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Rapid fragmentation of neuronal networks at the onset of propofol-induced unconsciousness

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The neurophysiological mechanisms by which anesthetic drugs cause loss of consciousness are poorly understood. Anesthetic actions at the molecular, cellular, and systems levels have been studied in detail at steady states of deep general anesthesia. However, little is known about how anesthetics alter neural activity during the transition into unconsciousness. We recorded simultaneous multiscale neural activity from human cortex, including ensembles of single neurons, local field potentials, and intracranial electrocorticograms, during induction of general anesthesia. We analyzed local and global neuronal network changes that occurred simultaneously with loss of consciousness. We show that propofol-induced unconsciousness occurs within seconds of the abrupt onset of a slow (<1 Hz) oscillation in the local field potential. This oscillation marks a state in which cortical neurons maintain local patterns of network activity, but this activity is fragmented across both time and space. Local (<4 mm) neuronal populations maintain the millisecond-scale connectivity patterns observed in the awake state, and spike rates fluctuate and can reach baseline levels. However, neuronal spiking occurs only within a limited slow oscillation-phase window and is silent otherwise, fragmenting the time course of neural activity. Unexpectedly, we found that these slow oscillations occur asynchronously across cortex, disrupting functional connectivity between cortical areas. We conclude that the onset of slow oscillations is a neural correlate of propofol-induced loss of consciousness, marking a shift to cortical dynamics in which local neuronal networks remain intact but become functionally isolated in time and space.

General anesthesia is a drug-induced reversible coma commonly initiated by administering a large dose of a fast-acting drug to induce unconsciousness within seconds (1). This state can be maintained as long as needed to execute surgical and many nonsurgical procedures. One of the most widely used anesthetics is propofol, an i.v. drug that enhances GABAergic inhibitory input to neurons (2–4), with effects in cortex, thalamus, brainstem, and spinal cord (5–7). Despite the understanding of propofol’s molecular actions, it is not clear how these effects at molecular targets affect single neurons and larger-scale neural circuits to produce unconsciousness.

The effects on macroscopic dynamics are noticeable in the EEG, which contains several stereotyped patterns during maintenance of propofol general anesthesia. These patterns include increased delta (0.5–4 Hz) power (8, 9); increased gamma (25–40 Hz) power (9); an alpha (~10 Hz) rhythm (10–12) that is coherent across frontal cortex; and burst suppression, an alternation between bursts of high-voltage activity and periods of flat EEG lasting for several seconds (13, 14). In addition, slow oscillations (<1 Hz) have been well characterized in deeply anesthetized animals and are associated with an alternation of the neuronal membrane potential between UP (depolarized) and DOWN (hyperpolarized) states (8, 15).

Although these patterns are observed consistently, it is unclear how they are functionally related to unconsciousness under general anesthesia. Most studies have focused on a deep steady state of general anesthesia and have not used a systematic behavioral measure to track the transition into unconsciousness. This steady-state approach cannot distinguish between patterns that are characteristic of a deeply anesthetized brain and those that arise at the onset of unconsciousness. Unconsciousness can occur in tens of seconds (4), but many neurophysiological features continue to fluctuate for minutes after induction and are highly variable between different levels of general anesthesia (1, 16). Therefore, identifying the specific dynamics associated with loss of consciousness (LOC) requires an examination of the transition into unconsciousness, linking neurophysiology with behavioral measures.

In addition, the dynamic interactions between cortical areas that underlie these EEG oscillations are not well understood, because few studies have simultaneously recorded ensembles of single neurons and oscillatory dynamics from sites distributed across the brain. Consequently, how propofol acts on neural circuits to produce unconsciousness remains unclear. A leading hypothesis suggests that anesthetics disrupt cortical integration (17, 18). Identifying the mechanism by which this disruption might occur requires a better understanding of how the spatial and temporal organization of neural dynamics evolves during induction of unconsciousness.

To address this question, we investigated both neuronal and circuit-level dynamics in the human brain during induction of unconsciousness with propofol. We obtained simultaneous recordings of single units, local field potentials (LFPs), and intracranial electrocorticograms (ECoG) over up to 8 cm of cortex, enabling us to examine neural dynamics at multiple spatial scales with milli-
second-scale temporal resolution. We used a behavioral task to identify within seconds the time at which patients became unresponsive to auditory stimuli, which we defined as LOC.

Our results reveal a set of neurophysiological features that accompany LOC that, together with previously reported effects (8, 9, 15), enable a multiscale account of this profound shift in brain state. We find that LOC is marked by the abrupt onset of slow oscillations (0.1–1 Hz) in the LFP. Power in the slow oscillation band rises sharply at LOC and maintains this increase throughout the post-LOC period. Neuronal spiking becomes coupled to the local slow oscillation within seconds of LOC: Spiking occurs only in short intervals of activity that are interspersed with suppression lasting hundreds of milliseconds, periodically interrupting information processing. These periods in which activity may occur are not simultaneous across the brain, implying that information transfer between distant (>2 cm) cortical networks is impaired. Cortical networks therefore are fragmented both temporally and spatially, disrupting both local and long-range communication. However, small-scale (<4 mm) functional connectivity measures remain similar to the conscious state, and neuronal spike rates can recover to baseline levels after LOC despite continued unresponsiveness. This result demonstrates that short periods of normal spike dynamics still can occur during unconsciousness. We conclude that the slow oscillation is a fundamental component of propofol-induced unconsciousness, marking a functional isolation of cortical regions while significant connectivity is preserved within local networks.

Results

We recorded single units (n = 198), LFPs, and ECoG in three patients undergoing intracranial monitoring for surgical treatment of epilepsy. Single units and LFPs were recorded from a 96-channel microelectrode array (19) implanted in temporal cortex for research purposes. We recorded throughout induction of general anesthesia by bolus administration of propofol before planned neurosurgery to remove the electrodes. Patients performed an auditory task requiring a button press in response to stimuli. All patients completely ceased responding to the task within 40 s of propofol administration and remained unresponsive for the remainder of the recording period, lasting 5–10 min after LOC. LOC was defined as the onset of the period of unresponsiveness to auditory stimuli. To acknowledge the fact that LOC could have occurred at any point between the last response and the failure to make the next response, LOC was defined as the interval beginning 1 s before the first missed stimulus up until the second missed stimulus (5 s total). We then compared spectra across all ECoG channels in the pre- and post-LOC periods. In agreement with previous scalp EEG studies of healthy subjects (9), we found that average spectra in the post-LOC period differed significantly from those in the pre-LOC period: Slow (0.1–1 Hz) and gamma (25–40 Hz) power increased in the unconscious state (Fig. S1). These results suggested that propofol acted as expected in these patients and did not reveal any gross disruption of GABA networks.

Spike Rates Are Highly Variable After LOC. To determine the relationship between changes in spike rate and LOC, we first examined the overall spike rate in a local network of cortical neurons. Consistent with propofol’s enhancement of GABAergic signaling, widespread suppression of spiking was observed after LOC. In each patient, the spike rate across the population of units decreased significantly 0–30 s after LOC (Fig. 1 and Table S1; the range of values is the range across patients). Mean spike rates across all units reached a minimum 35–85 s after LOC, having decreased 81–92% from the baseline awake state. However, spike rates subsequently recovered over several minutes (Fig. 1B). At 4 min after LOC, the rate across the entire population of units varied widely, ranging from 33% of baseline in patient A to 117% of baseline in patient B. At this 4-min post-LOC period, individual units also displayed a wide range of spike rates, with some as high as or higher than baseline; only 35.2% of units still had spike rates significantly below baseline, 55.1% of units were not significantly different, and 9.7% of units had significantly increased spike rates. We conclude that propofol rapidly causes a nearly complete but transient suppression of cortical spiking, and after several minutes many individual neurons recover to baseline spike rates. The fluctuation in spike rates across time, which could have come about from changing propofol blood levels, demonstrates that brain state is dynamic after LOC. However, subjects remained unconscious throughout this period despite widely varying spike rates, suggesting that unconsciousness is not strictly associated with gross changes in spike rate.

Spiking Activity Is Organized into Periods of Activity and Quiescence After LOC. Given that mean spike rates did not exhibit a fixed relationship with state of consciousness, we examined whether unconsciousness was associated instead with a change in the temporal structure of spiking. We observed that spiking activity across the population of units occurred in short periods of activity that were interrupted by periods of silence. To estimate conservatively the amount of time with no spike activity, we binned spikes from all units into 400-ms bins. We found that 63% of bins contained no spikes, significantly more than simulated neurons with a constant rate (33%, P < 0.001 for each patient, Pearson’s χ² test). Therefore we concluded that cortical
networks can be highly active during unconsciousness, but this activity is concentrated in short periods that are followed by profound suppression.

**Unconscious State Is Marked by a Rapid Increase and Stable Maintenance of Power in the Slow Oscillation Band.** The slow oscillation is known to modulate neuronal spiking (8, 15), and therefore we examined the time course of its onset relative to LOC. Before LOC, power in the slow oscillation band (0.1–1 Hz) was stable (SD <7% in each patient before LOC). At LOC, power in the slow oscillation band increased abruptly by 35–70% (Fig. 2), and this power increase occurred within one 5-s window of LOC in all patients (Table S1). The slow oscillation power then persisted at this high level for the remainder of the recording, with 99.0% of the post-LOC time bins having higher slow oscillation power than occurred in any time bin during baseline (Fig. 2A). We therefore concluded that power in the slow oscillation band is modulated simultaneously with LOC and is preserved thereafter despite large fluctuations in spike rate.

We next examined other frequency bands to investigate whether the power change at LOC was specific to the slow oscillation band or whether other frequency bands showed a similar relationship. Although power in the >10 Hz range increased slowly after LOC, theta (3–8 Hz) power showed the opposite trend, decreasing 20–30% after LOC (Fig. 2B and Fig. S1). In addition, power in all these bands continued to undergo modulations for several minutes rather than maintaining a consistent change after LOC (Fig. S1), perhaps as the result of differences in propofol dosage during the maintenance phase. The stable increase in power at LOC therefore was specific to the slow oscillation band. These results demonstrated that both spike rates and many oscillatory features (gamma, alpha, theta) are highly variable after LOC. In contrast, slow oscillation power increased abruptly at LOC and remained elevated throughout the rest of the recording (Fig. 2A). Therefore we concluded that onset of power in the slow oscillation band is associated with the transition into unconsciousness, whereas other oscillatory features do not reach a steady state until minutes later and may reflect dynamic neural shifts at varying concentrations of propofol.

**Neuronal Spiking Becomes Phase-Coupled to the Slow Oscillation at LOC.** Studies of deeply anesthetized animals have shown that neuronal spike activity is coupled to the phase of the slow oscillation (8, 15, 20). We examined whether this spike–phase relationship developed immediately at LOC and whether it was maintained consistently thereafter. In each patient, population spike activity after LOC was significantly phase-coupled to the LFP slow oscillation (0.1–1 Hz), with 46.6% of spikes from all units occurring near the trough of the slow oscillation, during a phase of 0 to π/2 (maximum spiking at a phase of π/20–4π/20). Phase-coupling developed within seconds of LOC (between ~2.5 and 7.5 s; Table S1) and persisted throughout the ensuing changes in spike rate (Fig. 3 and Fig. S2). Spikes also were phase-coupled to the slow oscillation in the nearest ECoG channel but at a significantly different phase (maximum phase = 0 to π/10; P < 0.001, Kolmogorov–Smirnov test), suggesting that the LFP slow oscillation has a different relationship to spiking than the nearby, larger-scale ECoG recording. These results support the hypothesis that spikes become phase-coupled to the slow oscillation at LOC.

When examining individual units, most (67.2% of the 183 units with post-LOC spiking) were significantly phase-coupled to the LFP slow oscillation (P < 0.05, Pearson’s χ² test). When this analysis was restricted to units with post-LOC spike rates over 0.1 Hz, 94.0% of units had significant phase coupling (P < 0.05, n = 50, Pearson’s χ² test). Of the units without significant phase-coupling, 65.0% also showed peak spiking activity within a phase of 0 to π/2, demonstrating that most units had the same phase-coupling trend. These results demonstrated that after LOC nearly all spiking activity is tightly coupled to the slow oscillation phase and is suppressed for a large portion of the slow oscillation cycle. We refer to these periods of high spiking as “ON” states and the silent periods as “OFF” states to remain consistent with previous work using only extracellular recordings (21, 22). Because of the alternation of ON and OFF states, spike activity was limited to periods of a few hundred milliseconds, interrupted by periods of silence that also can last hundreds of milliseconds. Therefore we concluded that the slow oscillation marks a temporal fragmentation of cortical spiking that occurs at LOC.

**Slow Oscillation Impairs Information Transfer Between Distant Cortical Regions.** Given that post-LOC spiking is interrupted periodically within a cortical region, we investigated whether communication across distant areas also was affected. We examined slow dynamics across the grid of ECoG electrodes in the two patients (A and B) for whom we had at least 3 min of post-LOC ECoG data. Because spiking was strongly coupled to slow oscillation phase, we examined how this phase varied across the brain to infer the relative timing of neuronal activity in different cortical regions.

We quantified the phase relationships between different cortical regions using the phase-locking factor (PLF), which characterizes the phase offset between two oscillations over a period (23). The PLF magnitude ranges between 0 and 1 and quantifies the stability of the phase offset (1 reflects constant phase offset; 0 represents variable phase offset). The PLF angle indicates the average phase offset. We calculated the PLF between every pair of ECoG channels on the grid (8 × 4 or 8 × 8 cm, n = 96 total
electrodes) to determine the relationship between local and distant slow oscillations. We found that the PLF magnitude was conserved between the pre- and post-LOC states (correlation coefficient $R = 0.66$, patient A; $R = 0.88$, patient B; $P < 10^{-50}$ for each, t test), with a small but significant increase in PLF magnitude after LOC (mean increase $= 0.02 - 0.07$, $P < 0.01$, Wilcoxon signed rank test) (Fig. 4A). This result was consistent with previous findings that low-frequency correlations in neural activity are maintained after LOC (24, 25) and suggests that LOC is associated with only a slight shift in the strength of phase relationships between slow oscillations in different areas.

We next examined how the PLF varied with distance to determine whether slow oscillations in different cortical areas were at different phases. The PLF magnitude dropped significantly with distance ($R = -0.61$, patient A; $R = -0.82$, patient B; $P < 10^{-16}$ for each) (Figs. 4B and 5C and Fig. S3), demonstrating that the phase offsets between distant slow oscillations were variable. We also examined the mean phase offsets (PLF angle). Mean phase offsets between distant channels varied across a wide range, spanning 0 to $\pi$ (Fig. 4C and Fig. S3). Because a phase offset of just $\pi/4$ corresponds to a lag of ~250 ms, slow oscillations in distant ECoG channels had substantial timing differences. These results demonstrated that distant slow oscillations often were at different phases than the local oscillation, and these phase differences were not stable across time.

To examine how these phase offsets would affect neuronal activity, we examined the phase relationship between local spiking and slow oscillations measured across the ECoG grid. We measured spike phase-coupling as a modulation index (MI) quantifying the Kullback–Liebler divergence, in bits, between the observed phase distribution and a uniform distribution (Methods). A large MI indicates a strong relationship between local spiking and ECoG phase, whereas an MI of zero indicates no relationship. In the pre-LOC period, MI values were consistently small across all ECoG channels (MI range: 0.001–0.04 bits) (Fig. 5D), demonstrating that slow oscillation phase was not associated with strong suppression of spiking in the pre-LOC period. After LOC, the MI was significantly more variable across channels (range: 0.006–0.62 bits, $P < 0.01$ in each patient, Levine’s test). Spikes were strongly phase-coupled to the slow oscillation in the nearest ECoG channel, and this relationship declined significantly with distance ($R = -0.40$, patient A; $R = -0.68$, Patient B; $P < 0.001$ in each patient) (Fig. 5B and D and Fig. S4).

Taken together, our analysis of phase–phase and spike–phase coupling show that the post-LOC state is characterized by periodic and profound suppression of spiking coupled to the local slow oscillation phase and that this phase is not consistent across cortex. Given the strong relationship between phase and ON/OFF periods, this result suggests that, after LOC, ON periods in distant (>2 cm) cortical regions occur at different times (Fig. 5E, Right). In contrast, low-frequency oscillations in the pre-LOC state are not associated with strong suppression of spiking, so neurons are able to fire at any phase of local or distant slow oscillations despite the presence of phase offsets (Fig. 5E, Left). The combination of phase offsets and strong phase-coupling of spikes that occurs at LOC therefore is expected to disrupt communication between distant cortical areas, because one cortical area frequently will be profoundly suppressed when another area is active.

Although spikes were not strongly phase-coupled to distant slow oscillations during the post-LOC period, several electrodes located more than 3 cm from the spike recording site showed a statistically significant relationship (Fig. S5). In these cases, phase-coupling was weak, and the phase of maximal spiking was shifted, consistent with our conclusion that distant cortical regions are unlikely to have simultaneous ON periods. However, this finding raises the possibility that, despite the asynchrony of slow oscillations across the brain, there might still be a link between slow oscillations in distant cortical regions. Given the observed phase offsets (which ranged up to $\pi$), such coupling would occur frequently over hundreds of milliseconds and would not reflect precisely timed inputs and interactions. Overall, these analyses support the conclusion that distant cortical regions frequently were at a suppressed phase of the slow oscillation when the local
network was active. Therefore activity within a cortical area was isolated, impairing communication between distant regions.

**Local Network Structure Is Preserved After LOC.** Having observed interruptions in local activity and disruption of long-range communication, we examined whether connectivity within the local cortical network also was impaired. We fit a generalized linear model (GLM) to spike activity from the ensemble of units to test whether spiking could be predicted by the slow oscillation phase alone or whether the history of local network activity also contributed (26). We used the Bayesian Information Criterion (27) to select the number of covariates to include in the model. In each patient, we found that this model included $>30$ ms of population spike history (Fig. 6A and Fig. S6). Ensemble spike history therefore predicted future spiking, demonstrating that, although cortical activity was limited to brief ON periods, interunit structure existed within these periods. This pattern resembled the pre-LOC state, in which recent spike history (0–48 ms) was predictive of future spikes and more distant spike history contributed less (Fig. S7). This result suggests that, after LOC, cortical activity is not reduced to disordered spiking during ON periods. Instead, significant structure is maintained between nearby neurons during their brief periods of activity.

Structure between single units was reflected further in a peak in the cross-intensity function between several pairs of units, demonstrating millisecond-scale synchronization of spike activity (Fig. 6B and C). To examine whether pairwise synchronization persisted after LOC, we analyzed the cross-correlation between the 15 units with the highest spike rates in patient A. Of the 103 pairs (excluding pairs recorded on the same electrode), 21 were significantly correlated before LOC ($P < 0.05$, exact Poisson test relative to baseline from shuffled spikes, Bonferroni correction for multiple comparisons). After LOC, 71.4% of these pairs remained significantly correlated. In contrast, a significantly smaller number (only 18.3%) of pairs that were not correlated before LOC became correlated after LOC ($P < 10^{-3}$, Fisher’s exact test). This result demonstrated that pairs of units tended to retain the same correlation structure after LOC that they had before LOC, whether it was the presence or absence of a correlation. Taken together, both the GLM and paired correlation results show that significant interunit connectivity is maintained within post-LOC ON periods. This result suggests that the dominant change after LOC is the isolation of cortical networks, whereas aspects of local network structure may remain unaltered.

**Spiking Activity Is Associated with Modulations in Slow Oscillation Shape and Higher Frequency Power.** The mechanisms underlying the slow oscillation are debated (20, 28–31). Therefore, we examined the relationship between spike activity and slow oscillation shape in greater detail. We calculated an average LFP triggering at the beginning of ON periods. The triggered average demonstrated that ON periods begin at the minimum of the LFP slow oscillation (Fig. 7B and C). In addition, the LFP slow oscillation was asymmetric (Fig. 7C), with a higher peak after spiking than before spiking (mean difference $= 40.7 \mu V$, $P < 10^{-3}$, $P < 0.005$ for each patient, $t$ test). We tested whether this asymmetric shape occurred on all cycles of the slow oscillation or was specific to cycles with high spike activity. We compared cycles of the LFP slow oscillation that contained spikes with cycles that did not, matching the amplitudes of the slow oscillation minimum. Cycles that were not associated with spikes were symmetric (mean difference $= 0.3 \mu V$, $P > 0.9$, $t$ test), whereas those that were associated with many spikes produced a higher peak after spiking (mean difference $= 32.2 \mu V$, $P < 0.001$, $P < 0.05$ for each patient, $t$ test) (Fig. 7D). This asymmetry did not extend to the nearby ECoG recording (Fig. 8S), suggesting that the relationship between spike activity and slow oscillation shape is a highly local effect limited to less than 1 cm (i.e., the spacing in the ECoG grid). These results demonstrated that high spike rates are associated with an increased slow oscillation peak in the LFP, potentially reflecting enhanced suppression after spike activity.

Because low gamma ($25–50$ Hz) power also increased after LOC (Fig. S1 and Table S1), we examined its relationship to spike activity as well. ON periods were associated with significantly increased broadband ($<50$ Hz) power in the LFP and ECoG ($P < 0.05$, F-test, Bonferroni correction for multiple comparisons across frequencies) (Fig. 7A and Fig. S8). LFP power in alpha, beta, and gamma bands was significantly higher in slow oscillation cycles with high spike activity than in cycles with low spike activity.

![Figure 4](Image)
(P < 0.05, F-test, Bonferroni correction for multiple comparisons across frequencies) (Fig. 7E). These results showed that, in addition to the slow changes in gamma power occurring over minutes (Fig. S1), gamma power also fluctuated at the timescale of the slow oscillation (0.1–1 Hz) and was higher during ON periods. Therefore we concluded that after LOC power in the low gamma range is associated with high local spike rates. This result suggested that the gradual increase in gamma power after LOC may be related to the post-LOC fluctuations in spike rate rather than reflecting dynamics induced specifically at LOC.

**Discussion**

In summary, we found that rapid induction of unconsciousness using propofol causes the human brain to undergo an abrupt change in network dynamics, tipping into a new state in which neuronal activity is coupled to slow oscillations in the LFPs. Neural dynamics can be highly variable during the unconscious period, as spike rates and most oscillatory patterns continue to fluctuate for minutes after LOC (Fig. 1 and Fig. S1). The slow oscillation has a markedly different pattern: It develops simultaneously with LOC and maintains this increase thereafter (Figs. 2 and 3). Spiking activity is constrained to brief time periods coupled to the phase of the slow oscillation (Figs. 3 and 5), interrupting information processing within a cortical area. Moreover, these brief activity periods are phase-shifted across cortex (Figs. 4 and 5), limiting activity spatially, because different cortical areas are likely to be active at different times. However, functional connectivity within the local network is preserved...
The slow dynamics... 

Potential Circuit Mechanisms Underlying the Slow Oscillation. The mechanisms underlying the slow oscillation are unclear. One hypothesis is that the slow oscillation is cortically generated (20, 28, 29), but others suggest that it results from an interaction between cortex, thalamus, and thalamic reticular nucleus (30, 31). The relationship identified here between spike activity and slow oscillation shape suggests that cortical spiking may have a causal role in the slow oscillation. Spikes predict a high-amplitude peak in the LFP slow oscillation (Fig. 7D), but this effect does not extend to the ECoG recordings, which integrate activity from a larger population of neurons. The highly local nature of this effect suggests that cortical spiking may affect the local slow oscillation directly. One possible mechanism is that pyramidal neuron spiking during ON periods excites GABAergic interneurons, whose inhibitory actions are enhanced by propofol, driving the local network into a more hyperpolarized state. Another possibility is that spike activity may drive disfacilitation of cortical neurons, a mechanism that has been demonstrated in slow-wave sleep (33). These effects could be consistent with either the cortical or corticothalamic hypothesis.

Slow Oscillations in General Anesthesia and in Sleep. The slow oscillation during propofol-induced unconsciousness shares several features with slow waves during sleep: In both states, spike activity is coupled to a local slow oscillation that is not synchronous across the brain (21). The asynchronicity observed here (Figs. 4 and 5E) contrasts with previous observations in anesthetized animals (34) and is most likely caused by the increased spatial sampling provided by the 8-cm grid of intracranial electrodes. In addition, the preservation of pre-LOC neuronal network properties after LOC (Fig. 6) is consistent with the hypothesis that cortical UP states during sleep have dynamics similar to the waking state (35).

However, the patterns observed under propofol also show striking differences from patterns during sleep. The onset of the slow oscillation during induction of general anesthesia was abrupt (Figs. 2 and 3), accompanying rapid LOC caused by the bolus administration of propofol. Because general anesthesia typically is induced with a bolus of propofol, this abrupt transition into the slow oscillation is likely to occur in the majority of clinical patients when they lose consciousness during general anesthesia. During sleep, the slow oscillation develops over minutes, consistent with the gradual nature of the transition into sleep (36, 37). In both cases, slow oscillation dynamics temporally track LOC, further supporting the proposal that the slow oscillation represents a breakdown of cortical communication.

In addition, we found that periods of spike activity were brief (Fig. 5E), whereas sleep is characterized by persistent spiking with brief periods of suppression during slow-wave events (21, 38–40). This observation was corroborated recently in a study of sleeping and anesthetized cats (41). A difference in the ratio of UP and DOWN states could provide one explanation for why propofol creates a more profound disruption of consciousness than sleep: There is less temporal overlap in neuronal spiking between different cortical regions, more reliably preventing the organization of large-scale population activity. Furthermore, recent findings that isolated OFF states in sleep-deprived rodents are associated with behavioral impairment (42) are consistent with the hypothesis that the spatial and temporal properties of OFF states affect cortical function.

Potential Role of Slow Oscillations in Unconsciousness. Our results show that the slow oscillation appears abruptly at the onset of propofol-induced LOC. In addition, we demonstrate that the slow oscillation marks a state in which neuronal networks are
slow oscillation in unconsciousness. Furthermore, this experiment defines LOC as loss of voluntary response but cannot disambiguate whether the ability to respond may be suppressed before LOC. Given the slow oscillation’s association with prolonged and asynchronous periods of nearly complete suppression of neuronal activity, it seems unlikely that it is compatible with conscious processing. However, other possible mechanisms for propofol-induced unconsciousness include coherent frontal alpha rhythms that limit thalamocortical function (11, 12, 43). Future work in animal models could test whether the slow oscillation is sufficient to produce unconsciousness.

**Generalizability of Propofol-Induced Slow Oscillations.** A limitation of this study is that we enrolled patients with epilepsy, and it is possible that their cortical networks differed because of seizure foci or medication history. However, several factors support the hypothesis that these results generalize to the healthy brain. First, the microelectrodes were located at least 2 cm from the seizure focus in each patient, and histology did not reveal any disruption of the local network, suggesting that the LFPs and single units were recorded from healthy cortex. Second, the overall effects of propofol were highly consistent with those observed in healthy subjects: Unconsciousness was associated with increased slow oscillation power and increased gamma power, in strong agreement with previous studies (9). These results suggest that propofol acts similarly in healthy and patients’ brains. Finally, we report statistics for each individual patient and show that the timing of the slow oscillation onset and its relationship to spiking were replicated across patients despite their individual clinical profiles. Because epilepsy is a heterogeneous disease with different cortical origins, the high consistency of these results suggests that the effects reported here are not caused by the presence of epilepsy. These three observations suggest that our results are not a product of an epileptic brain but rather reflect a true neural correlate of LOC that is likely to generalize to the healthy brain. Future studies in patients with pathologies other than epilepsy could address this issue further.

**Role of Slow Oscillations in Other Brain States.** Other anesthetic drugs, such as ketamine and dexmedetomidine, operate through molecular and neural circuit mechanisms different from those of propofol (4). The study presented here provides a framework for studying these other drugs further and identifying how they influence neural dynamics to produce altered states of arousal. In addition, further work will be needed to explore how these findings may relate to other conditions, because slow-wave activity is a common feature in conditions such as coma (44) and complex-partial seizures (45), and those slow waves share some characteristics with the slow oscillations studied here. Although a unitary mechanism for unconsciousness under general anesthesia is technically possible, it is more likely that a variety of mechanisms exist that can produce unconsciousness (2). We have shown here that the slow oscillation is a fundamental component of propofol-induced unconsciousness, marking an impairment of cortical integration at both the local and global scale.

**Methods**

**Data Collection.** Three patients with epilepsy intractable to medication were implanted with intracranial subdural electrocorticography electrodes for standard clinical monitoring (AdTech). Informed consent was obtained from all patients in accordance with the Partners Human Research Committee. ECoG electrode placement was determined solely by clinical criteria, and the electrodes were located in temporal, frontal, and parietal cortices. Individual ECoG electrodes within a grid were spaced 1 cm apart. In addition, a 96-channel NeuroPort microelectrode array with 1.0-mm-long electrodes (BlackRock Microsystems) was implanted into the superior (patient B) or middle (patients A and C) temporal gyrus to record LFPs and ensembles of single units for research purposes. In each patient, the Neuroport array was located at least 2 cm from the seizure focus. There was no evidence of fragmented, impairing both local and long-range communication. However, the scope of this study, performed in human subjects, does not allow us to test explicitly the causal role of the
disruption in local network structure based on the firing properties of the neurons or the postsection histological examination of the area around the LFP recordings were collected at the beginning of surgery to ex.
plant the electrodes. Anesthesia was administered as a bolus dose of pro-
profol according to standard clinical protocol. All propofol doses were based
on the anesthesiologist’s clinical judgment rather than the research study
considerations. Patient A received three boluses (130 mg, 50 mg, and 20 mg),
patient B received one bolus (200 mg), and patient C received one (150 mg).
After induction, patients were transferred to a continuous i.v. infusion of propofol to maintain anesthetic levels. Throughout the induction period,
patients responded to auditory stimuli (prerecorded words and the patient’s
name) with a button press, and stimuli were presented every 4 s to obtain
precision for LOC time on the order of seconds. LOC time was defined as the
period from −1 to 4 s surrounding the first stimulus after the patient com-
pletely ceased responding. Spike sorting was carried out according to stan-
dard procedures (46) with Offline Sorter (Plexon) and produced 198 single
units for further analysis. LFPs were referenced to a wire distant from the
microelectrode array and were collected with hardware filters bandpassing
between 0.3–7.5 kHz with a sampling rate of 30 kHz. LFPs then were low-
pass-filtered at 100 Hz and resampled to 250 Hz. For display, raw time-series
were low-pass-filtered with a finite-impulse response filter with 4,464 co-
efficients, achieving unit gain between 0 and 40 Hz and attenuation of more
than −300 dB above 40 Hz. ECoG recordings were collected with a sampling
rate of either 250 Hz (patients B and C) or 2,000 Hz (patient A), in which case
it was low-pass-filtered at 100 Hz and resampled to 250 Hz. ECoG recordings
were referenced to an intracranial reference strip channel when available
(patient A) and otherwise to an average reference. In patients A and B, ECoG
recordings were collected throughout. In patient C, the microelectrode
recordings were collected throughout, but the ECoG recording ended −100 s
after IND. Significant changes of spike rate could not be assessed in ECoG channels, because the spike rate was nearly zero during this time. Two ECoG grid channels were rejected in patient A
because of large artifacts. All data were exported to Matlab (Mathworks) for
analysis with custom software.

Spike Rate Analysis. Spike rates and confidence intervals were computed with
Bayesian-space estimation (47). To minimize any error caused by un-
stable recordings, the spike rate analysis excluded units that were not con-
fidently detected throughout the entire baseline period (8.1%). The
computed spike-rate effects were similar when these units were included.
Periods of silence were compared with a simulated Poisson distribution of
equal rate over each 10-s period, and significance was assessed for each
patient with a χ^2 test relative to that distribution.

Spectral Analysis. Spectrograms were calculated with multitaper methods
using the Chronux toolbox (http://chronux.org/; for bandwidth settings, see
SI Methods). Power changes after LOC were computed as the percent change
in the period 30–60 s after LOC relative to the period 30–60 s before
LOC. Ranges in the text reflect ranges across patients. The slow oscillation
was extracted by applying a symmetric finite impulse-response bandpass
filter, bandpassing between 0.1 and 1.25 Hz. The spectral power contribution
below 0.3 Hz was minimized. Phase was extracted with a Hilbert transform.
Statistical testing of triggered spectrograms was done by taking a ratio of each
χ^2 distribution, and significance was calculated as an F-test with a Bonferroni correction for multiple comparisons across frequencies. For
comparing spectra during and before an ON period, power spectra from 250 ms
after ON period onset were compared with spectra from 250 ms before
the ON period onset. Averaged LFP waveforms were compared by pre-
selecting a time period and performing a t test on the mean amplitude
distribution of the peak values within that interval. When comparing the waveform height before and after spiking, a t test was performed comparing the mean amplitude in
the time window from −750 to 500 ms and in the time window from 500–
750 ms locked to ON period onset or slow oscillation minimum. The

Phase Modulation. Significance for single-unit phase-coupling was computed with
a χ^2 test on the binned phase distribution. The analysis was performed
a second time on only cells with spike rates above 0.1 Hz, ensuring that there
were at least five expected spikes per phase bin. Strength of phase modu-
lation was computed as an MI (48) adapted to quantify the Kullback–Li
dier divergence of the phase histogram from the uniform distribution, measured
in bits. Spike phase was split into a phase histogram (p) of 10 bins, and MI
was computed as \(\sum_i p_i \log p_i + \log 10\). We also computed the χ^2
statistic as an alternative measure, yielding similar results. MI significance for each
ECoG channel was calculated by shuffling the entire spike train randomly
between 2 and 10 s and calculating a shuffled MI over 2,000 random shifts.
The empirical MI then was compared with the shuffled MI with a signifi-
cance level of 0.05 and a Bonferroni correction for multiple comparisons
across channels. For LFP phase analysis, each single unit was compared with
its local LFP channel. The time-varying phase modulation was computed
with a window of 20 s sliding every 5 s. To assess the phase of maximal
spiking relative to the ECoG slow oscillations, the phase of spiking was di-
vided into 20 bins, and then the mode of the phase histogram was reported.

Timing of Spike Rate and Spectral Power Changes Relative to LOC. We tested
spike rates and spectral power to determine the first time bin in which these
features differed significantly from the baseline period before propofol
administration. We compared every time point, starting 30 s before LOC, with
a baseline of spike rates or spectral features from the 3-min baseline period
immediately preceding it. To assess spike rate significance, we used a Bayesian
analytical approach in which each baseline time point was compared with
samples drawn from the Gaussian distribution of the baseline period and tested
for a significant difference. This baseline sampling distribution was computed
with the same state-space algorithm used to calculate spike rates (47). To de-
terminate the time at which power at a given frequency differed significantly,
we used an analogous method but replaced the Gaussian sampling distribution
with a χ^2 distribution, which is the appropriate distribution for power mea-
ures. The time bin in Table S1 lists the earliest point at which the value had
a 95% probability of being higher than the baseline period. We could not
construct a similar sampling test for the MI because its distribution is not
known, so we did not resample the baseline and instead reported the time
at which MI became higher than the mean of the baseline period plus two SDs.
For all these measures, 5-s nonoverlapping bins were used to identify the time at
which changes occurred relative to LOC, which is the period from −2.5 to 2.5 s.

PLF. The PLF was computed to obtain a time-varying measure of phase offsets
between slow oscillations. The phase of the slow oscillation was extracted as
described in Spectral Analysis. For each time point, we then computed \(z(t) = \exp(-i\phi(t) - i\phi(0))\) where \(\phi(t)\) is the phase of one ECoG slow oscillation
at each time point and \(\phi(0)\) is the phase of another ECoG slow oscillation. The
PLF magnitude \(|z(t)|\) was calculated as the magnitude of \(z(t)\) across the peri-
period from 2.5 to 2.5 s. We tested for the significance of the PLF magnitude
using a Bonferroni correction for multiple comparisons across channels. For
the post-LOC period. To assess the variability of phase offsets, we calculated
the magnitude of the PLF. The distribution of PLF magnitude was assessed by plot-
ing the mean and SD of the PLF magnitude across each pair of ECoG channels
separated by a given distance (the distance between channels computed geo-
metrically across the grid). To determine the mean value of the phase offset
across time, we calculated the angle of the PLF. We then plotted the distribution
of mean phase offsets across all pairs of channels separated by a given distance,
by taking a 2D histogram of PLF angle values for all electrode pairs. Accompa-
nying reconstructed brain showed individually localized electrode positions (49).

GLM Fitting. A GLM was fit to ensemble spiking using custom software
that performed regression with Truncated Regularized Iteratively Reweighted
Least Squares (TR-IRLS) (26, 50) and using the Bayesian Information Criterion
to determine the best model. After Akaike’s Information Criterion (AIC) had yielded
a significant contribution of spike history. The GLM was constructed to predict
elementary spiking, which was defined as a series of 12-ms bins that
contained a 1 if any spikes from any units occurred in that period and
a 0 otherwise. Ten covariates were used to represent the range of possible
LFP phase values. Amplitude was normalized to range between 0 and 1.
Because individual unit spike rates are low, the history-dependent terms in
this model predominantly reflect interactions between units. The version
presented is with 12-ms bins of spike history; similar results were obtained
when using 4- or 8-ms bins. We excluded the minute surrounding LOC to
ensure that any correlation between pre-LOC and post-LOC analyses did not result
from bias from adjacent recordings during the LOC transition. For
detailed equations, see SI Methods.

Single-Unit Correlations. Single units with high post-LOC spike rates were
selected for correlation analysis to ensure sufficient spikes to assess
the significance of their correlations. The minute surrounding LOC was excluded
to reduce bias that could result from comparing adjacent recordings. Cor-
relations between single units were computed relative to a shuffled baseline,
to examine fine time-scale synchronization beyond the changes in population
spike rate induced by the slow oscillation. Spike times were randomly shuffled
200 times, between 50 and 500 ms, to obtain a baseline of correlated spike
rate without millisecond-level timing information. Paired correlations then
were tested for significance between −100 and 100 ms, with \(P < 0.05\) using
a Bonferroni correction for multiple comparisons across lags. Correlations

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were judged significant if they had a \( P < 0.05 \) departure from the Poisson distribution of spike occurrence predicted by the shuffled baseline. The relationship between pairs of single units was visualized with the square root of the estimate of the cross-intensity function (51). Fisher’s exact test was performed in R statistical software (http://www.r-project.org/).

Detecting Initiation of ON Periods. ON periods were detected by binning spikes from all units in 50-ms time bins and then setting a threshold to detect local peaks in the spike rate. The threshold was determined manually for each patient by visually checking to ensure adequate detection, because the number of units and thus expected population spike rates differed in each patient. After detection, the first spike within 300 ms of ON period detection was taken as the initiation time, and spike histograms verified that these times represented initiation of spiking. These ON period initiation times then were used for subsequent analysis of slow oscillation spectra and waveform morphology.

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