Neurophysiological Correlates of Sevoflurane-induced Unconsciousness

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ABSTRACT

Background: Recent studies of anesthetic-induced unconsciousness in humans have focused predominantly on the intravenous drug propofol and have identified anterior dominance of alpha rhythms and frontal phase–amplitude coupling patterns as neurophysiological markers. However, it is unclear whether the correlates of propofol-induced unconsciousness are generalizable to inhaled anesthetics, which have distinct molecular targets and which are used more commonly in clinical practice.

Methods: The authors recorded 64-channel electroencephalograms in healthy human participants during consciousness, sevoflurane-induced unconsciousness, and recovery (n = 10; n = 7 suitable for analysis). Spectrograms and scalp distributions of low-frequency (1 Hz) and alpha (10 Hz) power were analyzed, and phase–amplitude modulation between these two frequencies was calculated in frontal and parietal regions. Phase lag index was used to assess phase relationships across the cortex.

Results: At concentrations sufficient for unconsciousness, sevoflurane did not result in a consistent anteriorization of alpha power; the relationship between low-frequency phase and alpha amplitude in the frontal cortex did not undergo characteristic transitions. By contrast, there was significant cross-frequency coupling in the parietal region during consciousness that was not observed after loss of consciousness. Furthermore, a reversible disruption of anterior–posterior phase relationships in the alpha bandwidth was identified as a correlate of sevoflurane-induced unconsciousness.

Conclusion: In humans, sevoflurane-induced unconsciousness is not correlated with anteriorization of alpha and related cross-frequency patterns, but rather by a disruption of phase–amplitude coupling in the parietal region and phase–phase relationships across the cortex. (Anesthesiology 2014; XXX:00-00)
Materials and Methods

This study was conducted at the University of Michigan Medical School and was approved by the Institutional Board Review (HUM00061087); written consent was obtained from all participants after a careful discussion of risks and benefits.

Study Population

We recruited 10 healthy volunteers for the study based on similar studies of propofol-induced unconsciousness; two volunteers withdrew during administration of sevoflurane due to discomfort and one subject's electroencephalographic data were suboptimal due to excessive motion. Thus, data from seven healthy volunteers (four male, 20 to 23 yr of age) were included in the analysis. Participants were American Society of Anesthesiologists class I physical status, body mass index less than 30, with Mallampati 1 or 2 airway classifications, and no other factors predictive of difficult airway. We excluded subjects who were pregnant, or with a history of obstructive sleep apnea, gastroesophageal reflux, cardiac conduction abnormalities, asthma, epilepsy, history of problems with anesthesia, family history of problems with anesthesia, history of drug use, and any neurologic or psychiatric history. Pregnancy and illicit drug use were ruled out through urine and blood analyses.

Anesthetic Protocol

Standard intraoperative monitors (noninvasive blood pressure measurement, electrocardiogram, pulse oximetry, and capnography) and temperature probes were employed; at least two fully trained anesthesiologists conducted all studies. Participants were instructed to keep their eyes closed during waking states throughout the experiment and received sevoflurane anesthesia by a secured face mask with an initial concentration of 0.4% in high-flow oxygen (8 l/min). Before anesthetic delivery, adequate face-mask seal was confirmed by analyzing inspired and expired concentrations of oxygen while delivering an FiO₂ of 1. Sevoflurane was administered at a given level for a 15-min equilibration period to achieve steady state, followed by 10 min at the target concentration. Sevoflurane concentration was increased by increments of 0.2% until loss of consciousness (LOC) was achieved. This protocol is based on that of Suzuki et al., who found that 0.7% was the approximate concentration of sevoflurane at which consciousness was lost. After at least 10 min of unconsciousness, the reverse protocol (equilibration followed by steady-state period) was employed until participants regained consciousness. After recovery of consciousness (ROC), anesthetic concentration was still titrated downward until end-tidal values were 0% (fig. 1A). Before sevoflurane exposure, participants had objects placed in each hand that would emit a sound when squeezed. At the beginning of sevoflurane administration, participants were instructed to squeeze the object in either their left hand or right hand; instructions were delivered every 30 s through an audio loop and left/right commands were randomized. Positive or negative responses were observed and recorded in the data collection software by two investigators throughout the experimental period.

Behavioral State Analysis

Each participant’s experimental session was divided into nonoverlapping 5-min epochs. Within each epoch, the time-series of positive and negative responses to auditory commands (every 30 s) was used to calculate the percentage of time the participant was responsive. Changes in responsiveness were assumed to reflect changes in levels of consciousness over the course of the experiment. To compare and pool data across participants, we defined two behavioral time markers for each subject: (1) LOC—the first negative response to an auditory command during induction that was followed by at least 5 min of 0% responsiveness; and (2) ROC—the first positive response to an auditory command after LOC. We further defined

Electroencephalography Data Acquisition and Preprocessing

The electroencephalogram was acquired using a 64-channel sensor net from Electrical Geodesics, Inc. (Eugene, OR); all channels were referenced to the vertex. Electrode impedance was reduced to below 50 kΩ before data collection (as per manufacturer recommendations) and electroencephalographic signals were collected using a 64-channel biosignal amplifier at a rate of 500 Hz. The electroencephalogram was visually monitored in real-time to ensure continued signal integrity throughout the data collection period. After the experiment, electroencephalographic signals were high-pass filtered at 0.1 Hz, and re-referenced to an average reference. An investigator (S.B.M.) experienced in reading electroencephalograms visually inspected the data to reject epochs and channels with noise or nonphysiological artifacts.
six 5-min “alignment epochs” based on experimental conditions and individual responsiveness profiles: (1) Baseline—before sevoflurane exposure, participants at rest with their eyes closed (100% responsiveness); (2) Effect onset of sevoflurane (EOn)—during induction, immediately after the first negative response to an auditory command (≤90% responsiveness); (3) Transition to unconsciousness (TrUN)—during induction, immediately before LOC (>0% responsiveness); (4) Unconscious—between LOC and ROC (0% responsiveness); (5) Transition to recovery (TrCON)—during emergence, immediately after ROC (>0% responsiveness); and (6) Recovery—after emergence, participant at rest with their eyes closed (100% responsiveness). These defined epochs are illustrated in figure 1B.

Electroencephalographic Analysis

Spectral Analysis. Spectrograms were computed in Chronux,* using the multitaper method, with window lengths of $T = 2$ s, step size of 0.1 s, time-bandwidth product $NW = 2$, number of tapers $K = 3$. These parameters were chosen to replicate the spectral analysis in a recent study of propofol-induced unconsciousness.2 Electroencephalographic data from all 64 channels were used in the spectral analysis for each participant. To compute group-level spectrograms, we aggregated all electroencephalogram channels from each participant across the six alignment epochs described in the Behavioral State Analysis section.

Topographic Analysis. Scalp power distributions were calculated using the topoplot function in EEGLab.12

Phase–Amplitude Coupling Analysis. Phase–amplitude coupling was conducted using the Phase–Amplitude Coupling Toolbox13 in EEGLab. We used band-pass filters to extract low-frequency (0.1 to 1 Hz) and alpha (8 to 14 Hz) oscillations from the electroencephalogram for each participant. Next, we used a Hilbert transform to extract the instantaneous amplitude and phase from each bandwidth. We computed a phase–amplitude modulogram6 by assigning each temporal sample to one of $n = 18$ equally spaced phase bins based on the instantaneous value of the low-frequency phase, and then averaging the corresponding instantaneous amplitude of alpha within a 1-min epoch. The modulogram was calculated for five frontal electrodes near Fz and four parietal electrodes near Pz, and the results were averaged across all participants.

PLI Analysis. A surrogate of functional connectivity across frontal–parietal, frontal–temporal, and frontal–occipital brain regions was calculated using PLI, which is sensitive only to nonzero phase lead/lag relationships to avoid the problem of volume conduction.14 The instantaneous phase of each channel of the electroencephalogram was extracted

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using a Hilbert transform. Phase difference $\Delta \phi_i$ was calculated between channels, where $\Delta \phi_i = \phi_{i,t} - \phi_{j,t}$, $t = 1, 2, \ldots$, $n$, $n$ is the number of samples within one epoch, and $i$ and $j$ were set to include all channels in frontal, parietal, temporal, and occipital brain regions. We then calculated PLI as follows:

$$\text{PLI}_{ij} = \left[\text{sign}(\Delta \phi_i)\right]$$

In this equation, the sign() function results in 1 if $\Delta \phi_i > 0$, 0 if $\Delta \phi_i = 0$, and 1 if $\Delta \phi_i < 0$. A PLI value close to 1 indicates that the instantaneous phase of one signal is consistently ahead of another signal and the phases are locked, whereas a PLI value close to 0 indicates that there is no consistent phase lead or lag relationship between signals. We controlled for spurious PLI values by performing a surrogate data analysis as follows. Surrogate data sets were generated for each combination of channel pairs $i$ and $j$ by maintaining the phase time series (calculated using a Hilbert transform) of channel $i$, and interchanging the phase time series from 0 to $n/2$ with the phase time series from $n/2$ to $n$ of channel $j$, where $n$ is the number of samples within one epoch. This method ensured that the true phase relationships between channels were eliminated although maintaining the spectral properties of each epoch. As with the empirical datasets, the PLI was calculated for surrogate datasets derived from frontal–parietal, frontal–temporal, and frontal–occipital combinations.

**Methodological Control**

To establish a methodological control condition that validated the algorithms employed in this study, we tested our methods on two sets of electroencephalographic data collected from participants undergoing propofol-induced unconsciousness. Dataset $P_1$ was obtained using the same 64-channel electroencephalograph and was collected over 15 min from a single human volunteer. Propofol infusion began at 100 $\mu$g kg$^{-1}$ min$^{-1}$ and increased by this amount every 5 min until LOC, defined by the same criteria as used in sevoflurane experiments. Dataset $P_2$ consisted of 8-channel electroencephalographic data previously collected from nine surgical patients.$^{15}$ Propofol was administered at an initial target controlled infusion of 2.0 $\mu$g/ml and increased at a rate of 1.0 $\mu$g/ml every 20 s until LOC. Five minutes of electroencephalographic activity were analyzed from five periods: (1) baseline; (2) propofol induction, ending in LOC; (3) unconsciousness; (4) recovery, beginning with cessation of propofol delivery and ending with return of consciousness; and (5) postrecovery. Note that periods four and five reflect the effects of surgery as well as drugs other than propofol. The identification of increased alpha power, alpha anteriorization, and expected cross-frequency coupling changes confirmed the ability of our algorithms to detect these patterns when present (Supplemental Digital Content 1, fig. 1, http://links.lww.com/ALN/B104).

**Statistical Analysis**

**Spectrogram.** We determined whether or not there was a significant increase from baseline in low-frequency and alpha power in the group-level spectrogram by performing a two-tailed sign test at each time-frequency location in these bandwidths, treating each participant as an independent sample. Values were considered significant at $\alpha$ less than 0.05.

**Phase Amplitude Coupling.** Canolty’s phase–amplitude coupling Modulation Index was calculated for each minute of the phase–amplitude modulogram.$^{16,17}$ To test the null hypothesis that the phase and amplitude are decoupled, we used the surrogate data method proposed in Canolty et al.$^{16}$ For each minute of data, we generated a random time shifts $\Delta t$ that ranged between 0 and 60 s. We then shifted the amplitude series by $\Delta t$ although keeping the phase series fixed, and calculated the Modulation Index for the resulting surrogate time series. We repeated this procedure 2,000 times to produce distributions of values of the Modulation Index for datasets in which the null hypothesis should hold. The Modulation Index of the nonshifted time series was then compared to this distribution and was considered significant if it exceeded 95% of the surrogate values ($P < 0.05$).

**Phase Lag Index.** phase lag index values were compared among baseline, unconscious, and recovery epochs using a one-way ANOVA with a post hoc Tukey correction for each of the following channel combinations: frontal–parietal, frontal–temporal, and frontal–occipital. For each of these combinations, a Student $t$ test (two-tailed) was used to compare PLI values from baseline, unconscious, and recovery epochs with the PLI from the randomized counterpart of surrogate datasets. All statistical analysis were conducted in MATLAB (Natick, MA) and differences were considered significant at $\alpha$ less than 0.05.

**Results**

**Pharmacological and Behavioral Characteristics of Sevoflurane-induced Unconsciousness**

Six of the seven participants experienced LOC that lasted for at least 10 min as sevoflurane levels were being increased from 0.6 to 0.8%, which is consistent with the protocol described by Suzuki et al.$^{11}$ The sevoflurane concentration profile over the experimental session for these six participants is illustrated in figure 1A. One participant lost consciousness at a steady-state concentration of 0.4% sevoflurane; for this participant, unconsciousness was maintained for 10 min, and sevoflurane was subsequently reduced to 0%. Each participant demonstrated a unique behavioral responsiveness profile to the auditory commands as sevoflurane levels changed, with some losing and recovering consciousness once, and others many times throughout the experimental session. One such responsiveness profile is presented in figure 1B, with an
consciousness or unconsciousness (fig. 4A). In parietal channels, significant coupling was observed during any period of correlation phase and alpha amplitude. In frontal channels, no significant coupling was observed in 60% of the conscious epochs but in none of the unconscious epochs (fig. 4B). The average modulation index over time for all frontal and parietal channels is presented in figure 4C.

**Sevoflurane-induced Unconsciousness Is Correlated with Decreased Phase Lag Index in the Alpha Bandwidth across Frontal–Parietal, Frontal–Temporal, and Frontal–Occipital Regions**

We calculated PLI between frontal–parietal, frontal–temporal, and frontal–occipital channel combinations in the alpha bandwidth during baseline, unconscious and recovery alignment epochs. In each comparison, a one-way ANOVA yielded significant differences between epochs (frontal–parietal: $P = 0.013$; frontal–temporal: $P = 0.0043$; and frontal–occipital: $P = 0.0002$; fig. 5A). A post hoc Tukey test showed that the unconscious epoch had significantly lower PLI values than baseline or recovery epochs (frontal–parietal: $P = 0.025$; frontal–temporal: $P = 0.01$; and frontal–occipital: $P = 0.0006$); PLI values in baseline and recovery epochs were not significantly different. Decreased PLI during sevoflurane-induced unconsciousness was observed for each individual participant between frontal and occipital brain regions in the alpha bandwidth, despite the fact that dominance of alpha power did not shift from occipital to frontal regions in the majority of subjects (fig. 5B). Surrogate PLI analysis did not, as expected, result in nonzero values; empirical PLI values were significantly higher than all surrogate PLI counterpart values across baseline (frontal–parietal: $P = 0.003$; frontal–temporal: $P = 0.001$; and frontal–occipital: $P = 0.0004$), unconsciousness (frontal–parietal: $P = 0.0005$; frontal–temporal: $P = 0.0003$; and frontal–occipital: $P = 0.0004$), and recovery (frontal–parietal: $P = 0.003$; frontal–temporal: $P = 0.001$; and frontal–occipital: $P = 0.0006$).

**Discussion**

The current study demonstrates that the stepwise, steady-state induction of unconsciousness with sevoflurane is not characterized by high-power slow oscillations, anterior dominance of alpha rhythms, or a specific phase–amplitude relationship between these frequencies in the frontal cortex. Instead, sevoflurane-induced unconsciousness was correlated with disrupted phase–amplitude relationships in parietal cortex and disrupted phase–phase relationships across frontal–parietal, frontal–temporal, and frontal–occipital regions of the brain.

**Past Studies of Anteriorized Power and Cross-frequency Coupling**

In the 1970s, Tinker et al.\(^{18}\) were the first to test the hypothesis (in Java monkeys) that anteriorization or “frontal dominance” of electroencephalographic power was correlated with the loss of ability to respond to a stimulus, as induced by various anesthetic drugs. In the mid-1990s, the reduction of the expected posterior alpha in the awake, eyes-closed state, and the shift to more anterior structures was identified during...
propofol sedation and during unconsciousness induced by a combination of isoflurane and nitrous oxide. In 2001, John et al. suggested that electroencephalographic anteriorization of alpha and slow-wave activity was an agent-independent feature of general anesthesia, based on the study of 176 surgical patients undergoing general anesthesia with a wide variety of intravenous and inhaled anesthetics. In 2004, Feschenko et al. demonstrated the disappearance of occipital alpha oscillations and the appearance of high-power frontal alpha oscillations during propofol-induced unconsciousness in 52 healthy human participants.

More recently, Purdon et al. used high-density electroencephalography to demonstrate that anteriorized alpha rhythms are associated with propofol-induced unconsciousness. During propofol-induced unconsciousness, slow oscillations have also been associated with anterior structures; peak slow-wave activity correlates with gray matter volume of the frontal cortex. Cross-frequency coupling analysis of alpha rhythms and slow oscillations revealed characteristic phase–amplitude relationships across states. The “trough-max” pattern is associated with transitions around the loss or recovery of consciousness and is defined by the maximal alpha amplitude occurring at the trough of the slow oscillation. Conversely, the “peak-max” pattern is associated with deeper levels of propofol-induced unconsciousness and is defined by the maximal alpha amplitude occurring at the peak of the slow oscillation. This trough-max and peak-max relationship had been previously demonstrated in a study of desflurane, but at higher relative doses than those used in the current study.

**Alpha Rhythms and Sevoflurane-induced Unconsciousness**

Using methodology that is comparable to recent studies, we were not able to demonstrate a consistent association...
of unconsciousness induced by sevoflurane with the anterior dominance of alpha oscillations. Gugino et al. compared the electroencephalographic topography of propofol and sevoflurane, finding an increased tendency to anteriorization of alpha with propofol at induction and before recovery of consciousness. Findings related to sevoflurane in this study were possibly confounded by polypharmacy, since volunteers received fentanyl premedication and remifentanil during administration of propofol and sevoflurane. The addition of remifentanil to an anesthetic can potentially alter level of sedation as well as the characteristics of frontal electroencephalography. By contrast, the current investigation was conducted using sevoflurane exclusively.

Fig. 4. Cross-frequency coupling between low-frequency (0.1 to 1 Hz) phase and alpha (8 to 14 Hz) amplitude. (A) Frontal channels. No significant coupling is observed during conscious epochs or sevoflurane-induced unconsciousness. (B) Modulation index for frontal (purple) and parietal (green) channels. Values above the dotted line are significant at P value less than 0.05. (C) Parietal channels. Significant coupling is observed during conscious epochs but not during sevoflurane-induced unconsciousness. LOC = loss of consciousness; ROC = return of consciousness; Trans<sub>CON</sub> = transition to consciousness; Trans<sub>UN</sub> = transition to unconsciousness.

Fig. 5. Phase lag index (PLI) analysis. A PLI value of 1 reflects perfectly consistent phase lead/lag relationship and a value of 0 reflects no consistent phase relationship. (A) Average PLI in the alpha bandwidth across aligned baseline, unconscious and recovery epochs. PLI values significantly decrease during sevoflurane-induced unconsciousness across three brain region combinations. (B) Individual PLI values across baseline, unconsciousness, and recovery alignment epochs between frontal and occipital channels. Each individual participant demonstrates a reversible decrease in PLI upon sevoflurane-induced unconsciousness, despite the lack of a consistent shift of alpha power from occipital to frontal cortices. Note that for all empirical data analyses shown in this figure, counterpart surrogate datasets suggested no spurious phase–phase relationships.
There are several reasons why our findings could be discrepant with past studies of propofol and other anesthetics. First, it is possible that anteriorization occurs at higher concentrations of sevoflurane than those tested in the current study. Indeed, Gugino et al.\textsuperscript{24} reported a dose-dependent effect on anteriorization during administration of sevoflurane but not propofol (which had already reached maximal levels at LOC). However, we were specifically testing the hypothesis that anterior dominance was a marker of sevoflurane-induced unconsciousness, rather than the hypothesis that sevoflurane can—at some dose—cause anteriorized electroencephalographic activity in the alpha bandwidth.

Second, the data presented in the current study were acquired during anesthetic-induced unconsciousness in the absence of a noxious stimulus. By contrast, the protocol of Gugino et al.\textsuperscript{24} included graded attempts to arouse the participants. As such, higher concentrations of anesthetic were required in order to suppress a response to command compared to the simple auditory stimulus that we employed. As noted, opioids were also included in the Gugino et al. study, which can facilitate sedation and alter electroencephalographic patterns in the frontal cortex.

Third, it is possible that distinct underlying molecular mechanisms account for the distinct neurophysiological profile. Ching et al.\textsuperscript{4,26} and Vijayan et al.\textsuperscript{5} have developed models for the anteriorization of alpha rhythms induced by propofol—and potentially other anesthetics—based on agonism of the GABA\textsubscript{A} receptor in the thalamus. Accordingly, the phenomenon of anteriorization in relationship to unconsciousness should be determined by an anesthetic’s (1) selectivity for the GABA\textsubscript{A} receptor, and (2) dependence on modulating GABA transmission to achieve the behavioral endpoint of unconsciousness. Sevoflurane is a halogenated ether that is less selective for the GABA\textsubscript{A} receptor and less dependent than propofol on GABA\textsubscript{A} agonism to achieve anesthetic endpoints.\textsuperscript{27,28} For example, in a rat hippocampal slice model, propofol primarily enhanced inhibitory pathways (mediated by GABA\textsubscript{A} receptors), whereas sevoflurane primarily depressed excitatory pathways (mediated by glutamate receptors).\textsuperscript{24} Furthermore, changes in ambient GABA concentrations result in altered cellular and behavioral responses to propofol but only minimally affect responses to sevoflurane.\textsuperscript{25} Based on these considerations, we suggest that, as would be predicted by recent modeling studies, the electroencephalographic anteriorization associated with propofol reflects a certain threshold of GABA\textsubscript{A} agonism, which is a critical molecular mechanism of propofol-induced unconsciousness. By contrast, sevoflurane is less selective than propofol for GABA\textsubscript{A} receptors and less dependent on GABA\textsubscript{A} agonism for the induction of unconsciousness. Therefore, anteriorization does not appear consistently at the state transition of sevoflurane-induced unconsciousness but rather at higher doses of sevoflurane\textsuperscript{24} when the reduced selectivity for the GABA\textsubscript{A} receptor is overcome by increased concentrations of the drug. This hypothesis is further supported by comparing our results to a past study of unconsciousness induced by ketamine, for which the GABA\textsubscript{A} receptor is not a major target.\textsuperscript{30} Lee et al.\textsuperscript{31} demonstrated a significant decrease in relative alpha power during ketamine-induced unconsciousness. These parallel findings reinforce the hypothesis that the electroencephalographic characteristics associated with propofol-induced unconsciousness (e.g., anterior dominance of alpha, increased alpha power) are mediated by GABA\textsubscript{A} agonism.

**Phase–Amplitude Relationships and Sevoflurane-induced Unconsciousness**

We did not observe significant phase–amplitude coupling in the frontal electroencephalogram during any of the alignment epochs. Our results are distinct from phase–amplitude relationships observed during propofol-induced unconscious, where the loss and recovery of consciousness have been characterized by a trough–max coupling pattern.\textsuperscript{2} Mukamel et al.\textsuperscript{6} associated this trough–max pattern with anterior structures such as the anterior cingulate and the frontal cortices; this coupling occurs at approximately the same time as an increase in alpha power in the frontal electrodes. We did not observe consistent alpha anteriorization or the emergence of a trough–max pattern, which suggests that the two phenomena are coupled. We also did not observe the emergence of a peak–max phase amplitude coupling pattern. This result is to be expected, as peak–max coupling has been associated with profound unconsciousness and we did not increase sevoflurane concentrations beyond what was necessary to achieve LOC. Again, it is possible that the characteristic changes of cross-frequency coupling in frontal channels are mediated specifically by changes in GABA\textsubscript{A} receptor function. Blain-Moraes et al.\textsuperscript{32} recently demonstrated that the pattern of phase–amplitude modulation observed during propofol-induced unconsciousness does not occur during ketamine-induced unconsciousness.

An important positive finding of our cross-frequency coupling analysis was the significant trough–max coupling observed in parietal channels during 60% of epochs before LOC and after recovery of consciousness. By contrast, there was no significant coupling observed during the unconscious epoch. This observed coupling disruption in parietal but not frontal regions is consistent with studies that have demonstrated that parietal networks are more affected by anesthetics than frontal networks.\textsuperscript{33,34} The fact that significant coupling only occurred 60% of the time outside of the unconscious epoch may potentially be explained by the fact that participants were only conscious for a fraction of the time during transitions into and out of unconsciousness.

**Sevoflurane-induced Unconsciousness Is Correlated with Disrupted Phase–Phase Relationships across the Brain**

LOC during exposure to sevoflurane was characterized by a significant and reversible decrease of PLI, a surrogate functional connectivity measure based on phase lead/lag relationships observed during propofol-induced unconsciousness. Indeed, Gugino et al.\textsuperscript{31} demonstrated a significant decrease in relative alpha power during ketamine-induced unconsciousness.
relationships that preferentially reflects longer-range phase relationships. Importantly, the frontal–occipital PLI in the alpha bandwidth was reduced in each volunteer during the period of unconsciousness. The fact that this anterior–posterior functional disconnection was observed consistently despite the absence of uniform alpha anteriorization suggests that disrupted phase–phase relationships may be mechanistically independent and a more sensitive indicator of sevoflurane-induced unconsciousness. Furthermore, this phase–phase disruption across the cortex may be independent of GABA_A modulation. In support of this, we have recently demonstrated that ketamine-induced unconsciousness is also associated with a decrease in PLI. Thus, unlike anesthesia, anesthetic actions at distinct molecular targets may nonetheless result in a functional disconnection across the cortex, suggesting a state-specific versus a drug-specific phenomenon. Functional disconnections between the frontal cortex and other brain regions may represent a “top–down” mechanism for anesthetic-induced unconsciousness, which is supported by the fact that molecularly diverse anesthetics can disrupt long-range coupling in a cortical slice model that does not include a thalamus. However, it is also possible that anesthetic actions in higher-order nuclei of the thalamus could destabilize corticocortical or corticothalamic-cortical interactions. In either case, the results of this and past investigations in humans suggest that multiple classes of anesthetics can disrupt phase relationships across the cortex resulting in a fragmentation of networks that might represent the common mediator and proximate cause of anesthetic-induced unconsciousness.

Limitations
This study has numerous limitations. First, this was a small study of young healthy volunteers and, despite the significant differences identified in parietal phase–amplitude relationships and cortical phase–phase relationships, we cannot extrapolate our data to a broad population of surgical patients of varying ages and comorbidities. Second, we did not study doses of sevoflurane beyond those required to achieve a consistently unconscious state. Third, we cannot extend our findings to other volatile anesthetics since we did not study all halogenated ethers or haloalkanes in current use only, 6 months from the cover date of the issue. A

Conclusions
Unconsciousness during administration of sevoflurane is not characterized by the anterior dominance of alpha rhythms and slow oscillations, or specific phase–amplitude relationships between these frequencies in the frontal cortex. Instead, reversible disruptions of parietal phase–amplitude coupling and cortical phase–phase relationships in the alpha bandwidth are correlated with sevoflurane-induced unconsciousness.

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Competing Interests
The authors declare no competing interests.

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