

Brain-state-independent neural representation of peripheral stimulation in rat olfactory bulb

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It is critical for normal brains to perceive the external world precisely and accurately under ever-changing operational conditions, yet the mechanisms underlying this fundamental brain function in the sensory systems are poorly understood. To address this issue in the olfactory system, we investigated the responses of olfactory bulbs to odor stimulations under different brain states manipulated by anesthesia levels. Our results revealed that in two brain states, where the spontaneous baseline activities differed about twofold based on the local field potential (LFP) signals, the levels of neural activities reached after the same odor stimulation had no significant difference. This phenomenon was independent of anesthetics (pentobarbital or chloral hydrate), stimulating odorants (ethyl propionate, ethyl butyrate, ethyl valerate, amyl acetate, n-heptanal, or 2-heptanone), odor concentrations, and recording sites (the mitral or granular cell layers) for LFPs in three frequency bands (12–32 Hz, 33–64 Hz, and 65–90 Hz) and for multiunit activities. Furthermore, the activity patterns of the same stimulation under these two brain states were highly similar at both LFP and multiunit levels. These converging results argue the existence of mechanisms in the olfactory bulbs that ensure the delivery of peripheral olfactory information to higher olfactory centers with high fidelity under different brain states.

The ability to perceive the ever-changing external world accurately and precisely is a basic brain function. It is critical for daily life, survival, and higher brain functions, such as making decisions, plans, and judgments. However, the brain itself operates in ever-changing states (1, 2) or baselines (3) that are constantly modulated by external and internal factors, such as physical, physiological, psychological, clinical, and metabolic conditions. Therefore, elucidation of how sensory systems represent the same sensory information to the brain under different operational states is fundamental to understanding the related brain functions (1, 4–6).

The olfactory system, as the most direct and intrinsic sensory module (1, 7, 8), provides a unique model to study the principles and functional mechanisms of the brain. The peripheral olfactory information takes just one synapse to enter the olfactory bulb (OB) in the brain. The OB outputs directly to the olfactory cortices that mostly belong to the limbic system. As a center to code and process the peripheral olfactory inputs and a hub to distribute the processed information to the olfactory cortices (9–11), the OB consists of laminar structures, from the outer surface to the inner core: the olfactory nerve, glomerular, external plexiform, mitral cell layer (MCL), and granular cell (GCL) layer. All peripheral olfactory input is received by a few thousand glomeruli, which generate an odor-specific spatiotemporal activity pattern across the glomerular layer (10, 12, 13). The information is mainly processed in the external plexiform layer that contains a dense dendrodendritic network formed between the secondary dendrites of the mitral/tufted (M/T) cells, whose somas are located in the MCL, and the dendrites of granular cells, whose somas are located in the GCL, then sent to higher olfactory centers by M/T cells (11, 14, 15). Under given conditions, the representation and process of the olfactory in-

formation in the OB and the regulation of OB activity by the other brain regions have been extensively studied, and have greatly advanced our understanding of the olfactory system (6, 16–18). However, how the OB processes, represents, and transmits information of the same olfactory stimulus in different brain states has not been systematically investigated (19–21).

In this study, we examined the OB activities at rest and after stimulation in different brain states. We found that in two resting states with rather different baseline activities, the levels of peak activities of the same stimulation had no significant difference. This phenomenon was observed for both local field potential (LFP) and multiunit activity, independent of anesthetics, stimulating odorants, odorant concentrations, and recording sites in the OB. These results suggest that in the OB, the same peripheral event is represented by a similar population of neurons with similar roles and activity levels, independent of brain states. We speculate that this invariability of neural activity and representation for a specific event can ensure the peripheral olfactory information is sent to the higher olfactory cortices with high fidelity in different operational states of the brain.

Results

Baseline LFP Signals in the GCL. In the OB, the LFP signals in different frequencies (Fig. 1*A*) have different origins, amplitudes, and functions (22–25). The GCL contains a large amount of local neurons whose dendrites radiate almost exclusively toward the more superficial external plexiform layer, generating the strongest LFP signals in the OB (24, 25). Comparing the high brain state (HBS) with the low brain state (LBS) generated by chloral hydrate, the baseline activities for all three LFP bands are significantly increased by about 50% (Fig. 1*B*). Because the baseline activities and the increases in HBS are band-specific (Fig. 1*A* and Table S1 for detailed statistics), all LFP and multiunit signals are normalized against these in HBS to represent the neuronal activity changes in different brain conditions.

Effects of Brain States on GCL Responses. When the animal was stimulated with amyl acetate, the neuronal activity was enhanced in both LBS and HBS. Although the baseline activities are significantly different, the activity levels reached for a given frequency band in LBS and HBS are similar (Fig. 1*A*). The time courses show some band-specific features. For example, the highest frequency band has the smallest activity increase and slowest recovering kinetics (Fig. 1*B*). However, statistical analyses reveal that there is no significant difference between the maximum LFP signals in LBS and HBS for a given band (Fig. 1*C*

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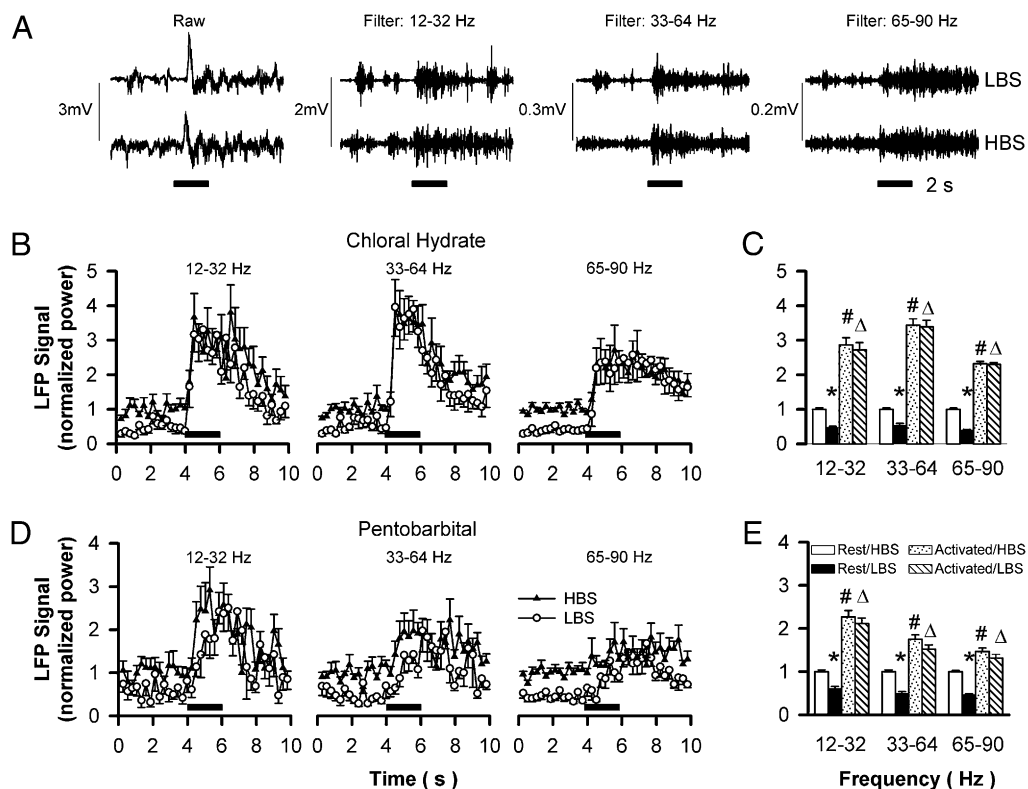


Fig. 1. LFP signals in the GCL under different brain states generated by different anesthetics. (A) LFP signals in three frequency bands were separated from the raw data in LBS and HBS (Upper and Lower). (B and D) The averaged time courses of LFP signals in the GCL for the three bands with chloral hydrate ($n = 12$) and pentobarbital ($n = 13$) as anesthetics, respectively. (C and E) Statistical analyses of the data in B and D, respectively. *HBS vs. LBS at rest; #rest vs. stimulated in HBS; Δ rest vs. stimulated in LBS; $P < 0.001$ for all comparisons; no significant differences between the peak activities in LBS and HBS for any bands any anesthetic.

and Table S1), indicating that peak neuronal activity of a given event is independent of brain state.

Effects of Anesthetics on GCL Responses. Different anesthetics exert differential effects via distinct mechanisms (26). To test whether the observed phenomenon above is anesthetic specific, we examined the LFP signals with a different anesthetic in the same OB layer. Chloral hydrate targets both glycine and GABA_A receptors, but pentobarbital targets only the GABA_A receptor and, furthermore, with different subunit compositions targeted by chloral hydrate (27). In the LBS and HBS generated by pentobarbital similar to those by chloral hydrate, some anesthetic-specific effects are revealed, including smaller activity increases and slower rising rates after odor stimulation for all frequency bands. However, the peak activities elicited by the same stimulus are not significantly different, as in the case of chloral hydrate (Fig. 1 D and E, and Table S1), indicating that the observed phenomenon above is independent of anesthetics.

Effects of Brain States on the LFP Signals and Representations of Different Odors. It is generally accepted that a given olfactory stimulation elicits a specific activity pattern in the OB (12, 13, 28). To test whether the observed phenomenon above is odor-specific, we examined the GCL responses to five other odors at nine recording sites. Although Fig. 2A shows odor-specific responses at one site during one exposure, statistical analyses revealed that the activity pattern of each odor, formed by the LFP signals across all sites, is odor-specific (Fig. 2B and Table S2), and that the activity pattern of each site, formed by the LFP signals of all odorants, is site-specific (Fig. S1 and Table S3). Interestingly, the activities in the LBS and HBS have no signif-

icant difference for a given odor at a given site, indicating that the phenomenon of brain-state-independent representation is not odor-specific (43 of 45 pairs) (Table S4). Because the activity at each point is highly similar under LBS and HBS, the corresponding two activity patterns of an odorant have no significant difference ($P < 0.005$ for all five odors, Fig. 2B and Table S3).

Effects of Odor Concentration on GCL Responses. Although the nominal concentration used above for all odorants is the same, 9.1% of the air stream flowing over pure odorants, the actual concentration is different because of the very different vapor pressures of these odorants. To assure that the observed phenomenon is not concentration-specific, LFP signal in the GCL elicited by different concentrations were recorded (Fig. 3). The signals of all LFP bands are significantly correlated with odor concentrations under both LBS and HBS ($P < 0.01$) (Table S5). More interestingly, the peak activities after stimulation under LBS and HBS in any of the frequency bands and concentrations are not significantly different ($P > 0.05$) (Table S5), indicating that the brain-state-independent representation is not concentration-specific.

Effects of Brain States on MCL Responses. Different bulbar layers have different cellular compositions and functions (11). The MCL contains M/T cell bodies and granular cell dendrites that are perpendicular to the layer, thus also generating strong LFP signals in both the LBS and HBS (Fig. 4A and B, Top). In the same conditions as in Fig. 1B, the time courses show that different bands had different activation rates (Fig. 4C). Although some layer-specific properties are revealed, including smaller activity increases, and slower rising and recovering rates in the

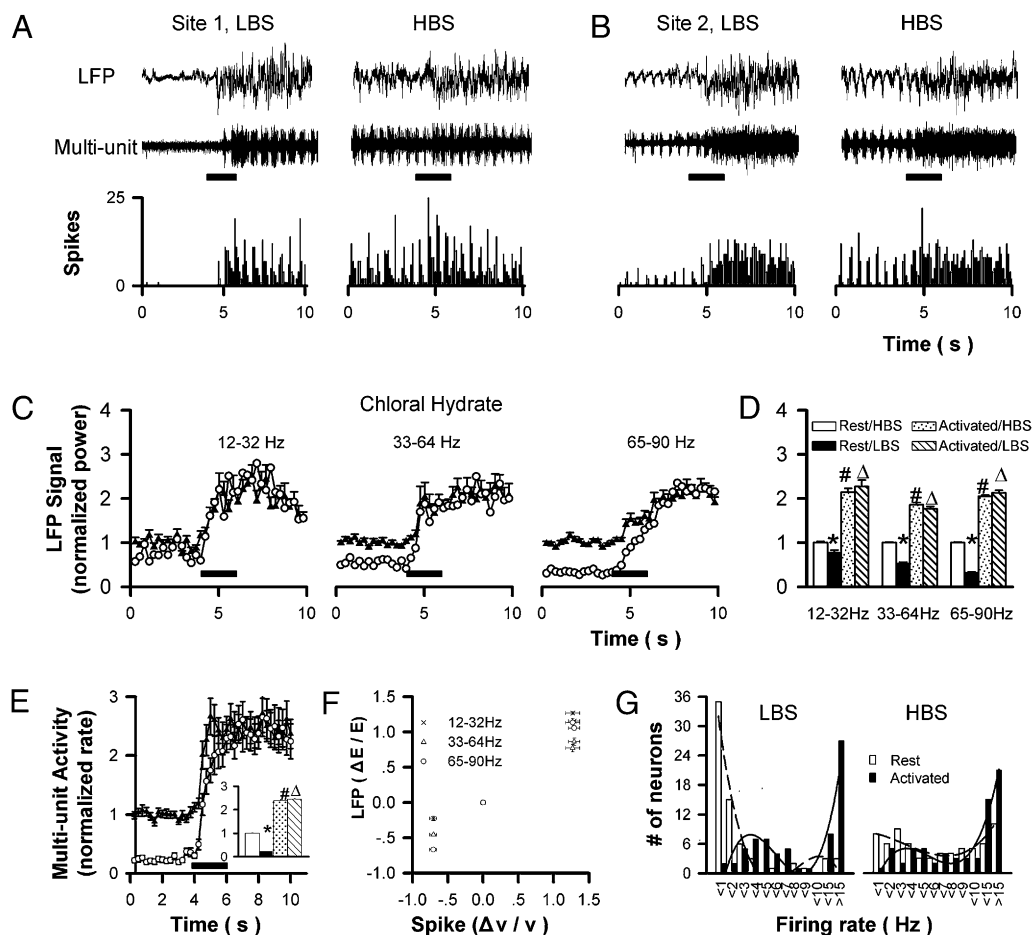


Fig. 4. Electrical recording in the MCL under LBS and HBS. (*A* and *B*) LFP signals (*Top*), multiunit spiking (*Middle*), and histogram of spiking (*Bottom*) at two recording sites with 2-s stimulation in LBS and HBS. (*C*) The averaged time courses of LFP signals in three bands (29 recording sites from 19 rats). (*D*) The statistical comparisons of data in *C*. (*E*) Multiunit signals and the statistical analyses. (*F*) Correlation between LFP and multiunit signals. (*G*) The distribution patterns of firing rate at rest and after stimulation in LBS and HBS, respectively; the last three bars were the number of neurons with firing frequency from 11 to 15, and > 15, and > 15, respectively. *, #, and Δ are the same as in Fig. 1; $P < 0.001$ for all situations; no significant difference between the peak activities in LBS and HBS for all three LFP bands and multiunit signals; circles and filled circles in *C* and *E* are LBS and HBS, respectively.

when the same visual stimulus is presented (29, 30). By using anesthetics to alter the baseline activity to a broader range, multiunit spiking rates in the primary somatosensory cortex during forepaw stimulation are also independent of the baseline levels (31–33). The state-independent total neuronal activity described above, however, is not always observed, as the responses under deeper anesthesia are stronger in many reports (33–36), but weaker in some others (37–39). These differences might be a result of differences in experimental conditions (anesthetics, dosages, animals, and so forth), or studied regions (primary or higher cortices in the sensory systems).

It has been reported that in the aspects of LFP signals and spiking rates, task-related learning processes can modify the neural representation (40, 41). However, the objective modulation does not contradict the results in this study, because the task-related modified representations are also stable after the animals learned the task (40). Therefore, the modification just redefines the stimulus to generate a new representation and perception accordingly, conferring the system with higher efficiency for complex olfactory functions. It will be interesting to test whether the modified representation after task-related learning is also brain-state-independent (e.g., fed vs. fasted).

Neuronal Basis for the Larger Increment at LBS in the OB. The state-independent neuronal activity and representation for a given

stimulus means that the increment is larger at LBS. The peripheral and centrifugal inputs and the intrinsic circuits together decide the OB activity. Although the effect of anesthesia on the peripheral inputs is negligible (42), it is more significant for the central processes (1, 19). The OB receives modulatory inputs from several nuclei (16–18, 43) and heavy feedback inputs from many cortices (11, 43, 44). Because multiple receptor types of these pathways spread over the OB, the effect of anesthesia on modulatory inputs is difficult to predict. However, with glutamate as a neurotransmitter, the feedback inputs are suppressed by anesthesia (6). Because their targets are the granular cells, the LBS will disinhibit the dendrodendritic network in the external plexiform layer, enhancing the activity increment.

Energetic Basis for the Enhanced Responses at LBS. The human brain makes up 2% of body mass, but consumes 20% of the cardiac output in adults and even more in babies (45, 46). Although blood supply to a given brain region can increase significantly during activation (3, 47, 48), the total amount to the brain is increased only slightly, even when the brain is performing difficult tasks (45, 46, 49). Therefore, even in a “rest” state, the brain needs large amounts of energy, close to its maximum quota. As revealed by comprehensive magnetic resonance spectroscopy studies, these energies are predominantly (>80%) used for neural activity (50, 51). The constraint of global energy avail-

ability in the brain would limit the total brain activity and determine how the brain performs its functions, as shown in ever-increasing studies that the two major systems in the brain for internal and external worlds cannot be activated simultaneously (52, 53). At the level of individual neurons, the same olfactory stimulation elicited stronger responses from M/T cells in anesthetized rats (19). This so-called “sparse coding” in the awake state, a mode taking advantage of timing to save energy, has been observed in several sensory systems (54, 55). In LBS, inhibition is stronger and the energy restraint is weaker, and the enhanced activity increment might be a balanced result. The energy restriction, combined with the interactions among the involved neuro-circuits discussed above, could lead to the observed larger increment in enhanced responses under LBS in the early sensory stages, although additional mechanisms are also likely involved.

Similarity of Activity Patterns Under Different Brain States. For the same olfactory stimulus, the activity patterns under LBS and HBS are highly similar on the levels of both LFP, which reflects the activity of many neurons close to the electrodes, and multiunit spiking, which reflects the activity of a few individual neurons (Figs. 2*B* and 4*G*). Furthermore, the LFP and multiunit spiking signals are well correlated, as shown in Fig. 4*F*. These coherent results indicate that in different brain states, the neurons in the OB activated by a given stimulus consist of similar populations with similar activities. In other words, under different brain states the same information is processed similarly in the OB, and the information sent to high olfactory cortices is also highly similar, as indicated by the highly similar patterns of multiunit activity (11, 15, 28, 56). For the same odor at different concentrations under the same brain state, previous studies using a variety of methods have revealed that their patterns have similar topography, but with concentration-correlated intensities (13, 57–60), leading to the speculation that the pattern topography represents stimulation quality and the intensity represents strength (13). The results in this study revealed that the intensity not only correlated with concentration (Fig. 3), but can vary with brain states. However, the total activity for a given stimulus is invariable in different brain states, consistent with the speculation that the total activity represents the event itself (61). However, the neurochemical and neurophysiological basis for these observations are barely known. Revelation of these mechanisms will be helpful in elucidating how the OB processes peripheral information and presents the processed information to higher stages.

Summary. The sensory systems should provide accurate information about the environment to the brain, and the brain has to form accurate perceptions from the input sensory information, independent of brain states. Our results showed that in the OB, the total activity elicited by the same odor stimulation was independent of global brain states, suggesting that the OB requires total, not incremental, neuronal activity to code, process, and convey the related sensory information. Furthermore, the activity patterns elicited by the same stimulus under different brain states were highly similar at both LFP and spiking levels, suggesting that similar populations of neurons are involved, each with similar roles and firing properties. This brain-state-independent neural activity and representation might be the mechanisms for the OB to present the peripheral olfactory information with high fidelity to the higher brain centers under ever-changing operational states.

Materials and Methods

Animal Surgery. All animal procedures were approved by the Chinese Academy of Sciences. Adult Sprague-Dawley rats (200–300 g) were anesthetized with intraperitoneal chloral hydrate (35 mg/kg), or pentobarbital (1 mg/kg). The rat was on a heat blanket and the core temperature was maintained at a constant (36 ± 1 °C) throughout the studies. Respiration was monitored with a piezoelectric device recording the chest wall movements.

Manipulation of Brain States. To reveal how the OB responds to odor stimulation under different brain states, we had to generate two levels of baseline activity reliably. Anesthetics, broadly used to alter global brain activity and baseline metabolism (26), are suitable for this purpose. After animal surgery and the placement of electrodes were finished, a supplementary dosage (30% of the original) was applied so both LFP and EEG recordings showed iso-electrical patterns (Fig. S2*A*). This deep anesthesia level was defined as LBS. The anesthesia level became lighter with time, so both LFP and EEG recordings showed more high-frequency signals (Fig. S2*B*). The lighter anesthesia level, at which the total LFP power was increased by $100 \pm 10\%$ compared with LBS, was defined as HBS. The time required to reach HBS was animal-dependent, but it generally took about 45 min. For pentobarbital, supplementary dosage (30% of the original) was applied to induce LBS. The HBS, which also had a $100\% \pm 10\%$ increase of LFP signal compared with LBS, was generally reached about 30 min later.

Odor and Its Delivery. To reveal the representation of different odors in the OB under different brain states, amyl acetate, ethyl acetate, ethyl butyrate, ethyl valerate, heptanal, and 2-heptanone were selected based on the facts that their specific activity patterns overlap significantly in the general regions (where recordings were made in this article) in the middle-posterior area of medial side or the middle area of the lateral side of the OB (<http://gara.bio.uci.edu/odorants.jsp?start=1&end=1&action=null> and <http://senselab.med.yale.edu/OdorMapDB/search.aspx> for 2-deoxyglucose and functional MRI, respectively). The odorants were delivered to freely breathing animals by passing a stream of charcoal-filtered, humidified air over pure odorants, then diluted to one-tenth by an olfactometer. Different dilution ratios were used for concentration effects. Odor stimulation, triggered by the recording program, lasted for 2 s with an interstimulation interval >100 s to prevent habituation and a flow rate of 1 L/min.

LFP and Multiunit Recordings. Electrophysiological signals were used as an indicator of neuronal activity. The rat was placed prone on a stereotaxic holder on a vibration-free table inside a Faraday cage. After the skull over the OB was exposed and covered with mineral oil to prevent drying, micro-electrode (~ 1 - μm tip diameter tungsten with impedance of 2–4 M Ω ; FHC) was inserted using a stereotaxic micromanipulator (Stoelting) to the GCL or MCL for recording. Within the general recording regions (± 1.5 mm lateral to midline, -7.0 mm anterior to Bregma), responses to all selected odorants can be readily obtained. The recording site was identified by histology after the experiments (SI Text and Fig. S3). A steel screw was fixed on the parietal lobe (± 2.5 mm from midline, 4 mm posterior to Bregma) to record EEG signals. The LFP and EEG signals were amplified ($\times 2,000$; Dagan) before digitization ($\mu\text{-}1401$; CED). The raw recording data were separated into LFP and multiunit signals (0.1–300 Hz and 300–3,000 Hz, respectively). At each recording site for each brain condition, four to six repeats were performed.

Data Process. Off-line analysis (Spike-2; CED) was used to create a spike template that was then used to create a temporal history of the ensemble by converting to spiking frequency (ν) with 0.256-s bins. The datasets in which excitatory multiunit spiking was not identified after odor stimulation under deep anesthesia were excluded for further data analysis. Spike analysis from 29 recording sites identified 69 individual neurons in the neuronal ensemble (SI Text). Raw data 4 s before and 6 s after the onset of the odor stimulus were selected for the processing of both LFP and multiunit signals, with the former period as the baseline and the latter as the activated. These 10-s data were binned with a width of 0.256 s (512 sample points for LFP). Time courses of LFP, multiunit, or spiking rate were obtained for each recording. Spectrum analysis and calculation of the spectrum power of LFP and the count of multiunit spikes were both performed using spike software (Spike-2; CED). After time-frequency transformation analysis of the LFP, three frequency bands of LFP, 12 to 32 Hz, 33 to 64 Hz, and 65 to 90 Hz, were filtered for further process. Student *t* test comparisons between resting and stimulated data were used to calculate significance of change. Pearson correlation coefficients are used to reflect the relationship between the activities obtained under different conditions, including recording sites, odorants, concentrations, brain conditions, and LFP and multiunit signals.

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