" Matlab function with alpha 0.05 for 20,022 voxels (https://www.mathworks.com/matlabcentral/fileexchange/ 71734-fdr-false-discovery-rate). Thereby, the whole-brain maps show voxels of positive correlation with t-value >2.54 and negative correlation with t-value < -2.54 (Fig. 4c, Supplementary Fig. 11).

fMRI versus ACC Ca²⁺

T-values for correlation were calculated between the power of 1–4 Hz ACC Ca²⁺ and each fMRI voxel as described above. In addition, the correlation was corrected by the false discovery rate using the "FDR" Matlab function with alpha 0.05 for 20,022 voxels as described above. Thereby, the whole-brain maps show voxels of positive correlation with t-value > 2.25 and negative correlation with t-value < -2.25 (Fig. 3f and Fig. 4f, Supplementary Fig. 10a and b).

Results

Simultaneous pupillometry and electrophysiology recordings in the ACC and LH

A cycle of brain dynamic stages in anesthetized rats appears with a 10–13 min period on average (Clement et al. 2008; Whitten et al. 2009; Pagliardini et al. 2012), also comparable to the cyclic period of different brain dynamic stages in sleeping rats (Borbély 1976;

Trachsel et al. 1991). In this study, LFP was recorded at ACC and LH simultaneously with pupillometry for an epoch duration of 15 min to study the brain state-dependent pupil dynamics in anesthetized rats (Fig. 1a). The LFP power spectrum fluctuation was well-aligned with concurrent pupil diameter changes in a representative trial (Fig. 1b). The correlation between pupil diameter and LFP power in various frequency bands in the LH and ACC varied across trials from negative to positive as well as no coupling, as shown in (89 trials from 9 animals, Fig. 1c and d, Supplementary Fig. 1). The varied correlation patterns were also detected within the same animal (Supplementary Fig. 2). In particular, among all frequencies, the oscillation of the delta (1-4 Hz) power was most prominent throughout the entire recording period (Fig. 1b). Therefore, correlations of delta power with pupil dynamics were further explored to investigate the LFP coupling between LH and ACC.

The correlation between pupil dynamics and LFP delta power fluctuation was more evenly distributed in the LH (positive correlation: 52%, n = 46; no correlation: 4%, n = 4; negative correlation: 44%, n = 39; Fig. 1e left) than the ACC, which showed primarily negative correlation to pupil dynamics (positive correlation: 13%, n = 12; no correlation: 8%, n = 7; negative correlation: 79%, n = 70, Fig. 1e right). The correlation-based distribution patterns between the LH and ACC differed significantly, as assessed using the two-sample Kolmogorov-Smirnov test (Marsaglia et al. 2003) $(P = 5.1 \times 10^{-7})$; Fig. 1f). Based on the positive versus negative correlation features, we specified two groups, which were statistically determined by calculating the 95% confidence interval from 10,000 correlation measures computed between randomized pupil size and LFP delta power signals (Supplementary Fig. 3). When the LH delta power and pupil dynamics were negatively correlated (LFP-Group 1), ACC delta power fluctuation and pupil dynamics were negatively correlated in 100% of cases (Fig. 1g). However, when LH delta power fluctuation and pupil dynamics were positively correlated, possibly indicating a different brain dynamic stage (LFP-Group 2), the correlation patterns between ACC delta power and pupil dynamics exhibited both positive and negative correlation (Fig. 1g). Two-sample t-tests showed that correlations of ACC delta power and pupil dynamics between LFP-Group 1 and 2 occurred differently. (P = 5.2×10^{-10} ; Fig. 1g), possibly suggesting differences in brain dynamic stages based on the relationship between LH and pupil dynamics.

Characterizing distinct brain dynamic stages through cross-frequency coupling analyses

To further investigate two brain dynamic stages identified by distinct coupling features between the LH delta power fluctuation and pupil dynamics (Fig. 1g), we performed cross-frequency coupling analyses. In LFP-Group 1, the LH and ACC exhibited the peak power at 2–2.5 Hz (Fig. 2a, Supplementary Fig. 4). In contrast, in LFP-Group 2 (when the LH delta power fluctuation was positively correlated with pupil dynamics), the LH and ACC exhibited the peak power at 3 Hz (Fig. 2a, Supplementary Fig. 4). Similarly, analyses of both coherence and phase-locking values between the LH and ACC showed the highest values at 2 Hz in LFP-Group 1 but at 3 Hz in LFP-Group 2 (Fig. 2b-d, Supplementary Fig. 5, Supplementary Table 1). Two-sample t-tests between LFP-Group 1 and 2 phase-locking values showed significant difference at 2 Hz ($P = 4.8 \times 10^{-7}$) and 3 Hz ($P = 8.8 \times 10^{-7}$). These results indicate that the neuronal coupling between LH and ACC altered between two different states. Moreover, the modulation index, a measurement of the phase-amplitude coupling, between the phase of delta band and amplitudes of all other LFP frequency bands at the ACC was higher in LFP-Group 1 than LFP-Group 2 (Fig. 2e, Supplementary Table 2). Similar observation was also confirmed at the LH that the modulation index in LFP-Group 1 was higher than LFP-Group 2 except low gamma band (Supplementary Fig. 6, Supplementary Table 3). Two-sample t-tests between LFP-Group 1 and 2 showed significant difference on theta band (P= 1.9×10^{-12} for ACC, P= 2.6×10^{-8} for LH), alpha band ($P = 4.6 \times 10^{-12}$ for ACC, $P = 2.4 \times 10^{-7}$ for LH), beta band $(P=1.9 \times 10^{-11} \text{ for ACC}, P=3.2 \times 10^{-8} \text{ for LH})$, low gamma band $(P = 1.5 \times 10^{-5} \text{ for ACC})$, and high gamma band $(P = 5.9 \times 10^{-8} \text{ for }$ ACC, P = 0.0013 for LH) except low gamma band at the LH (P = 0.97). In addition, the results for the separation of two different groups and cross-frequency coupling analyses were consistent with those obtained when a higher threshold (correlation coefficients = 0.15) was used (Supplementary Figs. 7 and 8), suggesting that the use of +/-0.065 correlation coefficients provided physiologically meaningful results. These results validate the presence of two distinct brain dynamic stages involving both LH and ACC under anesthesia, which resembled previous sleep studies (Benoit et al. 2000; Nair et al. 2016; Olbrich et al. 2017) (see Discussion).

Verifying coupling features of neuronal Ca²⁺ signals with simultaneous pupil dynamics and fMRI

To assess the relationship between brain state-dependent pupil dynamics and whole-brain activity, blood oxygen level dependent (BOLD) fMRI with simultaneous pupillometry was performed as previously reported (61 trials from 10 animals) (Pais-Roldan et al. 2020). Meanwhile, we performed fiber photometry to record the GCaMP6-mediated neuronal Ca²⁺ signals in the ACC of these animals, as an alternative to LFP recordings, with better MRI compatibility (Fig. 3a) (Pais-Roldan et al. 2020). Similar to LFP results, the power of ACC Ca²⁺ signals also co-varied with spontaneous pupil size fluctuations (Fig. 3b). The power of 1-4 Hz Ca²⁺ signal in the ACC was primarily negatively correlated with pupil dynamics (positive correlation: 11%, n=7; no correlation: 20%, n = 12; negative correlation: 69%, n = 42; Fig. 3c and d), which was consistent with our electrophysiology results (Fig. 1e). The clustering of varied correlation features between the 1–4 Hz Ca²⁺ power fluctuation and pupil dynamics among different trials was determined based on the 95% confidence interval calculated by permutation (Supplementary Fig. 9). Besides pupil dynamics, ACC neuronal Ca²⁺ signal also showed positive and negative correlation with fMRI signals across different brain regions and trials (Fig. 3e, Supplementary Fig. 10a). Interestingly, in trials with a negative correlation between 1 and 4 Hz Ca²⁺ power fluctuation and pupil dynamics, the Ca²⁺ power fluctuation showed a positive correlation with fMRI fluctuation in cortical and thalamic areas but a negative correlation in subcortical areas, including the LH (Fig. 3e and f). In other trials, no significant correlation was detected between fMRI and the ACC Ca²⁺ power fluctuation (Supplementary Fig. 10b). These multimodal fMRI results indicated that correlation features between Ca^{2+} -based neuronal fluctuation of the ACC and BOLD fMRI signals vary among different brain regions according to different anesthetized brain dynamic stages.

Classifying distinct brain dynamic stages based on LH BOLD fMRI and pupil dynamics

Next, we examined if LH BOLD signal could be used to differentiate brain state-dependent pupil dynamics across different trials, as evaluated with LFP data in previous experiments. First, we created a mean correlation map between fMRI and pupil dynamics, showing a global negative correlation pattern as previously reported (Supplementary Fig. 11) (Pais-Roldan et al. 2020). Moreover, similar to the fMRI-Ca²⁺ correlation, the trialspecific correlation patterns between pupil dynamics and fMRI showed varied signs across different brain regions (Fig. 4a, Supplementary Fig. 11). In particular, at the LH, the t-value of correlation coefficient varied extensively across individual trials, showing positive (33%, n = 20) or no (67%, n = 41) correlations (Fig. 4b). We statistically determined this positive correlation by calculating the 95% confidence interval from 10,000 repetitions of correlation measures between randomized pupil signals and each LH fMRI voxel (Supplementary Fig. 12). In trials showing positive signs, robust positive correlation was also observed between the pupil dynamics and fMRI signal in the brainstem and basal forebrain regions but negative correlation was detected in thalamic and cortical regions (Fig. 4c), which was not detected in trials with no correlation between LH fMRI and pupil dynamics (Fig. 4c). Notably, when exploring the correlation of pupil dynamics and ACC 1–4 Hz Ca²⁺ power from the trials with the positive LH fMRIpupil correlation sign, all of the ACC Ca²⁺-pupil correlation coefficients were negative (Fig. 4d and e; fMRI-Group 1). This observation was consistent with the electrophysiological results when differentiating LFP-Group 1 and 2 based on LH LFP-pupil relationship (Fig. 1f). Two-sample t-tests for correlations of ACC 1–4 Hz Ca²⁺ power with pupil dynamics between fMRI-Group 1 and 2 showed a significant difference ($P = 3.7 \times 10^{-4}$; Fig. 4d). It should be noted that in LFP-Group 1 the negative LH LFP (delta power)-pupil correlation predicted the negative correlation of ACC LFP (delta power)-pupil correlation, while in the fMRI-Group 1 it was the positive LH-pupil correlation which predicted the negative ACC Ca²⁺-pupil correlation. This might be due to an anti-correlation between BOLD fMRI and delta power in the LH (Jaime et al. 2019) (see Discussion). In addition, whole-brain fMRI BOLD and ACC Ca²⁺ power coupling was higher in fMRI-Group 1 than fMRI-Group 2 (Fig. 4f), indicating distinct brain dynamic stages associated to the different groups. Overall, these results indicate that correlation of LH fMRI-pupil dynamics can identify distinct brain dynamic stages, similar to the LH-ACC observations of LFP-based brain dynamic stages.

Discussion

In the present study, we reported distinct coupling features between brain activity and pupillometry in anesthetized rats, presenting either positive or negative correlation based on multimodality measurements. When the LH LFP delta power fluctuation was negatively correlated to pupil size changes, the ACC LFP-pupil correlation was always negative. In contrast, varied correlation features of ACC-pupil were detected when the LH was positively correlated to pupil dynamics. Based on the LHpupil relationship, we identified distinct cross-frequency coupling features of the LFP signals detected in the LH and ACC, defining two different anesthetized brain dynamic stages (LFP-Group 1 vs. 2). In parallel, the trial-specific ACC Ca²⁺-pupil correlation was associated with the fMRI-based LH-pupil relationship similar to the LFP results. In summary, two distinct brain dynamic stages could be identified based on the LH-pupil relationship, which could be verified by analyzing the site-specific LFP crossfrequency coupling or whole-brain fMRI-based $\rm Ca^{2+}$ and pupil correlation patterns.

The brain state embraces a broad number of distinctive features physiologically and behaviorally that can be observed across multiple scales (Kringelbach and Deco 2020; McCormick et al. 2020). These can be as evident as those differentiating sleep



Fig. 2. a) Power spectrum density of the LH (left) and ACC (right) in LFP-group 2 (blue line) and LFP-group 1 (orange line). The shaded areas represent the standard error of the mean. The peak frequency in both LH and ACC is at 3 Hz in LFP-group 2 (blue line) but at 2 Hz in LFP-group 1 (orange line). b) The radial plots of the trial-specific phase angles ([degrees], gray) and phase-locking value (black) between LH and ACC at 2 Hz and 3 Hz in LFP-group 1 (orange) and LFP-group 2 (blue). c) Phase locking value between the LH and ACC in both groups. The highest phase synchrony occurred at 3 Hz in LFP-group 2 (blue) but at 2 Hz in LFP-group 1 (orange). d) Magnitude squared coherence between LH and ACC. The highest coherence occurred at 3 Hz in LFP-group 2 (blue) but at 2 Hz in LFP-group 1 (orange). The shaded areas represent the standard error of the mean. e) Phase-amplitude coupling using the modulation index (MI) at the ACC between the phase of delta (1–4 Hz) band and amplitude of other frequency bands. The Mi in LFP-group 1 (orange) is higher than LFP-group 2 (blue) on all frequency bands. Error bars indicate the standard deviation. Note: ***P < 0.001, **P < 0.01, *P < 0.05.

from wakefulness (e.g. readily observed as electrophysiological transitions) (Adamantidis et al. 2007; Yuzgec et al. 2018) or much less apparent, which is the case when assessing varying brain dynamic stages under anesthesia. An example of the latter scenario is provided here in the form of changes that we reliably observed only after multi-modal analysis in our data. In

order to detect the features of complex brain states, exploratory analyses complementary to routine electrophysiology assessments are commonly used, such as cross frequency analysis in electrophysiology and correlation analysis between pupil diameters and fMRI/electrophysiology signals (McGinley et al. 2015; Yellin et al. 2015; Yuzgec et al. 2018; Migliorelli et al. 2019).



Fig. 3. a) The experimental setup for multimodal resting-state BOLD fMRI recordings simultaneously with measurements of optical fiber Ca^{2+} signals and pupil dynamics. b) Power spectrum of ACC Ca^{2+} signals during 15 min of resting-state along with changes in pupil diameter (pink trace) in a representative rat. A.u: Arbitrary units. c) Examples of positive (right, green) and negative (left, red) correlations between pupil dynamics (pink) and ACC 1–4 Hz Ca^{2+} power fluctuation (navy blue) during 15 min of resting state. d) Distribution of correlation coefficients between pupil dynamics and ACC 1–4 Hz Ca^{2+} power fluctuation for all trials (n=61, 10 animals). These correlation coefficients are negative (correlation coefficient < -0.064) in 69% (n=42, red), none (-0.064 < correlation coefficient < 0.064) in 20% (n=12, gray), and positive (correlation coefficient > 0.064) in 11% (n=7) of the trials. The pie chart represents ratios of the positive, no, and negative correlations. e) Examples of positive and negative correlation between fMRI signal (brown) and ACC 1–4 Hz Ca^{2+} power (navy blue) fluctuations during 15 min of resting state. f) t-value maps of correlation between ACC 1–4 Hz Ca^{2+} power and each fMRI voxel when ACC 1-4 Hz Ca^{2+} power fluctuation and pupil dynamics were negatively correlated during 15 min of resting state (n=42.69%). The overlay shows voxels with t-value < -2.25 and t-value > 2.25. A coronal slice at coordinates A/P = -3.2 mm is shown magnified on the right, where the location of the LH is indicated with white arrows and black polygons.

In this study, we investigated the coupling features between pupil size and LFP power and between pupil size and fMRI signal during 15 min of recording sessions to evaluate different brain dynamic stages in anesthetized animals. Our results showed two opposite correlation patterns between pupil dynamics and LFP measured at the LH and ACC as well as different fMRI-pupil synchronization across trials (Fig. 5), which we interpreted as surrogates of distinct states of the brain. Importantly, our definition of brain state is based on our available multimodal observations (pupil size, two-point electrophysiology, fMRI, and calcium recordings); it should

be noted that more complex or even a mixture of several brain states could give rise to the states here-identified.

It is known that the neural activity in the LH plays an important role in changing and maintaining an ongoing brain state during wakefulness, sleep, and anesthesia (Saper et al. 2001; Kelz et al. 2008; Herrera et al. 2016; Yin et al. 2019). Awake-promoting neurons in the LH (e.g. orexin-producing neurons) reduce their neural firing during both slow-wave sleep (Lee et al. 2005) and anesthesia (Kelz et al. 2008). The anesthetized brain mimics different sleep stages (Vacas et al. 2013), suggesting common regulatory



Fig. 4. a) Examples of positive (left) and negative (right) correlations between pupil dynamics (pink) and fMRI signal fluctuations (green) during 15 min of resting state. b) The representation of two distinct brain states. Trials are sorted into two groups characterized by positive (fMRI-group 1) or no (fMRI-group 2) correlation between the LH BOLD fMRI signal and the pupil size fluctuations. On the top, two fMRI-pupil correlation maps at A/P = -3.2 mm from fMRI-group 1 and 2 are shown. The location of LH is indicated with a white arrow. The bottom graph represents the t-value distribution of the LH-fMRI and pupil dynamics correlations across all trials (n = 61), on which the state differentiation is based. fMRI-group 1 (positive correlation) was observed in 33% of the trials (n = 20, t > 1.96), and fMRI-group 2 (no-correlation) in 67% of trials (n = 41, t < 1.96). c) Whole-brain maps and their sagittal plane view of the correlation between fMRI and pupil dynamics from trials associated with fMRI-group 1 (top) and 2 (bottom). d) Correlation coefficients between pupil dynamics and ACC 1–4 Hz Ca²⁺ power fluctuation sorted by fMRI-group 1 (orange ellipse) and 2 (but ellipse). The thresholds of correlation coefficients are set at +/- 0.064. e) the plot shows the correspondence between two kinds of analysis in the same data: Correlation t-values between LH-fMRI and pupil (left) and the correspondent ACC 1–4 Hz Ca²⁺ power and pupil dynamic correlation for each trial (right). The color indicates the groups (orange for fMRI-group 1 and blue for fMRI-group 2) associated with each trial. f) t-value whole-brain maps showing the correlation between fMRI and ACC 1–4 Hz Ca²⁺ power from trials assigned to fMRI-group 1 (top) and 2 (bottom). The overlay shows voxels with t-value < -2.54 and t-value > 2.54. Note: ***P < 0.001, **P < 0.05.



Fig. 5. A schematic diagram showing the sign of correlations across multimodal measurements. a) Correlations between pupil diameter and electrophysiology at the LH and ACC are described. Negative correlations between ACC LFP delta power and pupil diameter were always identified when LH LFP delta power was negatively correlated with pupil diameter. b) Correlations between pupil diameter and 1–4 Hz Ca²⁺ power at the ACC as well as fMRI signals at the LH are described. Negative correlations between ACC 1–4 Hz Ca²⁺ power and pupil diameter were always identified when the LH fMRI signal was positively correlated with pupil diameter. Negative correlation between LH fMRI and LH 1–4 Hz LFP power is also expected from previous works mentioned in the figure.

mechanisms to control the brain dynamic stages. In addition, optogenetic activation of awake promoting neurons in the LH produces cortical arousal and behaviorally awaking from anesthesia (Herrera et al. 2016) or sleep (Adamantidis et al. 2007). Under anesthesia, the neural firing of melanin-concentrating hormone neurons, promoting sleep (Konadhode et al. 2013), are unaffected in spite of the reduced neural firing in awake promoting neurons (Kelz et al. 2008). Therefore, different neural populations in the LH may contribute to the varied coupling features of pupil dynamics at different arousal states even under anesthesia. Moreover,

the LH activation causes pupil dilation (Ranson and Magoun 1933) due to inhibition of Edinger Westphal Nucleus (EWN) (Koss et al. 1984; Szabadi 2018). These attributes of the LH support our observation of two opposite correlation patterns (positive and negative) between the neuronal oscillation in the LH and the pupil dynamics (Fig. 1c and d).

We also revealed a unique LH-ACC interaction underlying the varied brain state-dependent pupil dynamics. The ACC is an important hub of the default mode network in both humans and rats (Lu et al. 2012), and its involvement in the resting-state

connectivity has been observed to differ between high and low arousal states, showing a decreased participation of the ACC in the default mode network during the lower arousal state compared to higher state (Horovitz et al. 2009; Tarun et al. 2021). The neural activity in the ACC has been linked to arousal fluctuation and associated with pupil size changes, which can be explained by its bidirectional connection to the noradrenergic LC (Gompf et al. 2010; Ebitz and Platt 2015; Joshi et al. 2016; Pais-Roldan et al. 2020). The opposite correlation features between ACC activity and pupil dynamics (Fig. 1c-e, Supplementary Fig. 2) could not be solely explained by noradrenergic projections from the LC. Given the functional and anatomical interaction between LH and ACC (Saper et al. 2001; Morgane et al. 2005; Gompf et al. 2010; Stevens et al. 2011; Jin et al. 2016), our work suggested that LH can play a pivotal role to mediate subcortical regulation of the ACC activity underlying brain state-dependent pupil dynamics. In the state when the LH delta power fluctuation was negatively correlated with pupil dynamics, the ACC LFP-pupil showed negative correlation (Fig. 1g). This LFP-based Group 1 likely corresponds to a lower arousal level based on evidence from the previous electrocorticogram (ECoG) study with pupil size measurements (Yuzgec et al. 2018); they demonstrated that cortical ECoG-based delta amplitude and pupil dynamics showed very low correlation during wakefulness but strong negative correlation during NREM sleep or moderate negative correlation during REM sleep. We also performed cross-frequency analyses of the LFP signals detected in the ACC and LH, which showed distinct LFP coupling features between LFP-Group 1 and 2 (Fig. 2b). Of note, this spectral-based differentiation of brain dynamic stages is in agreement with sleep EEG studies showing that during NREM sleep, i.e. the deepest state of unconsciousness, the cortical activity peaks ~2 Hz (Benoit et al. 2000; Rodenbeck et al. 2006; Olbrich et al. 2017). In contrast, during REM sleep, i.e. a state of relative awareness, the frequency peak of the EEG shifts to higher values (e.g. 2.5-3 Hz in frontal-central cortical regions) (Olbrich et al. 2017; Bernardi et al. 2019). We also observed the shift of power spectrum between the two LFP groups. The peak of power spectrum in the ACC was at 2 Hz in the trials where we observed negative correlations between the LH activity and pupil size but 3 Hz in the trials where we observed positive correlations (Fig. 2a, right). This shift indicates the changes of spontaneous activity in the ACC. Whether the observed changes in the ACC reflect a direct interaction of the ACC and LH remains unclarified. Future studies using optogenetics or an equivalent technique combined with calcium imaging/electrophysiology could potentially help clarifying the direct influence of one area on another upon the brain dynamic stages. Besides the frequency-specific coupling features of neural firing in the ACC and LH, the magnitude-squared coherence and phase locking analyses also showed distinct values between LFP-Group 1 and 2, which is similar to previous reports of brain state switches from conscious wakefulness to unconsciousness (e.g. anesthetized or deep sleep state) (Ishizawa et al. 2016; Miyamoto et al. 2016; Migliorelli et al. 2019). Since previous studies report that phase-amplitude coupling appears stronger in deeper anesthetized or sleep states (Mukamel et al. 2011; Nair et al. 2016; Pal et al. 2017), our work strongly suggests that the trials in LFP-Group 1 are at a lower arousal level than those in LFP-Group 2.

Pupil dynamics track changes in brain states, in particular, when combined with fMRI as a non-invasive arousal indicator (Yellin et al. 2015; Chang et al. 2016; Schneider et al. 2016). It has been shown that pupil dilation occurs concurrently with global negative BOLD-fMRI signal in anesthetized rats (Pais-Roldan et al. 2020), which coincides with decreased low-frequency power

recorded in the cortex in both sleeping and anesthetized animals (Blasiak et al. 2013; Yuzgec et al. 2018). It should be noted that the correlation features between pupil dynamics and fMRI or electrophysiological signals depend on the underlying brain dynamic stages (Yellin et al. 2015; Yuzgec et al. 2018; Pais-Roldan et al. 2020). Based on the fMRI signal detected in the LH, we have defined fMRI-Group 1 for positive correlation with pupil dynamics (33% of trials) and fMRI-Group 2 without significant correlation (~67% of trials) (Fig. 4b). Our results also showed brain-wide cortical negative fMRI correlation with pupil dynamics in fMRI-Group 1 but the negative correlation was not observed in fMRI-Group 2 (Fig. 4c). It is plausible that fMRI-Group 1 corresponds to a lower arousal state than fMRI-Group 2, similar to what has been defined by the LFP-Groups. Indeed, under this positive correlation between LH fMRI and pupil dynamics (fMRI-Group 1), we observed that Ca²⁺-based neuronal signals in the ACC and pupil dynamics were always negatively correlated, a result that is consistent with our electrophysiology findings (Fig. 4d and Fig. 1g). Furthermore, our BOLD fMRI and Ca²⁺-based neural correlations were higher in fMRI-Group 1 than in fMRI-Group 2 (Fig. 4f), indicating a change in brain dynamic stages based on LH fMRI-pupil relationships.

A healthy brain during resting-state transits between different states that can be identified based on correlation patterns (Tagliazucchi and Laufs 2014; Barttfeld et al. 2015; Zhou et al. 2019), and their relevance to clinical studies has been demonstrated (Ahmadi et al. 2021; Wu et al. 2021), implying that states identified through correlation changes are biologically meaningful. Although fMRI-pupil interaction shows a promising indicator to estimate brain dynamic stages noninvasively, more efforts remain needed to better elucidate the dynamic coupling features of fMRI and neuronal signals during brain state fluctuation across different brain regions (Mukamel et al. 2005; He et al. 2008; Conner et al. 2011; Magri et al. 2012). In the cortex, He et al. (He et al. 2008) demonstrated varied correlation between electrophysiological and BOLD signals in awake state and slow wave sleep. Furthermore, depending on eye open/closed arousal states in monkeys, correlation features between fMRI and LFP signals vary (Scholvinck et al. 2010), and these eye behaviors can track arousal states with fMRI (Chang et al. 2016). The fiber photometry-based Ca²⁺ (GCaMP) recording with simultaneous fMRI (He et al. 2018) revealed the brain state-dependent correlation features of BOLD signal with neuronal and astrocytic Ca²⁺ signaling, relying on the neuromodulation from thalamic or brainstem nuclei (Wang et al. 2018; Pais-Roldan et al. 2020). In our work, ACC 1-4 Hz Ca²⁺ power fluctuation was positively correlated with BOLD fMRI signals of the entire cortex (Fig. 3f, Supplementary Fig. 10a). In contrast, it was negatively correlated with BOLD fMRI in various subcortical regions including the LH, brainstem, and striatum (Fig. 3f, Supplementary Fig. 10a). Given the positive coupling feature between LH and ACC LFP data as described in LFP-Group 1, the positive correlation of LH fMRI with pupil dynamics in fMRI-Group 1 suggested negative correlation between LH fMRI and LFP. This negative correlation between BOLD fMRI and LFP low-frequency power, in particular at the delta band, has been previously observed in several brain regions including subcortical areas (e.g. striatum) (Hutchison et al. 2015; Jaime et al. 2019), suggesting a non-linear relationship between the neuronal and the vascular network, which is not totally understood.

In this study, we used alpha-chloralose anesthesia to obtain higher quality of BOLD fMRI responses (Hyder et al. 1994; de Celis et al. 2011). The use of alpha-chloralose induces slow-wave activities in the animal's brain (He et al. 2018; Pais-Roldan et al. 2020). Although this anesthetic stably maintains unconsciousness of animals, the state of brain dynamics fluctuates across time (Wang et al. 2018). In fact, our pupillometry shows that pupil size and LFP signals simultaneously change across time (Fig. 1b). Previous studies showed that different anesthetics (e.g. Urethane and isoflurane) also induced different dynamics of electrophysiological signals and simultaneous pupil size changes (Blasiak et al. 2013; Kum et al. 2016), similar to state-dependent changes of pupil size during different sleep stages (Yuzgec et al. 2018). Moreover, a recent study has also claimed state-dependent correlations between pupil size and both cortical blood volume and LFP, showing negative correlations during a naturally occurring deep unconscious state (NREM sleep) but positive during a shallow unconscious state (REM sleep) (Turner et al., 2023). These observations are comparable with our study that both LFP and BOLD fMRI signals show positive and negative correlations with pupil size depending on different brain dynamic stages. Although the mechanism of how different anesthetics or naturally occurring brain dynamic stages work can differ, our study presents a basis of common systemic features on pupil size and brain dynamics. This above content is now added in the discussion of the manuscript.

In summary, we reported varied brain state-dependent coupling features which are orchestrated in multiple regions (e.g. LH and ACC) and cross multiple scales (frequency and amplitude of neuronal firing, inter-region coherence, brain-wide fMRI, pupillary coupling, etc). This work offers a reference framework to understand the LH-ACC relationship, and, more importantly, it presents a novel way to track brain dynamic stage transitions by bridging pupillary changes with the LH activity. We demonstrate that LH LFP and fMRI signals provide a strong indication of the vigilance level; hence, we anticipate that their assessment may constitute a useful approach to evaluate the ongoing brain states in arousalrelated clinical research.

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Author contributions

Kengo Takahashi (Conceptualization, Data curation, Formal analysis, Investigation, Methodology, Software, Writing—original draft, Writing—review & editing), Filip Sobczak (Software, Writing review & editing), Patricia Pais-Roldán (Data curation, Writing review & editing), and Xin Yu (Conceptualization, Funding acquisition, Methodology, Supervision, Writing—review & editing).

Supplementary material

Supplementary material is available at Cerebral Cortex online.

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Data availability

The data are available from the corresponding author upon request.

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