Different effects of propofol and dexmedetomidine sedation on electroencephalogram patterns: Wakefulness, moderate sedation, deep sedation and recovery

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Abstract

Sedation induces changes in electroencephalography (EEG) dynamics. However, the distinct EEG dynamic characteristics at comparable sedation levels have not been well studied, resulting in potential interpretation errors in EEG monitoring during sedation. We aimed to analyze the EEG dynamics of dexmedetomidine and propofol at comparable sedation levels and to explore EEG changes with increased sedation levels for each agent. We measured the Bispectral Index (BIS) and 20-channel EEG under dexmedetomidine and propofol sedation from wakefulness, moderate sedation, and deep sedation to recovery in healthy volunteers (n = 10) in a randomized, 2-day, crossover study. Observer’s Assessment of Alertness and Sedation (OAA/S) score was used to assess sedation levels. Despite similar changes in increased delta oscillations, multiple differences in the EEG spatiotemporal dynamics were observed between these two agents. During moderate sedation, both dexmedetomidine and propofol induced increased spindle power; however, dexmedetomidine decreased the global alpha/beta/gamma power, whereas propofol decreased the alpha power in the occipital area and increased the global spindle/beta/gamma power. During deep sedation, dexmedetomidine was associated with increased fronto-central spindle power and decreased global alpha/beta/gamma power, but propofol was associated with increased theta/alpha/spindle/beta power, which was maximized in the frontal area. The transition of topographic alpha/spindle/beta power distribution from moderate sedation to deep sedation completely differed between these two agents. Our study demonstrated that there was a distinct hierarchy of EEG changes with increased sedation depths by propofol and dexmedetomidine. Differences in EEG dynamics at the same sedation level might account for differences in the BIS value and reflect the different sedation mechanisms. EEG-based clinical sedation monitoring should consider the effect of drug types on EEG dynamics.
Introduction

Sedation is a state of altered consciousness induced by different kinds of anesthetics. Sedation can provide a comfortable experience for the patient and a better operating condition for the clinician during unpleasant diagnostic or therapeutic procedures, and it is now widely performed in operating rooms and non-operating sites[1–3]. Sedation ranges from minor anxiety, amnesia, and hypnosis to unconsciousness according to the clinical requirements. Conscious sedation is a state that allows patients to tolerate unpleasant procedures when they are awake and collaborative, while deep sedation provides a condition of no response to auditory stimuli or noxious stimulation[4, 5]. The graded fashion of conscious experience might reflect hierarchical brain dynamics. Electroencephalogram (EEG) measures cortical brain activity. However, more studies have focused on the transition from wakefulness to unconsciousness, and the different levels of sedation are not well characterized by EEG[6, 7].

Sedatives might act at different molecular targets and neural circuits to produce distinct EEG traces[8]. The common sedatives that alter arousal states include gamma-amino butyric acid type A (GABA\textsubscript{A}) receptor agonists, opioid receptor agonists, N-methyl D-aspartate receptor (NMDA) antagonists and \(\alpha\)2 receptor agonists. Previous studies have demonstrated a drug-dependent EEG trace during sedation. Propofol, an agonist of GABA\textsubscript{A} receptors, induces loss of consciousness (LOC) and is characterized by an abrupt anteriorization of alpha rhythms[6,9–11]. This distinct EEG variation is also observed during halothane, isoflurane, sevoflurane, and desflurane anesthesia[12–14]. Ketamine, an NMDA antagonist, results in “gamma bursts” and markedly increases the theta power across the cortex[7, 15]. Dexmedetomidine is a highly selective \(\alpha\)2-adrenergic receptor agonist. Dexmedetomidine-induced sedation resembles non-rapid eye movement (NREM) sleep with a characteristic spindle wave (12–16 Hz) in the frontal area[16, 17]. EEG dynamics also change with the agent dose and sedation level. Moderate propofol sedation increases EEG oscillations in the spindle range (12–15 Hz) and beta range (13–25 Hz) in frontal areas, and deep sedation is associated with delta oscillations across the cortex and alpha anteriorization[11]. Ketamine at a dose of 0.5 mg/kg is characterized by reduced posterior alpha power, and 1.5 mg/kg is associated with increased delta, theta and gamma power across the cortex[15]. However, whether the above EEG dynamics are comparable at the same clinical sedation level for different sedatives and whether EEG parameters are suitable for monitoring the depth of sedation remain unknown.

Describing EEG dynamics during a comparable sedation level for different sedatives might help explain the possibility of EEG monitoring during sedation and, to some extent, explore sedation mechanisms. Propofol and dexmedetomidine are two popular sedatives. Although the two agents can induce similar sedation levels by clinical evaluation, there is a significant difference in Bispectral Index (BIS) monitoring at the same sedation level[18]. This study aimed to determine the effect of propofol and dexmedetomidine on EEG spectral power parameters and EEG topography within specific frequency bands during different depths of sedation. We hypothesized that propofol and dexmedetomidine induce different brain dynamics at comparable sedation levels and that a distinct hierarchy of EEG changes exists with the increased sedation level for each agent. To explore these hypotheses, we measured and analyzed the EEG patterns of 10 healthy volunteers who received both propofol and dexmedetomidine sedation in a crossover study.

Materials and methods

Subjects and EEG data collection

The Ethics Committee of Beijing Tongren Hospital affiliated to Capital Medical University provided ethical approval for this study on 19, October 2015 (Chairperson Prof. Ningli Wang,
protocol number TRECKY2015-021). The study was registered with the Chinese Clinical Trial Registry (ChiCTR-OON-16008369). Ten healthy male volunteers (ASA physical status I, right-handed, non-smoking, normal weight) aged 23 to 30 were enrolled in this study after public advertisements. Each volunteer was informed of the consent policies, and written consent was obtained. All subjects underwent a physical examination, including an interview, physical status assessment, laboratory testing and a 12-lead ECG examination. Subjects were not allowed to use alcohol or any medication 48 hours before the study, and all fasted for at least 8 hours. All experimental procedures were scheduled from 9 am to 12 pm in a standard operating room to avoid the influence of circadian rhythm on sedation. We performed a randomized, 2-day crossover study design for propofol/dexmedetomidine sedation, with one sedation procedure on the first day and the other sedation procedure two weeks later. Volunteers were assigned to receive either propofol or dexmedetomidine according to a computer-generated randomization during the first sedation procedure. Oxygen supplement (2 l/min) was provided via a mask. Sedation monitoring included the BIS Index (BIS VISTA™ monitor, software 2.00, Aspect Medical Systems, Newton, MA), noninvasive arterial pressure (NIBP), a three-lead electrocardiogram, pulse oximetry, and respiratory rate.

After resting for 5 min, the volunteers were instructed to close their eyes and listen to a 2-min auditory stimulus. For dexmedetomidine sedation, dexmedetomidine (dexmedetomidine hydrochloride solution, 100 μg/ml, Hengrui, Jiangsu, China) diluted to 4 μg/ml was administered intravenously with a loading dose of 1 μg/kg over 10 min and a maintenance dose of 0.4–1 μg/kg/h. Propofol sedation was given via target-controlled infusion (TCI) with the pharmacokinetic parameters obtained from Schnider[19]. The target effect-site concentration of propofol was set beginning at 1 μg/ml and was then increased by 0.3 μg/ml per step. The 5-point Observer’s Assessment of Alertness/Sedation Scale (OAA/S) was used every 3 min to determine the sedation level in each case and to help titrate the infusion rate[18]. Wakefulness was defined as volunteers without any drug administration, a moderate sedation state was defined as OAA/S = 3 (responds only after name is spoken loudly and/or repeatedly), deep sedation was defined as OAA/S = 1 (does not respond to mild prodding or shaking), and recovery was defined as OAA/S = 5 (responds readily to name spoken in a normal tone)[18].

The CogniTrace 20-channels system (A.N.T., Netherlands) was applied to record the continuous EEG and play the auditory stimuli. The EEG signals from 20 sites were based on the international 10–20 system. All EEG electrodes were referred to the mastoids (M1 + M2 average). The impedance of each electrode was kept below 5 kΩ. The auditory stimuli (2000 Hz, 75 dB, with intervals of one s) were repeatedly displayed and applied via earphones at wakefulness, moderate sedation, deep sedation and recovery. The study design is provided in Fig 1.

![Figure 1](https://doi.org/10.1371/journal.pone.0199120.g001)
EEG spectral analysis and topographic mapping

After data collection, the EEG was analyzed offline with EEG lab 14.0 software as Zhang et al. described[20, 21]. Briefly, all EEG data were filtered offline with a 0.5–100 Hz bandpass filter for analysis. Filtered EEG data were then visually inspected for artifacts. The artifact rejection methods consisted of the exclusion of epochs with a large amplitude (over ±100 μV), DC bias, blinks, and slow eye movement. Bad electrodes were substituted with the extrapolated virtue values from the neighboring channels. After artifact rejection, each set of EEG data was subjected to a 2-s epoch, and each epoch was processed using fast Fourier transformation (FFT) analysis to obtain the absolute power at each electrode in the following six bands: delta (0.5–4 Hz), theta (4–8 Hz), alpha (8–12 Hz), spindle (12–15 Hz), beta (15–25 Hz) and gamma (25–40 Hz). In each 2-min period, EEG was analyzed in 2-s epochs, resulting in 60 epochs. On average, approximately 50–58 valid epochs in each study condition were subjected to further analyses. Spectral power was estimated by FFT, with window lengths of T = 1 s with a 0-s overlap and a spectral resolution of 0.25 Hz. The topographic power distribution was calculated with the EEGLab toolbox[22].

Statistical analysis

Sedation parameters are expressed as the mean ± SD. Statistical analyses of the sedation parameters were performed using GraphPad Prism 6.01 (GraphPad Prism Software Inc., La Jolla, California, USA). The sedation times of the two agents were compared with unpaired t-tests. Two-way repeated measures analysis of variance (RM-ANOVA) and Bonferroni’s multiple comparisons test were used to assess the variations in BIS values in the drug regimes (dexmedetomidine and propofol) and the sedation level (wakefulness, moderate sedation, deep sedation and recovery state). Low power spectral density, \(10^{\log_{10}(\mu V^2/Hz)}\), was used to describe the absolute spectral power. For each participant, the power spectrum density and statistical analyses were conducted with the Matlab-based EEGLab toolbox[22]. According to previous literature and the current topography results, sedation states induce peak spectrum activity at the frontal midline electrodes. Therefore, the Fz electrode spectra were analyzed with two-way RM-ANOVA followed by planned pairwise comparisons between moderate sedation and baseline, deep sedation and baseline, recovery state and baseline as well as between deep and moderate sedation conditions for both dexmedetomidine and propofol. To explore the long-range coordination of neural activity, all 20 electrodes were included in the topographic analysis. The signals were divided into six frequency bands: delta, theta, alpha, spindle, beta and gamma. For each band, ANOVAs and planned comparisons similar to those used for the Fz spectral analyses were performed to explore the topographical differences. All statistical significances of the Fz and topographical analyses were calculated using a nonparametric bootstrap approach with a 10000 resampling size at alpha = 0.05. Multiple comparisons were corrected by the false discovery rate (FDR) procedure, a commonly used method for high-dimension data analysis.

Results

Participant characteristics and sedation parameters

Ten right-handed male volunteers took part in this study, but two could not reach the deep sedation state during the dexmedetomidine infusion. We did not increase the dexmedetomidine dosage because of safety issues. Therefore, data from the remaining 8 volunteers were included in the final analysis. The times from wakefulness to moderate sedation, from moderate sedation to deep sedation and from deep sedation to recovery during dexmedetomidine
sedation were 10.8 ± 2.5 min, 15.4 ± 6.8 min and 52.6 ± 14.6 min, respectively. The times from wakefulness to moderate sedation, from moderate sedation to deep sedation and from deep sedation to recovery during propofol sedation were 10.4 ± 3.5 min, 13.5 ± 5.3 min and 22.8.6 ± 5.4 min, respectively. There were no significant differences in the time from wakefulness to moderate sedation or the time from moderate sedation to deep sedation between dexmedetomidine and propofol sedation (P = 0.81 and P = 0.55, respectively); however, the time from deep sedation to recovery during dexmedetomidine sedation was significantly longer than that of propofol (P<0.001).

The BIS values at wakefulness, moderate sedation, deep sedation and recovery during dexmedetomidine sedation were 88.8 ± 5.0, 65.6 ± 7.1, 43.8 ± 5.3 and 85.1 ± 8.1, respectively. The BIS values at wakefulness, moderate sedation, deep sedation and recovery during propofol sedation were 87.2± 5.8, 73.6 ± 3.7, 53.6 ± 7.6 and 86.7 ± 9.0 respectively. The BIS values at moderate and deep sedation by dexmedetomide were significantly lower than those by propofol (sedation effect: P<0.001, drug effect: P = 0.019, sedation*drug interaction: P = 0.016, Fig 2).

Frontal spectral power changes from wakefulness, moderate sedation, and deep sedation to recovery during dexmedetomidine/propofol sedation

One volunteer’s raw EEG of Fz and spectral analysis of Fz during the sedation procedures are presented (Fig 3 and Fig 4). We observed changes in the spectral analysis that were induced by dexmedetomidine and propofol from wakefulness to recovery, and the changes tended to differ between these two agents. Statistical analysis showed significant differences in the ranges 0–8 Hz, 9.25–16.25 Hz, and 31.5–40 Hz for the main sedation effect, 10.5–30 Hz for the main drug effect, and 10–40 Hz for the interaction effect. Multiple comparisons showed that compared with wakefulness, the dexmedetomidine-induced spectral power of moderate sedation was larger at 0.75–6.5 Hz and 12.5–15.5 Hz and smaller at 8.5–11 Hz and 17.5–40 Hz, whereas the propofol-induced spectral power of moderate sedation was larger at 0.25–3.75 Hz and 11.75–40 Hz and smaller at 9–10.25 Hz. Additionally, the dexmedetomidine-induced spectral power of deep sedation was larger at 0.25–7.5 Hz and 12.5–14.5 Hz and smaller at 9.25–10.75 Hz and 15.75–40 Hz, while the propofol-induced spectral power of deep sedation was larger at 0.25–8.75 Hz and 10–35.5 Hz. Further, the dexmedetomidine-induced spectral power of
recovery was smaller at 8.25–12. 5 Hz, 16.5–23 Hz and 38.75–39.25 Hz, whereas the propofol-induced spectral power of recovery was smaller at 10–11.25 Hz. Compared with moderate sedation, deep dexmedetomidine sedation was associated with increased spectral power in the range of 0.25–10.75 Hz and decreased spectral power in the range of 13.5–40 Hz, while the
Deep propofol sedation was associated with increased spectral power in the range of 0.25–16.5 Hz and decreased spectral power in the range of 26–40 Hz. Thus, for these two agents, the spectral powers of the alpha, spindle, beta and gamma bands changed differently from wakefulness to recovery. Moreover, the spectral power changes from moderate sedation to deep sedation were different in the alpha, spindle, and beta bands.

**Topographic changes in EEG power from wakefulness, moderate sedation, and deep sedation to recovery during dexmedetomidine/propofol sedation**

Both dexmedetomidine- and propofol-induced topographic changes in the EEG power of different bands during the sedation procedure are represented in Fig 5. As typically observed, both dexmedetomidine and propofol were correlated with increased global delta band power, but changes in other bands were topographically distinct, especially in the alpha, spindle, beta and gamma ranges. Statistical analysis showed significant differences in all electrodes of the six bands for the main sedation effect, in all electrodes of the spindle and beta bands for the main drug effect and in all electrodes of the alpha, spindle, beta and gamma bands for the interaction effect.

To identify the spectral changes, differences between the two states are displayed as changes in power with its associated t-statistic in Fig 6. Moderate dexmedetomidine sedation was associated with a reduction in global alpha, beta, gamma band power and an increase in global delta, theta power and spindle power at the frontal-vertex sites; by contrast, moderate propofol sedation was associated with a reduction in occipital alpha power and increases in delta, spindle, beta and gamma power at the global cortex. During deep dexmedetomidine sedation, global delta and theta power continued to increase, and global beta and gamma power continued to decrease, whereas deep propofol sedation was associated with the sustained increase in global delta power and increased theta/alpha/spindle/beta power maximally in the frontal area. The alpha/beta oscillation power was persistently lessened across the entire cortex at the recovery state from dexmedetomidine, while there was a minor reduction in alpha power at the frontal and occipital areas with the recovery state from propofol. Compared with moderate sedation, dexmedetomidine increased the delta and theta power but decreased the spindle.
beta and gamma power across the cortex region during deep sedation, whereas propofol increased the global delta, theta, alpha band power, decreased the global gamma power, and strengthened the spindle power in the fronto-central area. The above data suggest that although the topographic changes in delta power were similar to those from wakefulness to recovery for the two agents, the powers of the other bands differed. The transition of topographic alpha/spindle/beta power distribution from moderate sedation to deep sedation between these two agents is completely different.

Discussion

In this study, we visualized gradual changes in EEG dynamics from wakefulness, moderate sedation, and deep sedation to recovery during propofol/dexmedetomidine sedation. We demonstrated that there was a distinct hierarchy of EEG changes with increased sedation levels for these two agents. Despite a similar change in sustained increase in delta oscillation from wakefulness to deep sedation, multiple differences in the EEG spatiotemporal dynamics were observed between the two agents. 1) During moderate sedation, both dexmedetomidine and propofol induced increased spindle power; however, dexmedetomidine increased theta power and decreased alpha/beta/gamma power across the whole cortex, whereas propofol decreased alpha power in the occipital area and increased global beta/gamma power. 2) During deep sedation, dexmedetomidine was associated with increased global theta power and fronto-central spindle power and decreased alpha/beta/gamma power across the whole cortex, but propofol was associated with increased theta/alpha/spindle/beta power, which was maximized in the frontal area. 3) The transition of topographic alpha/spindle/beta power distribution from moderate sedation to deep sedation completely differed between dexmedetomidine and propofol.

Different frontal spectral powers and BIS differences at the same sedation levels

Like natural sleep, sedation is a state of decreased arousal. The clinical manifestation ranges from drowsiness to unconsciousness. Sleep is characterized into rapid eye movement (REM)
and three stages of NREM sleep by the distinct EEG patterns. However, the application of EEG-based monitoring to measure the depth of sedation is controversial.

The BIS value is calculated from three components: spectral analysis, bispectral analysis and temporal analysis[23]. The frequency domain-based relative $\beta$ ratio is measured by the spectral analysis. The $\beta$ ratio is defined as the logarithm of the power ratios in two empirically derived frequency bands, $\log(P_{30-47Hz}/P_{11-20Hz})$, and is a major parameter for the BIS calculation of the sedation level[24]. In our study, a substantial difference in the frontal spectral power was observed in the 10–40 Hz range during the sedation procedures between dexmedetomidine and propofol. The frontal beta (15–25 Hz) and gamma (25–40 Hz) powers were decreased by dexmedetomidine and increased by propofol. This result sufficiently explains why the BIS value of dexmedetomidine was lower than that of propofol at comparable sedation levels in our study and in previous report[18, 25]. Additionally, it explains the decreased BIS value when dexmedetomidine was added to propofol anesthesia as well as the reduction in propofol required during BIS-guided closed-loop anesthesia when dexmedetomidine was added[26–28].

Different spatiotemporal alpha/spindle/beta/gamma oscillations during dexmedetomidine/propofol sedation

As the targets of propofol, GABA$_A$ receptors are widely distributed in the cortex, the thalamus and the preoptic area (POA) of the hypothalamus. The mechanisms underlying propofol-mediated sedation include enhancement of GABA$_A$ inhibition from inhibitory interneurons to excitatory pyramidal neurons, from the thalamus to the cortex and from the POA to the arousal centers, including the midbrain, the pons, and the hypothalamus[29–31]. Dexmedetomidine selectively acts on the $\alpha_2$- receptors of the locus coeruleus (LC) projecting to the POA, which activates the inhibitory outputs to the arousal centers and results in a sedative state[30, 32]. Recently, the thalamocortical system was shown to contribute to both propofol and dexmedetomidine-induced altered arousal[10, 33, 34]. However, the effects of propofol and dexmedetomidine on the thalamocortical system have never been compared.

Alpha rhythms (8–12 Hz) are spontaneous EEG oscillatory activities over the occipital-parietal cortex of awake humans in a relaxed state with their eyes closed, and these rhythms change with the arousal state and cognitive activities. Spindles (12–15 Hz) are one of the most dominant EEG oscillations during NREM stage 2 sleep, and they have a lower density in deeper slow-wave sleep (SWS). Both the alpha power and the spindle power change in all parts of the cortex from wakefulness to different stages of sleep[35–38]. The origin of spindle waves shares a similar mechanism with alpha waves, as the GABAergic reticular nucleus in the thalamus is the pacemaker of this spindle rhythm, and the thalamocortical neurons and cortical neurons potentiate the genesis of spindle waves[39–42]. Propofol-induced LOC is characterized by an abrupt anteriorization of alpha rhythms[6,9–11]. Using a Hodgkin–Huxley-based model, Vijayan and Kopell suggested that alpha rhythms arise via a specialized class of thalamocortical cells (TCs), so-called high-threshold thalamocortical neurons (HTCs)[43]. Based on the above model, a computational model suggested the following: 1) Propofol attenuates the occipital alpha power by silencing HTCs via reducing the hyperpolarization-activated current $I_h$. 2) Propofol potentiates the GABA$_A$ synaptic current and decay time from interneurons onto pyramidal neurons cells and from reticular neurons (REs) onto TCs, thus bringing alpha activity to the cortex, creating alpha oscillations between REs and TCs, and enhancing the cortical inputs to the thalamus. These alternations result in a reciprocal excitation between the cortex and thalamus and form a strengthened cortical-thalamo-cortical loop alpha oscillation[10, 44]. In our study, we confirmed the hierarchical changes in the propofol-induced reduction of occipital alpha oscillations during moderate sedation and propofol-induced alpha anteriorization.
The topographic representation of alpha activity changes by dexmedetomidine is similar to that of sleep, with attenuation of alpha power across the frontal-posterior cortex [35–37]. The effect of dexmedetomidine on the HTCs has not been described. The neurotransmitter norepinephrine enhances the hyperpolarization-activated current $I_h$ in thalamocortical neurons in vitro [45]. Dexmedetomidine decreases norepinephrine release from the LC, which might reduce the hyperpolarization-activated current $I_h$ and lead to attenuated alpha oscillation. Whether reducing the hyperpolarization-activated current $I_h$ is involved in the mechanisms of dexmedetomidine-induced sedation need. The lack of alpha anteriorization might be possibly due to the short GABA$_A$ potentiation by dexmedetomidine.

Both dexmedetomidine and propofol sedation result in the appearance of spindle activity [17, 46]. In our study, dexmedetomidine–induced spindle oscillations increased dramatically during moderate sedation and attenuated during deep sedation, which shared a similar tendency with spindles across varying depths of NREM [38]. For propofol, increased spindle activity from frontal to posterior areas was observed during moderate sedation, and deep sedation dramatically increased the spindle activity in the frontal regions. These results are consistent with a previous report that described propofol-induced LOC being accompanied by increased spindle oscillations in the frontal area with/without auditory stimuli [11]. Although the increased spindle oscillations share a similar topographic pattern as alpha oscillations during deep sedation, their differential characteristics during wakefulness and moderate sedation indicate the impossibility of overlapping analysis. The distinct changes in spindle oscillation of the two agents also suggest the different thalamocortical sedation mechanisms. Recently, the slow spindles (centered at approximately 12 Hz) and fast spindles (centered at approximately 14 Hz) were found to differ in their frequency, topographic distribution and slow oscillation cycle points [38, 47–49]. The slow fast spindles are generated by different mechanisms and may reflect different physiological processes [38, 48]. One possible explanation may be based on the anatomical heterogeneity because spindles from the core and matrix thalamus system supposedly differ in their cortical projections [50]. The loss of thalamocortical functional connectivity presumably induces both propofol and dexmedetomidine-induced unconsciousness, and propofol confers differential changes in functional connectivity of the different parts of the thalamocortical systems [29, 33]. However, in our study, we could not deduce whether the differential effect of propofol and dexmedetomidine on spindle oscillations was attributed to their distinct effects on the fast/slow spindles or on the different anatomical locations of the thalamocortical systems. Further study combining electrophysiology and functional neuroimaging might explore their different thalamocortical mechanisms. The different spatiotemporal dynamics of alpha and spindle oscillations also suggest that propofol and dexmedetomidine act differently in the thalamocortical system.

The beta/gamma rhythms have been associated with cortical activities and higher levels of cognitive activities, such as sensory gating, attention, perception and motor control. In our study, propofol and dexmedetomidine had different effects on beta/gamma rhythms during sedation, and dexmedetomidine persistently decreased global beta/gamma oscillations during sedation and recovery. The decreased beta/gamma oscillations accounted for the lower BIS value of dexmedetomidine; however, elucidating these central mechanisms was beyond the scope of our study.

Clinical implications

The application of EEG-based monitoring to measure the depth of sedation is controversial. The BIS monitor displays a real-time EEG trace acquired from a frontotemporal montage. Good correlation exists between the BIS value and the dexmedetomidine/propofol-induced
However, the BIS value does not change with increased sedation depth with nitrous oxide or ketamine\[52–55\]. Moreover, when ketamine is used as an adjuvant of other anesthetics, such as propofol or sevoflurane, it reportedly has no effect or even increases the BIS value\[54, 56–58\]. The results of our study indicate that the different thalamocortical mechanisms might play an important role in propofol and dexmedetomidine-induced different EEG frontal spectral power, which results in their distinct BIS values at the same sedation level. We suggest that when BIS is used to monitor the depth of sedation or anesthesia, more consideration should be paid to the types of sedatives/anesthetics and their specific targets and neural circuits in the central nervous system. Furthermore, using the BIS value to compare the depths of sedation/anesthesia for different drugs might not be appropriate.

**Limitations**

In this study, we performed spectral power analysis and created topographic maps to demonstrate gradual brain dynamic changes from wakefulness, moderate sedation, and deep sedation to recovery. We did not apply global or local coherence analysis; thus, the degree of correlation between two cortical regions could not be reflected. Additionally, due to the limitation of EEG recording, we could not directly determine the different effects of these two agents on the thalamus or thalamocortical system, and further study with the combination of functional neuroimaging might refine our results. Furthermore, although the thalamo-cortical network should be involved in propofol/dexmedetomidine-induced sedation, the cellular and neurochemical basis has been studied less. Determining whether reducing the hyperpolarization-activated current $I_h$ is involved in the mechanisms underlying dexmedetomidine-induced sedation is a current research focus. Applying the different neuromodulators in microdialysis and the local field potential recording of animals might help further explore the sedation mechanisms at cellular and molecular levels.

**Conclusion**

In this study, we demonstrated the existence of a distinct hierarchy of EEG changes with the increased sedation levels induced by propofol and dexmedetomidine. Although both agents induced similar increases in delta oscillations, multiple differences in alpha/spindle and beta/gamma oscillations at comparable sedation levels were observed. Based on our results and discussion, these differences might account for the difference in BIS values at the same sedation level and reflect the different sedation mechanisms. EEG-based clinical sedation monitoring should consider the effect of drug type on EEG dynamics.

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