GABAergic ventrolateral pre-optic nucleus neurons are involved in the mediation of the anesthetic hypnosis induced by propofol

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Abstract. Intravenous anesthetics have been used clinically to induce unconsciousness for seventeen decades, however the mechanism of anesthetic-induced unconsciousness remains to be fully elucidated. It has previously been demonstrated that anesthetics exert sedative effects by acting on endogenous sleep-arousal circuits. However, few studies focus on the ventrolateral pre-optic (VLPO) to locus coeruleus (LC) sleep-arousal pathway. The present study aimed to investigate whether VLPO activation contributes to anesthetic-induced unconsciousness by propofol. The present study additionally investigated if the inhibitory effect of propofol on LC neurons was mediated by activating VLPO neurons. Microinjection, target lesion and extracellular single-unit recordings were used to study the role of the VLPO-LC pathway in propofol anesthesia. The results demonstrated that GABA\(_A\) agonist (THIP) or GABA\(_A\) antagonist (gabazine) microinjections into VLPO altered the time of loss of righting reflex and the time of recovery of righting reflex. Furthermore, propofol suppressed the spontaneous firing activity of LC noradrenergic neurons. There was no significant difference observed in firing activity between VLPO sham lesion and VLPO lesion rats. The findings indicate that VLPO neurons are important in propofol-induced unconsciousness, however are unlikely to contribute to the inhibitory effect of propofol on LC spontaneous firing activity.

Introduction

Approximately ten million patients receive general anesthesia for surgery in China every year. While intravenous anesthetics cause unconsciousness, the mechanism and neural basis of unconsciousness are poorly understood (1). At the molecular level, there are dozens of molecules known to be general anesthetic targets, including a number of ion channels (2), gap-junction channels (3), and G protein-coupled receptors (4). It’s remarkable that there is no single molecular target shared by all general anesthetics (1). Therefore, effects of general anesthetics must be comprehended in the context of network connectivity.

There are similarities between general anesthesia and natural sleep. Imaging studies have shown some parallels between the anesthetized brain and the brain during deep non-rapid-eye-movement (NREM) sleep (5,6). Electroencephalogram (EEG) studies have suggested that loss of consciousness caused by general anesthetics resembles the rapid transition from normal wakefulness to sleep (7). Sleep-related EEG waves that resembled gamma, delta and spindle waves have been observed during general anesthesia (8). These findings led an increasingly popular theory that anesthetics may induce unconsciousness by acting on endogenous sleep-arousal neural circuitry. But it remains unclear to what extent sleep-arousal pathway, such as the ventrolateral preoptic nucleus (VLPO)- locus coeruleus (LC) are involved in generating the hypnotic state.

Commonly used general anesthetic propofol exerts sedative effects by targeting GABA\(_A\) receptors. And GABA is the primary inhibitory neurotransmitter released by sleep-promoting neurons in the VLPO, which plays a critical role in inducing and maintaining sleep (9). The VLPO sends GABAergic inhibitory projections to several wake-promoting nuclei throughout the neuroaxis (10), including the LC, tuberomammillary nucleus (TMN) and orexinergic neurons in the lateral hypothalamus (11). Previous research has demonstrated that propofol and various barbiturates activate sleep-promoting VLPO neurons through different receptors (12-14). Moreover, the inhalational anesthetic isoflurane directly depolarizes VLPO neurons (15). Nevertheless, lesion of VLPO neurons could be expected to produce resistance to anesthesia because of the accrual of sleep debt (16). Thus, it remains controversial whether VLPO activation contributes to anesthetic-induced unconsciousness.

In the present study, we hypothesized that propofol may act on VLPO neurons to stimulate the release of GABA, thereby inhibiting wakefulness-promoting neurons in the LC.
To validate the hypothesis, we examined the LoRR (loss of righting reflex) and RoRR (recovery of righting reflex) time following GABA<sub>A</sub> receptor agonist or antagonist microinjection. We also recorded the firing activities of LC neurons under different concentrations of propofol. The results demonstrated that spontaneous firing of LC noradrenergic neurons was inhibited by propofol. Furthermore, the results also showed that firing activities were not significantly different between sham-lesion and VLPO lesion rats. Suggesting that VLPO neurons were not likely involved in propofol-mediated inhibition on LC neurons.

**Materials and methods**

**Animals.** Male Sprague-Dawley rats weighing 260-300 g (n=72) were housed in an isolated chamber at 20-22°C under a 12 h light/dark cycle (lights on at 8:30 AM). Food and water were available ad libitum. Twenty-four rats were used for righting reflex behavioral assays, and the remaining rats were used for *in vivo* electrophysiological recordings. Forty-eight neurons were recorded from forty-eight rats. All animals were supplied by the laboratory animal center of the Third Military Medical University (Chongqing, China). Animal experiments were approved by the Zunyi Medical College Animal Care and Use Committee (approval number: 2016126). All efforts were made to minimize the number of rats and their suffering.

**Surgical procedures and cannulas implantation.** For behavioral experiments, rats were anesthetized with sodium pentobarbital [50 mg/kg, intraperitoneally (i.p.)] and atropine sulfate (0.2 mg, i.p.). Anesthetized animals were mounted onto a stereotaxic device (68505; RWD Life Science, China) in a flat-skull position. The core body temperature was maintained at 37-38°C using a heat-controlled pad equipped with a rectal probe (Stoelting Co., Wood Dale, IL, USA). Sterilized guide cannulas were implanted as described previously (17,18). A pair of sterilized guide cannulas made of 23G stainless steel tubes and plugged internally with 30G stylets were stereotaxically implanted 2.0 mm above the VLPO (anteroposterior, -0.35 mm; left-right, ± 1.2 mm; dorsoventral, -7.0 mm from bregma). Stereotaxic positioning was defined according to the brain atlas of Paxinos and Watson (2007). Guide cannulas were fixed to the skull using dental cement.

**Lesion formation by ibotenic acid and saline injection.** To induce lesions in the VLPO, rats were anesthetized and kept in place as mentioned above. The skull of each rat was exposed, and a glass micropipette (10-12-µm tip diameter) was lowered into the VLPO region stereotaxically. Fifteen nanoliters of ibotenic acid solution (10 nmol; Sigma, St. Louis, MO, USA) (n=24) or saline (n=24) were injected into the VLPO bilaterally using a microinjection injector (Nanoliter 2000, World Precision Instruments, Sarasota, FL, USA) as described previously (16). The glass micropipette was slowly withdrawn after 5 min. The two holes above VLPO were filled with bone wax. The wound was stitched with sutures and closed with wound clips. A prophylactic dose of penicillin (50 ku/kg, intramuscular injection) was injected into each rat following surgery. After seven days of recovery, *in vivo* electrophysiology experiments were conducted.

**Righting reflex behavioral assays.** In rodents, loss of consciousness can be measured by the LoRR and resumption of consciousness can be analyzed by the RoRR (19,20). Fig. 1A illustrates the experimental design used to quantify the LoRR and RoRR after the VLPO was microinjected with artificial cerebrospinal fluid (ACSF) (126 mM Na<sup>+</sup>, 3 mM K<sup>+</sup>, 2 mM Ca<sup>2+</sup>, 1.2 mM Mg<sup>2+</sup>, 150 mM Cl<sup>-</sup>, pH=7.4) and a bolus injection of propofol (11 mg/kg, i.v.). After seven days of recovery from cannulas implantation surgery, rats (n=24) were put into a Plexiglas inhalation anesthesia induction chamber for 10-min acclimation (Fig. 1A; t=–40 min). Then, oxygen containing 2% isoflurane was introduced into the chamber at 2 l/min (Fig. 1A; t=–30 min). The 30G stylet from the guide cannula was removed from the isoflurane-anesthetized rats, and a 31G injection cannula was inserted. An injection cannula was attached to a dual-channel microinjection syringe with PE tubing under the control of an injection pump (302; RWD Life Science), and then a 24G intravenous catheter was implanted and fixed to the rat’s tail vein. Isoflurane administration was subsequently discontinued. The ACSF (t=0 min; 0.2 µl/side; n=24) (Fig. 1A) was microinjected into the bilateral VLPO at an infusion speed of 0.2 µl/min for 1 min (t=1 to t=2) (Fig. 1A). Five min later (t=7 min) (Fig. 1A), propofol (11 mg/kg, n=24) was bolus injected into the tail vein. The induction time of propofol was quantified as the time (sec) to LoRR; the resumption time of consciousness can be analyzed by the RoRR (19,20). Righting reflex behavioral assays.

**Extracellular single-unit recordings.** After a recovery period of seven days, the extracellular single-unit recording was performed from noradrenergic neurons of the LC (LC-NA) with VLPO lesions (n=24) and sham lesions (n=24) rats. Each group was subdivided into a consciousness group (n=6, without propofol administration), a low propofol concentration group (n=6; 20 mg/kg/h, i.v.), medium concentration group (n=6; 40 mg/kg/h, i.v.) and a high concentration propofol group (n=6; 60 mg/kg/h, i.v.).

The protocol of extracellular single-unit recordings in non-anesthetized rats was referred to previously report (21). Briefly, the rats were progressively habituated to the head-restrained position (7-14 days) by placing them inside a plastic chamber, painlessly restraining the head with a head holder and preventing large body movements with a cotton-coated plastic cover. The eyes were covered to reduce visual stimulation. After 7-14 days of habituation, the rats could be kept in a sphinx position for 3-6 consecutive h without showing any signs of discomfort. If any signs of discomfort were seen, the rats were freed from the restrained position.

Then A 24G intravenous catheter was implanted and fixed as described above. Rats were removed from the induction chamber and mounted onto a stereotaxic device with blunt ear bars. Propofol was continuously delivered via a syringe pump (WZS-50F6; Smith Medical, Plymouth, MN, US) connected
to the intravenous catheter in propofol administration groups. LC-NA extracellular recordings were obtained as previously described (21). A single barrel glass micropipette was filled with 0.5% sodium acetate and lowered into the LC (3.5 mm caudal to lambda, 0.85-1.0 mm lateral to the midline, 5-6.5-mm below the cortical surface) by using a micromanipulator (Mini 25; Luigs & Neumann, Ratingen, Germany) according to coordinates defined in the brain atlas of Paxinos and Watson (2007). The impedance of the micropipette was 10-20 MΩ, measured in physiological saline. Recording was performed during a period of at least 5 min, with only one cell measured from each rat. The recorded extracellular signals were amplified using a patch clamp amplifier (EPC10; HEKA Elektronik, Lambrecht, Germany), bandpass filtered (100-10,000 Hz) and saved to a computer installed with PATCHMASTER (Heka) and Mini Analysis System (Synaptosoft, Inc., Fort Lee, NJ, USA) for offline analysis. The body temperature of rats was maintained at 37-38˚C during the recording period.

Noxious cutaneous stimulation. The discharge patterns of NA-LC neurons were identified using standard criteria (22,23). They display a positive-negative action potential shape with a notch on the ascending limb (Fig. 2A) and a typical, biphasic excitation-inhibition response to noxious stimulation of the contralateral hind paw (Fig. 2B). We pinched the skin of the rat's hind paw using toothed forceps for cutaneous nociceptive stimulation to induce this characteristic firing response. The noxious stimulation procedure was only operated in anesthetized rats.

Histological localization of microinjection and lesion areas. After behavioral and electrophysiology experiments, rats were deeply anesthetized with sodium pentobarbital. Cold 100 ml physiological saline was perfused through the ascending aorta, followed by 200 ml 4% paraformaldehyde. The brains were then removed, post-fixed for 24 h at 4˚C in paraformaldehyde, and transferred to glass containers filled with 30% sucrose for 48 h at 4˚C. 20-µm coronal sections containing microinjection or lesion sites were cut and stained with hematoxylin-eosin for histological verification (Fig. 3). Stereotaxic coordinates were identified according to the brain atlas of Paxinos and Watson (2007).

Statistical analyses. Statistical analyses were performed using SPSS 17.0 for Windows. All data were expressed as means ± standard deviation. A P-value of <0.05 was considered statistically significant using paired-samples t-test in behavioral experiments. N refers to the number of rats studied. The
firing frequencies of noradrenergic neurons in different groups were compared using independent-samples t-test. The firing frequencies of noradrenergic neurons in the same group were compared using one-way ANOVA, and \( n \) refers to the number of neurons analyzed.

**Results**

**The effects of GABA\(_A\) receptors in the VLPO on LoRR and RoRR.** To determine whether GABAergic neurons are involved in unconsciousness induced by propofol. We microinjected GABA\(_A\) agonist (THIP) or GABA\(_A\) antagonist (gabazine) into VLPO and observed the LoRR and RoRR. Fig. 4A and B illustrates how ACSF and THIP microinjection into the VLPO of the same rats (n=12) affected propofol-mediated anesthesia and recovery in the same rats (n=12). Microinjection of gabazine into the VLPO increased LoRR (20.42±4.14 sec to 26.83±6.28 sec, \( P<0.05 \)) (Fig. 4C) and decreased RoRR (605.67±81.79 sec to 562.58±78.05 sec, \( P<0.05 \), Fig. 4D) before propofol administration.

**The effect of propofol on the spontaneous firing activity of LC neurons in rats with VLPO lesion.** To explore the effect of propofol on LC firing activity, we anesthetized the VLPO sham-lesion and VLPO lesion rats with different concentrations of propofol. Fig. 5 shows that propofol inhibited the spontaneous firing activity of LC neurons in a dose-dependent manner. In the VLPO sham-lesion group, the spontaneous firing activity of LC neurons in medium propofol concentration (40 mg/kg/h) was reduced to 2.98±0.6 Hz compared with
6.54±1.16 Hz of the low propofol concentration (20 mg/kg/h) group (P<0.05). The spontaneous firing activity of LC neurons in high propofol concentration (60 mg/kg/h) was reduced to 1.36±0.9 Hz compared with 2.98±0.6 Hz of the medium propofol concentration (40 mg/kg/h) group (P<0.05). The firing activity of LC neurons in medium propofol concentration was reduced to 2.98±0.6 Hz compared with 7.32±1.33 Hz of the consciousness group (P<0.05). And the firing activity of LC neurons in high propofol concentration was reduced to 1.36±0.9 Hz compared with 7.32±1.33 Hz of the consciousness group (P<0.05). However, there's no statistical significant differences in the firing activity between consciousness and low propofol concentration group, (Fig. 5A).

The similar inhibitory effect was observed in the VLPO lesion group. In this group, the spontaneous firing activity of LC neurons in medium propofol concentration rats was 2.80±0.46 Hz compared with 5.89±0.93 Hz in the low propofol concentration group (P<0.05). The spontaneous firing activity of LC neurons in high propofol concentration rats was 1.57±0.85 Hz compared with 2.80±0.46 Hz in the medium propofol concentration group (P<0.05). The firing activity of LC neurons in medium propofol concentration was reduced to 2.80±0.46 Hz compared with 6.99±2.03 Hz of the consciousness group (P<0.05). And the firing activity of LC neurons in high propofol concentration was reduced to 1.57±0.85 Hz compared with 6.99±2.03 Hz of the consciousness group (P<0.05). In addition, there's no statistical significant differences in the firing activity between conscious and low propofol concentration group, (Fig. 5B). Nevertheless, our experimental results showed that firing activities were not significantly different between sham-lesion and VLPO lesion rats after administration of the same concentration of propofol (P>0.05), (Fig. 5C).

Discussion

When we microinjected the GABA_A agonist THIP into the VLPO before propofol administration, results showed that LoRR decreased while RoRR increased. In contrast, microinjection of the GABA_A antagonist gabazine into the VLPO increased LoRR and decreased RoRR. The results indicate that GABA_A receptors in the VLPO are involved in propofol-induced unconsciousness, which is consistent with previous reports demonstrating that potentiating GABAergic transmission in the VLPO promotes sleep (15,24,25). The VLPO was the first nucleus to be identified as a sleep-regulating center containing GABAergic neurons (26). Two-thirds of VLPO neurons are inhibited by the wake-promoting neurotransmitter noradrenaline (NA). These
are putative sleep-promoting neurons and can be activated by general anesthetics. The remaining one-third of neurons can be excited by NA and inhibited by general anesthetics (14). Patch clamp experiments (27) have shown that propofol directly activates NA(-) neurons via GABA\(_A\) receptors. THIP is a specific GABA\(_A\) receptor agonist (28) and microinjection of THIP into the pontine reticular formation promotes wakefulness (29). On the other hand, microinjection of THIP into the VLPO improves sleep (30).

Our work suggests that early activation of VLPO GABAergic neurons enhances the anesthetic effect of propofol. This effect may be the result of more GABA releasing into the arousal system in response to THIP. Nelson (12) demonstrated that the sedative effects of GABA-responsive agents were attenuated when gabazine was microinjected into the TMN nucleus. In our study, the gabazine microinjection significantly increased LoRR and decreased RoRR. After the GABA\(_A\) receptor was blocked, propofol-induced anesthesia was attenuated. In conclusion, potentiating GABAergic transmission in the VLPO may represent a mechanism for propofol-induced anesthesia. In contrast, blocking GABAergic transmission may attenuate unconsciousness following propofol administration.

In the present study, we also found that the firing rate of LC neurons was dramatically decreased when the concentration of propofol was higher. It was reported that the persistent inward calcium current and the cAMP-activated inward sodium current formed the spontaneous firing activity of LC neurons (31). Previously \textit{in vitro} patch clamp experiments demonstrated that propofol inhibited the LC neurons by activation of outward chloride current, and the GABA\(_A\) receptor antagonists could block the inhibitory effect (32). Accordingly, in our \textit{in vivo} extracellular single-unit recordings experiments, we hypothesize that outward chloride current activated by propofol can balance the inward calcium current and the cAMP-activated sodium current. Thus, the spontaneous firing activity of LC will be inhibited. Furthermore, higher propofol concentration could increase the outward chloride current as to hyperpolarize the LC neurons.

The LC neurons fire continually during consciousness and the firing rate decreases during non-rapid eye movement (NREM) and rapid eye movement (REM) sleep (33). Halothane is an agent that could suppress such activity of LC neurons (34). Moreover, \(\alpha_2\)-adrenergic receptors in the LC are a major target for the sedative agent dexmedetomidine (35).
which directly inhibits LC neurons. Our data showed that the firing rate of LC neurons was not influenced by VLPO lesions, indicating that VLPO GABAergic neurons were not involved in the propofol-mediated inhibition on LC neurons. VLPO lesions could induce acute sleep loss in rats (16,36) and confer a short-term resistance to isoflurane (15). Our previous work (37) conducted acute lesions in VLPO culminated in opposite results, which indicated that VLPO is necessary for the propofol-induced inhibition of LC activity. Another thing to note is that sleep debt caused by sleep loss can amplify the sensitivity of propofol and isoflurane (38). In order to minimize the effect of sleep debt, we performed experiments seven days after the lesions. VLPO lesions did not affect the firing rate of LC neurons after propofol was administered. This could also be explained by the presence of different sleep-promoting neuron populations in the median preoptic nucleus (MNPO) (39), basal forebrain (BF) (40) and cortex (41). If one region is experimentally lesioned, other sleep-promoting systems remain intact. Thus, LC neurons also received inhibitory afferent projections from these nuclei, which may have compensated for the loss of VLPO GABAergic neurons in this study.

However, this study is limited in the following aspects. We aim to investigate the role NA(-) neurons play in propofol-induced anesthesia. Nevertheless, NA(+) neurons in the VLPO were also lesioned by ibotenic acid. Specific inactivation of NA(-) neurons by pharmacogenetics or genetic manipulation would be a better approach to conducting our study. Additionally, quantifying the changes of neurotransmitters GABA in the LC area would make our conclusion more persuasive.

In conclusion, we identified that VLPO GABAergic neurons play a crucial role in propofol-induced unconsciousness. Propofol could suppress the spontaneous firing activity of LC noradrenergic neurons in vivo, but VLPO neurons are not involved in propofol-mediated inhibition on LC neurons. Present findings are consistent with the hypothesis that general anesthetics act on endogenous sleep-wake circuitry to exert sedative effects.

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