Delta oscillations phase limit neural activity during sevoflurane anesthesia

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Understanding anesthetic mechanisms with the goal of producing anesthetic states with limited systemic side effects is a major objective of neuroscience research in anesthesiology. Coherent frontal alpha oscillations have been postulated as a mechanism of sevoflurane general anesthesia. This postulate remains unproven. Therefore, we performed a single-site, randomized, cross-over, high-density electroencephalogram study of sevoflurane and sevo-flurane-plus-ketamine general anesthesia in 12 healthy subjects. Data were analyzed with multitaper spectral, global coherence, cross-frequency coupling, and phase-dependent methods. Our results suggest that coherent alpha oscillations are not fundamental for maintaining sevoflurane general anesthesia. Taken together, our results suggest that sub-anesthetic and general anesthetic sevoflurane brain states emerge from impaired information processing instantiated by a delta-higher frequency phase-amplitude coupling syntax. These results provide fundamental new insights into the neural circuit mechanisms of sevoflurane anesthesia and suggest that anesthetic states may be produced by extracranial perturbations that cause delta-higher frequency phase-amplitude interactions.
General anesthesia is a drug-induced reversible state of unconsciousness, amnesia, antinociception, and immobility, with the maintenance of physiological stability. Since the first public demonstration of diethyl ether as a general anesthetic occurred at the Massachusetts General Hospital (MGH) in 1846, the practice of anesthesiology has revolutionized surgery and spurred advances in modern medicine. Now, over 170 years later, the administration of diethyl ether derivatives remains fundamental to anesthesiology. Fluoromethyl hexafluoroisopropyl ether, also known as sevoflurane, is a derivative of diethyl ether that is extensively used in current clinical practice. Basic science studies have helped to identify the molecular targets of sevoflurane. These targets include γ-aminobutyric acid type A (GABA<sub>A</sub>) receptors, glycine receptors, two-pore potassium channels, and N-methyl-D-aspartate receptors, among many others. However, a significant knowledge gap exists in relating how drug activity at molecular targets results in the neural circuit perturbations that cause anesthetic states.

Neural oscillations are a prominent feature of the anesthetized brain. These oscillations change systematically as a function of anesthetic drug dose and age, which suggests that they are generated from mechanisms that depend on cellular properties such as ionic currents, myelin integrity, and synaptic density. Neural oscillations are important for information processing in the brain. Thus, anesthetic-drug-induced oscillations may disrupt the brain’s capacity for information processing in cognitive, memory, and sensory circuits. Coherent frontal alpha oscillations—an oscillatory dynamic associated with sevoflurane—and the other modern-day derivatives of ether anesthesia—have been postulated as a fundamental mechanism by which the derivatives of diethyl ether cause unconsciousness. This postulate remains unproven, and many sevoflurane general anesthesia scenarios without coherent alpha oscillations have been identified. For example, alpha oscillations are not evident in the electroencephalogram (EEG) of patients until approximately 4 months of age and do not become coherent until approximately 10 months of age. Thus, the neural circuit mechanisms of sevoflurane general anesthesia are not clear.

To formally evaluate neural circuit mechanisms underlying sevoflurane general anesthesia, we conducted a study of sevoflurane anesthetic states in healthy human volunteers. Human neurophysiological studies of anesthetic-drug-induced states have typically used the loss of behavioral responsiveness to nonnal stimuli, such as auditory cues, as a shorthand for unconsciousness. This experimentally convenient approach may engender considerable confusion because the loss of responsiveness occurs during subanesthetic states that do not approximate general anesthesia. Further, states of internal processing such as dreaming during subanesthetic states may persist during subanesthetic states. We aimed to obtain fundamental new insights into anesthetic mechanisms of sevoflurane by targeting and studying subanesthetic and general anesthetic brain states using sevoflurane anesthetic concentrations that are consistent with current clinical practice and epidemiologically based characterizations.

Sevoflurane-induced lack of responsiveness is associated with loss of occipitally dominant alpha oscillations prior to the occurrence of frontal alpha oscillations. Here, we investigated neural circuit dynamics to explain sevoflurane-induced subanesthetic, general anesthetic, and deep general anesthetic brain states using high-density EEG recordings. We have previously suggested that ketamine, an anesthetic drug that is typically administered as an adjunct for antinociception during general anesthesia, is associated with decreased alpha oscillation power and coherence. Therefore, in a separate experiment, we specifically tested the hypothesis that coherent alpha oscillations are fundamental for sevoflurane general anesthesia by studying the effects of ketamine on sevoflurane-induced oscillations. Previously described neural correlates of propofol general anesthesia are based on the amplitude (power), frequency-dependent correlation, and phase—amplitude dynamics of neural oscillations. Therefore, we implemented multivariate spectral analysis to characterize the amplitude (power), global coherence analysis to characterize frequency-dependent correlation structure, and nonlinear cross-frequency coupling analyses to characterize the phase—amplitude dynamics of sevoflurane-induced oscillations. We also aimed to study whether dexmedetomidine, an anesthetic drug that modulates the alpha-2a receptor, shares neural correlates that approximate those of sevoflurane.

Results
No adverse events were reported during this study. Figure 1a is the schematic of our randomized, cross-over study design. Figure 1b–e illustrates data obtained from a subject during sevoflurane and sevoflurane-plus-ketamine study visits. In this manuscript, we use the term anesthetic depth to refer to the increased level of brain hyperpolarization associated with increased anesthetic drug administration.

Loss of responsiveness to behavioral stimuli is not synonymous with general anesthesia. We aligned the data (±10 min) to the probabilistic definition of LOR (loss of responsiveness) and ROR (return of responsiveness) (Supplementary Fig. 1). The end-tidal sevoflurane concentration at which LOR occurred was 1% (SD, 0.1). ROR occurred at 0.2% (0.1) (Supplementary Fig. 1a–d). During the sevoflurane-plus-ketamine visit in which sevoflurane was rapidly administered, the end-tidal sevoflurane concentration at which LOR occurred was 1.8% (0.2). ROR occurred at 0.2% (0.1) (Supplementary Fig. 1e–h). EEG oscillations during LOR were inconsistent with dynamics that have previously been described for sevoflurane general anesthesia.

Alpha oscillation power did not covary with anesthetic depth. We analyzed the neural oscillations associated with awake (baseline), subanesthetic (1.1% sevoflurane), anesthetic (2.1% sevoflurane), deep anesthetic (2.8% sevoflurane), and awake (emergence) states. We computed frontal spectrograms (Fig. 2a), power within canonical frequency bands of interest (Fig. 2b), and spectra differences (Fig. 2c). We also computed occipital spectrograms (Supplementary Fig. 2a), spectra differences (Supplementary Fig. 2b), and power spectral-spatial plots (Supplementary Fig. 2c).

When we analyzed data from frontal electrodes, we found that alpha oscillations did not covary with increasing anesthetic depth, such that EEG power was increased during the anesthetic state but decreased during the deep anesthetic state. However, slow oscillation power covaried with anesthetic depth. We implemented a noncanonical frequency band-based bootstrap procedure for statistical inference. These data are summarized in Fig. 2c. Data from occipital electrodes are summarized in Supplementary Fig. 2b.

Globally coherent alpha oscillation did not covary with anesthetic depth. Increased global coherence may occur despite decreases in oscillatory power. Therefore, we next analyzed the global coherence structure of the baseline, subanesthetic, anesthetic, deep anesthetic, and awake states. We computed global coherograms (Fig. 2d), global coherence within canonical frequency bands of interest (Fig. 2e), and global coherence differences (Fig. 2f). We also computed global coherence spatial plots (Supplementary Fig. 2d). We found that the anesthetic states were associated with globally coherent theta and alpha oscillations. However, these globally coherent oscillations did not covary with anesthetic depth such that coherence was increased during the
Ketamine reduced globally coherent oscillations during sevo-Ketamine induced relatively active electroencephalogram oscillations during sevo-flurane general anesthesia. Next, we tested the hypothesis that coherent alpha oscillations are fundamental for sevo-flurane general anesthesia by studying the effects of ketamine on sevo-flurane-induced oscillations. We analyzed the neural oscillations associated with awake (baseline), anesthetic (2.1% sevo-flurane), deep anesthetic (2.1% sevo-flurane plus ketamine; SevoKet), and awake (emergence) states. We computed frontal spectrograms (Fig. 3a), power within canonical frequency bands of interest (Fig. 3b), and spectra differences (Fig. 3c). We also computed occipital spectrograms (Supplementary Fig. 3a), spectra differences (Supplementary Fig. 3b), and power spectral-spatial plots (Supplementary Fig. 3c).

When we analyzed data from frontal electrodes, we found that delta oscillation power was decreased during the SevoKet deep anesthetic state. The SevoKet deep anesthetic state was associated with a paradoxical increase in beta oscillation power. These data are also summarized in Fig. 3c. Data from occipital electrodes are summarized in Supplementary Fig. 3b.

Ketamine reduced globally coherent oscillations during sevo-flurane general anesthesia. We next analyzed the effect of ketamine on the global coherence structure of the baseline, anesthetic, SevoKet deep anesthetic, and awake states. We computed global coherograms (Fig. 3d), global coherence within canonical frequency bands of interest (Fig. 3e), and spectra differences (Fig. 3f). We also computed global coherence spatial plots (Supplementary Fig. 3d). We found that the SevoKet deep anesthetic state was associated with decreased globally coherent theta and alpha oscillations. Thus, globally coherent oscillations did not covary with anesthetic depth. These data are summarized in Fig. 3f. Supplementary Movie 2 illustrates the oscillatory and global coherence changes associated with sevo-flurane anesthetic states.

Sevo-flurane anesthetic states were associated with a phase—amplitude coupling syntax. The power and global coherence of sevo-flurane-induced oscillations did not track with anesthetic depth. Therefore, we next investigated whether anesthetic states emerge from impaired information processing instantiated by cross-frequency coupling. First, we computed comodulograms to elicit the putative phase driver(s) associated with sevo-flurane anesthesia states. The comodulogram plots demonstrated that delta oscillation indices were highest during sevo-flurane anesthetic states (Fig. 4a). The spatial representation of comodulograms is presented in Supplementary Fig. 4a. We next analyzed phase—amplitude coupling dynamics between delta and higher frequency oscillations by computing phaseamplograms (Fig. 4b). We observed that the relative amplitude to higher-frequency oscillations was restricted to the $\pm \pi/3$ phase of delta oscillations during the subanesthetic state. The relative amplitude of oscillations $<8$ Hz remained restricted to the $\pm \pi$ phase of delta oscillations during sevo-flurane general anesthesia. However, oscillations $>8$ Hz were restricted to approximately $-\pi/3$
to \(\pi/3\) phase of delta oscillations. The relative amplitude of oscillations <6 Hz remained restricted to the \(\pm \pi/3\) phase of delta oscillations during the deep general anesthetic state. However, oscillations >6 Hz were restricted to approximately \(-\pi/3\) to \(\pi/3\) phase of delta oscillations.

To enable statistical inference, we computed circular phasor plots (Fig. 4c; top panel) for phase driver (2–4 Hz) and higher frequency oscillations (13–20 Hz). We observed that the mean phase angle shifted from approximately \(\pi\) phase towards \(\pi/4\) phase as a function of anesthetic depth. Alpha oscillation power did not covary with the anesthetic state. During the sevoflurane anesthesia states, the distribution of the circular mean of the mean amplitude vectors (Fig. 4c; bottom panel) was not uniformly distributed during the sevoflurane anesthesia states (baseline, \(p = 0.259\); subanesthetic, \(p = 0.011\); general anesthetic, \(p = 0.006\); deep general anesthetic, \(p = 0.011\); emergence, \(p =\)).
We also observed that the mean amplitude vector for delta oscillation phase became larger as a function of anesthetic depth (subanesthetic > baseline, \( p = 0.104 \); general anesthetic > subanesthetic, \( p = 0.0171 \); linear mixed-effects model). The anesthetic and deep anesthetic comparison was not significantly different (\( p = 0.1367 \)).

We also computed the spatial distribution of phaseamplograms (Supplementary Fig. 5a) and circular phasor plots and mean amplitude histograms of slow and delta driver frequencies (Supplementary Fig. 5b, c). Worse signal to noise ratio can be observed in the mean amplitude distribution of the slow driver.

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**Fig. 3** Spectral and global coherence analysis of the sevoflurane-plus-ketamine study visit. a Ketamine reduced the power of alpha oscillations during sevoflurane general anesthesia. b Power in the canonical slow-delta and alpha frequency bands change with respect to the anesthetic states. c Frontal spectra and bootstrapped difference of median spectra confirm that the ketamine-induced alpha oscillation power decrease was significant. Ketamine was also associated with a decrease in delta oscillation power and an increase in beta oscillation power. d The median global coherograms demonstrate ketamine reduced the global coherence of theta and alpha oscillations. e Global coherence in the canonical theta and alpha frequency bands change with respect to the anesthetic states. f Global coherence spectra and bootstrapped difference of median global coherence confirm that ketamine significantly reduced theta and alpha global coherence to suggest this dynamic is not fundamental for general anesthesia. Shaded regions represent the 99% confidence bounds of the bootstrapped median global coherence spectra. Black lines represent frequency bands that met our threshold for statistical significance.
Sevoflurane phase–amplitude coupling syntax tracks with the anesthetic state. We previously demonstrated that sevoflurane plus ketamine general anesthesia is associated with relatively activated EEG oscillations. Therefore, we computed comodulograms to assess whether the putative neural phase–amplitude syntax that we observed during sevoflurane general anesthetic states was conserved after the administration of ketamine. The comodulogram plots demonstrated that delta oscillation modulation indices were highest during sevoflurane general anesthetic states (Fig. 4d). Spatial representation of comodulograms is presented in Supplementary Fig. 4b. We analyzed phase–amplitude coupling dynamics between delta and higher frequency oscillations by computing phaseampograms (Fig. 4e). Similar to the findings described above, the relative amplitude of oscillations <8 Hz was restricted to the ±π phase of delta
oscillations during sevoflurane general anesthesia. However, oscillations >8 Hz were restricted to approximately $-\pi/3$ to $\pi/3$ phase of delta oscillations. Similar to findings described for the deep anesthetic state during sevoflurane visit, the relative amplitude of oscillations <6 Hz remained restricted to the $\pi$ phase of delta oscillations while oscillations >8 Hz were restricted to approximately $-\pi/3$ to $\pi/3$ phase of delta oscillations during the SevoKet deep general anesthesia state.

We also computed circular phasor plots (Fig. 4f; top panel) for phase driver (2–4 Hz) and higher frequency oscillation (13–20 Hz). We observed that the mean phase angle shifted from approximately $\pi$ phase towards $\pi/4$ phase as a function of anesthetic depth (baseline at $3\pi/5$; anesthetic at $11\pi/20$; deep anesthetic at $2\pi/5$; emergence at $3\pi/5$). The distribution of the circular mean of the mean amplitude vectors (Fig. 4f; bottom panel) was not uniformly distributed during the sevoflurane anesthesia states (baseline, $p = 0.645$; general anesthetic, $p = 0.006$; sevoKet deep general anesthetic, $p = 0.012$; emergence, $p = 0.527$; omnibus test). We also observed that the mean amplitude vector for delta oscillation phase became larger as a function of anesthetic depth. The median amplitude vector (red line) was increased from baseline during the anesthetic states. Mean amplitude distribution was not uniformly distributed during sevoflurane-induced anesthetic states. Circular phasor plots also demonstrated that neural activity systematically shifted from $\pi$ towards 0 phase of delta oscillations as a function of anesthetic depth. The median amplitude vector (red line) was increased from baseline during the anesthetic states. Mean amplitude distribution was not uniformly distributed during sevoflurane-induced anesthetic states. Colored circular dots on phasor plots represent subject level data. Error bars represent standard deviation.

**Discussion**

Our findings demonstrate that sevoflurane sedation, a subanesthetic state from which patients can be aroused to consciousness, is associated with phase restricted activity of neural oscillations to the trough ($\pi$) region of delta oscillations. During sevoflurane general anesthesia states, the phase-restricted activity of $-0.1$–6 Hz oscillations to the trough region was maintained. However, $-8$–$30$ Hz frequency oscillations became restricted to the peak ($-\pi/3$ to $\pi/3$) region of delta oscillations. This dynamic was maintained across all electrode sensor locations, although modulation indices were strongest in frontal electrode locations. The magnitude and phase of the mean normalized phase–amplitude coupling vector, not the power or global coherence of oscillations, tracked with anesthetic depth. Taken together, our results provide strong evidence that subanesthetic and general anesthetic brain states emerge from impaired information processing instantiated by a delta-higher frequency phase–amplitude coupling syntaxis.

At the neuronal level, slow oscillations are associated with an alternation between Up states where neurons are able to fire, and Down states where neurons are relatively silent$^{26,27}$. General anesthesia is associated with longer Down states in humans$^{26}$. The ionic currents underlying cortically generated delta oscillations are similar to those underlying the slow oscillation Up and Down states$^{28}$. Neural activity during the Down state may result from intrinsic and synaptic properties of pyramidal cells in layer 5 such as the summation of spontaneous action potential independent excitatory synaptic potentials$^{29–32}$ or from neurons that fire persistently during the Down state$^{33–36}$. Neural activity during the Up state may result from enhanced activity of inhibitory interneurons and activation of activity-dependent hyperpolarizing currents$^{37–39}$. Thus, the phase–amplitude coupling dynamics we describe suggest that membrane conductance and timescales governing well-timed neuronal activity are markedly prolonged by sevoflurane. Thus, the repertoire of possible neuronal firing and coupling dynamics are restricted to a rigid structure during anesthetic states. Neurophysiological recordings across various spatial scales are necessary to precisely define the micro-circuit dynamics underlying our findings.

A unifying mechanism to explain how anesthetic drugs from various drug classes produce general anesthesia may not be achievable. Theories of information processing propose that phase–amplitude coupling is critical for regulating neuronal activity and information processing across spatial and temporal scales$^{17,18,40,41}$. Phase–amplitude coupling between slow oscillations and thalamocortical alpha oscillations was recently described for propofol anesthesia$^{16}$. We also show that dexmedetomidine, an alpha-2a receptor agonist that patterns the activity of various arousal nuclei similar to sleep$^3$, is associated with phase–amplitude coupling between slow and spindle oscillations. Thus, although the center frequency of phase drivers may change as a function of the anesthetic drug class or drug dose, we speculate that phase–amplitude coupling may be consistent across anesthetic states produced by various anesthetic drug
classes. While our present work supports this conclusion, further research is required to more thoroughly explore this hypothesis.

Although functional imaging studies are not a surrogate for neuronal spiking activity, they provide a measure of the collective behavior of neurons. Functional imaging studies of subanesthetic sevoflurane and subanesthetic dexmedetomidine have previously demonstrated that cortico-cortical connectivity between brain regions that comprise the Default Mode Network is preserved during these states. Thus, the phase—amplitude pattern we describe during the subanesthetic state may reflect functionally coupled cortical neural activity that is primed for the response to external stimuli. Conversely, sevoflurane general anesthesia is associated with impaired cortico-cortical connectivity between brain regions that comprise the Default Mode Network. Thus, the phase—amplitude pattern we describe during this state may reflect functionally decoupled neural activity that is not receptive to external stimuli. We note that the phase—amplitude syntax we describe is consistent with asynchronous and hypersynchronous low-frequency oscillations causing altered arousal states. Further, our findings of decreased phase—amplitude coupling modulation indicate that anesthetic states may be produced by extracranial perturbations such as direct current stimulations that cause delta-higher frequency phase—amplitude interactions. Subcortical sources of low-frequency oscillations such as the thalamus is not shown higher frequency phase—amplitude interactions. The feasibility of this approach is supported by a recent report that used extracranial stimulations to produce theta-gamma phase—amplitude coupling and improved working memory in humans. This potentially non-drug-mediated approach may fundamentally advance anesthetic care by eliminating cardiovascular and respiratory morbidity that result from off-target drug binding.

Methods

The Partners Institutional Review Board approved this human research study. This study was registered on www.ClinicalTrials.gov (Identifier: NCT03503578).

Subject recruitment. This was a single-site, randomized, cross-over study conducted in healthy subjects. Subjects underwent a complete medical history and a pre-anesthesia assessment. The primary inclusion criterion was meeting American Society of Anesthesiology Physical Status I. Key criteria for exclusion were pregnancy, personal or family history of anesthesia-related complications, suspected history of drug abuse, and neuropsychiatric diagnoses. We performed the following screening laboratory tests: complete blood count, liver function, basic metabolic panel, urine toxicology, and urine pregnancy for females. We also recorded data from a 12-lead electrocardiogram. We obtained written informed consent from 12 subjects (7 males), mean age 25 (SD ± 4.7) years, mean weight 70 (11) kg, and mean BMI 24.1 (3) kg/m². We also reanalyzed data from a previously reported study of dexmedetomidine plus-ketamine anesthesia. Briefly, dexmedetomidine was administered intravenously as a bolus of 1 μg kg⁻¹ over 10 min followed by a 0.7 μg kg⁻¹ h⁻¹ infusion for 50 min to healthy volunteers, 18–36 years of age. The EEG data analyzed were obtained during the 0.7 μg kg⁻¹ h⁻¹ maintenance phase of dexmedetomidine.

Data acquisition. All study procedures were conducted at MGH, Boston, MA. We induced and allowed recovery from sevoflurane general anesthesia and sevoflurane—plus-ketamine general anesthesia in 12 subjects using a cross-over study design (Fig. 1a). Thus, each subject received both sevoflurane—induced general anesthesia (Fig. 1b, c) and sevoflurane—plus-ketamine—induced general anesthesia (Fig. 1d, e) on study visits separated by at least 48 h. Subjects were required to avoid food and water intake for at least 8 h prior to study onset. We reperformed urine toxicology screening, and urine pregnancy testing prior to the administration of sevoflurane. We monitored blood pressure using a standard noninvasive cuff, oxygen saturation using pulse oximetry, ventilation using capnography, and circulation using a 5-lead electrocardiogram.

We instructed all subjects to close their eyes throughout the data acquisition period. Eye closure reduces blinks, muscle artifacts and helps to distinguish between awake occipital alpha oscillations and sevoflurane—induced alpha oscillations. A tight facemask was applied, and subjects were acclimated to spontaneously breathing through the tight face mask before initiation of the study. We administered sevoflurane using the Dräger Fabius Tiro (Telford, PA, USA) machine. The sevoflurane concentration was analyzed using a General Electric standard multigas module analyzer in clinical use at our institution. During the sevoflurane—induced general anesthesia visit, after 10 min of baseline (awake) recordings, we increased the
end-tidal sevoflurane concentration in a stepwise fashion to subanesthetic (1.1%),
general anesthetic (2.1%), and deep general anesthetic (2.8%) states. Each
concentration level was maintained for 15 min. During the sevoflurane-plus-
ketamine-induced general anesthesia visit, after 10 min of baseline (awake) recordings,
we increased the sevoflurane end-tidal concentration to a general anesthetic (2.1%) state and
maintained the anesthetic concentration for 45 min. We administered an intravenous bolus of ketamine (0.75 mg·kg−1) after achieving 15 min steady-state
sevoflurane concentration. Additionally, we recorded 10 min of emergence (awake)
EEG data during both study visits. Two board-certified anesthesiologists were present
during all study procedures.

We recorded high-density EEG signals using the Waveguard system with a
standard 24 channels, ANT Neuro, Netherlands) and electrode impedances of <5 kΩ. We instructed subjects to click a mouse button when they
heard auditory stimuli. The auditory stimuli were either a verbal command to press
a button or auditory steady-state responses (40 and 80 Hz). We randomly
presented auditory stimuli every 4–8 s. All auditory stimuli were 1 s long and
were delivered using headphones (ER2; Etymotic Research).

Data preprocessing and epoch selection. We down-sampled the EEG data to 250
Hz, spline-interpolated corrupted data, and remontaged the data using a nearest-
neighbor Laplacian referencing scheme. For the sevoflurane visit, we selected 5-min
EEG epochs. These EEG epochs were selected 10 min after the sevoflurane con-
centration reached the desired steady-state concentrations of 1.1, 2.1, and 2.8%. For the
sevoflurane-plus-ketamine visit, we selected 5-min EEG epochs. These EEG epochs
were selected 10 min after sevoflurane reached the desired steady-state con-
centration of 2.1% and 2 min after the ketamine bolus was administered. We also
selected 5-min EEG epochs during baseline and emergence periods.

For spectral analysis, we averaged data from three channels to approximate frontal,
temporal, and occipital location corresponding to F3, F4, and O1, respectively. We also averaged data from the
channels to approximate occipital channel location corresponding to Oz. Two
investigators visually inspected the selected EEG data to ensure artifact-free epochs.

Probability of response analysis. Button press responses to auditory stimuli were
binarized. We computed subject level probability (response) curves from the binary
response data with a Bayesian state-space algorithm52. We defined the loss of
responsiveness (LOR) as the probability (response) <0.05 after the administration
of sevoflurane and if maintained for at least 5 min. We defined the return
of responsiveness (ROR) as the probability (response) >0.05 after the discontinuation
of sevoflurane and if maintained for at least 5 min.

Spectral analysis and spatial plots. We used the Chronux toolbox in Matlab
2018 (Mathworks, Natick, MA) to compute multiplexer spectral estimates. The
parametric and nonparametric tests were carried out by the Chronux toolbox.
We down-sampled the EEG data to 250 Hz, spline-interpolated corrupted data, and remontaged the data using a nearest-
neighbor Laplacian referencing scheme. For the sevoflurane visit, we selected 5-min
EEG epochs. These EEG epochs were selected 10 min after the sevoflurane con-
centration reached the desired steady-state concentrations of 1.1, 2.1, and 2.8%. For the
sevoflurane-plus-ketamine visit, we selected 5-min EEG epochs. These EEG epochs
were selected 10 min after sevoflurane reached the desired steady-state con-
centration of 2.1% and 2 min after the ketamine bolus was administered. We also
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For spectral analysis, we averaged data from three channels to approximate frontal,
temporal, and occipital location corresponding to F3, F4, and O1, respectively. We also averaged data from the
channels to approximate occipital channel location corresponding to Oz. Two
investigators visually inspected the selected EEG data to ensure artifact-free epochs.

Global coherence analysis and spatial plots. Global coherence is a multivariate
measure of synchrony. We estimated global coherence for all electrode locations
except the bilateral mastoids and electroc心ogram. First, we computed the
cross-spectral matrix for each subject over each nonoverlapping moving
window using multliplexer methods. Next, we computed the median of the
global coherence estimates across all subjects for group-level visualization.
We also computed the median of the group averaged spectral estimates
for each frequency band of interest. The topoplot function in EEGlab
was used to interpolate these data and to generate spatial head plots.

Modulation index. The modulation index, an adaptation of the Kulbacak–Leiber
distance, is a scalar measure of phase–amplitude coupling between two frequency
ranges of interest: phase modulating (phase driver) and amplitude modulated
frequency bands. However, the modulation index does not make clear the phase
–amplitude coupling relationship between the phase modulating and the amplit-
itude modulated frequency bands (i.e., whether the amplitude-modulated band
resides on the peak or trough of the phase-modulated band cannot be described
from the modulation index). The phaseamogram is used to reveal the phase
–amplitude relations between the phase-modulating and amplitude-modulated
frequency bands. The comodulogram, an extension of the modulation index, is
a principled approach to choosing the phase-modulating frequency band and
amplitude-modulated frequency bands of interest. This is because the comodul-
ogram depicts modulation indices across a range of phase-modulating and
amplitude-modulated frequency bands. We computed the comodulogram
on 60-s long nonoverlapping data segments. These data were averaged to
obtain a within-subject comodulogram. We next computed the median comodu-
logram across subjects. We calculated the modulation index between lower
(0.1–0.8 Hz, 0.32 Hz steps, 1 Hz bandwidth) and higher frequencies (1–30 Hz, 1 Hz bandwidth) as previously described41.

Phase–amplitude coupling and amplitude vector distribution. Phase–amplitu-
de coupling and amplitude vector distribution were computed on 60-s long,
nonoverlapping data segments. These data were averaged to obtain within-subject estimates.
We next computed the median across subjects.

To compute phase–amplitude coupling dynamics associated with low and high
frequencies of interest, we constructed phaseamograms by adapting a previously
described method to our dataset51. We calculated the phaseamograms between
delta frequencies (2–4 Hz) and higher frequencies (4.1–30 Hz, 3 Hz steps). We computed the median phaseamogram across all subjects for group-level
visualization.

We computed the normalized mean amplitude vector corresponding to the
higher frequencies of interest and the circular mean of the mean amplitude vectors.
This resulted in a vector whose length represents the mean amplitude of higher
frequency activity and phase represents the mean angular location of higher
frequency on the phase of the low oscillation driver frequency. We computed the mean amplitude vector for group-level visualization.

Statistics and reproducibility. We used an empirical bootstrap approach to enable
statistical inferences52. First, we bootstrapped the estimates of each non-
overlapping window. Next, we computed the median of the bootstrapped estimates
at the subject level and computed the group median of this estimate. We computed the median difference between groups and then iterated the above procedure 5000
times to obtain a distribution of the median difference between groups.
We computed the 99% confidence interval of this distribution. We used the omnibus circular test statistic to test phase uniformity53. We performed a linear mixed-
effects model with Tukey's HSD for post-hoc comparisons to test for differences
between the mean amplitude vectors JMP®, Pro 14 (SAS Institute Inc., Cary, NC,
1989–2007).

Reporting summary. Further information on research design is available in the
Nature Research Reporting Summary linked to this article.

Data availability

The data supporting the findings of this study will be made available from the
Corresponding authors upon reasonable request.

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