Sleep spindles are generated in the absence of T-type calcium channel-mediated low-threshold burst firing of thalamocortical neurons

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T-type Ca\textsuperscript{2+} channels in thalamocortical (TC) neurons have long been considered to play a critical role in the genesis of sleep spindles, one of several TC oscillations. A classical model for TC oscillations states that reciprocal interaction between synaptically connected GABAergic thalamic reticular nucleus (TRN) neurons and glutamatergic TC neurons generates oscillations through T-type channel-mediated low-threshold burst firings of neurons in the two nuclei. These oscillations are then transmitted from TC neurons to cortical neurons, contributing to the network of TC oscillations. Unexpectedly, however, we found that both WT and KO mice for Ca\textsubscript{2+}\textsuperscript{3.1}, the gene for T-type Ca\textsuperscript{2+} channels in TC neurons, exhibit typical waxing-and-waning sleep spindle waves at a similar occurrence and with similar amplitudes and episode durations during non-rapid eye movement sleep. Single-unit recording in parallel with electroencephalography in vivo confirmed a complete lack of burst firing in the mutant TC neurons. Of particular interest, the tonic spike frequency in TC neurons was significantly increased during spindle periods compared with nonspindle periods in both genotypes. In contrast, no significant change in burst firing frequency between spindle and nonspindle periods was noted in the WT mice. Furthermore, spindle-like oscillations were readily generated within intrathalamic circuits composed solely of TRN and TC neurons in vitro in both the KO mutant and WT mice. Our findings call into question the essential role of low-threshold burst firings in TC neurons and suggest that tonic firing is important for the generation and propagation of spindle oscillations in the TC circuit.

Sleep spindles are one type of several rhythmic brain waves detected by electroencephalography (EEG) during normal non-rapid eye movement (NREM) sleep. A spindle consists of characteristic waxing-and-waning field potentials grouped into 7- to 14-Hz oscillations that last for 1–3 s and recur once every 5–10 s in the thalamus and the cortex (1–3). Spindles are also visible under anesthesia, particularly with barbiturates but also with ketamine-xylazine combinations (4, 5). These oscillations are generated in the thalamus as a result of synaptic interactions between inhibitory [i.e., thalamic reticular nucleus (TRN)] neurons and excitatory thalamocortical (TC) neurons, and are propagated to the cortex. Corticothalamic projections back to the thalamus complete the cortico-thalamo-cortical loop.

In vivo data suggest that TRN neurons are spindle pacemakers, because spindles can be generated in deafferented TRN neurons (6) but disappear in TC regions after disconnection from TRN neurons (7). However, in vitro data suggest that an intact TC-TRN network is a necessity, because spindles are abolished after disconnection of TC and TRN neurons (8).

Two distinct firing patterns, tonic and burst, are displayed by both TRN and TC neurons. Burst firing is mediated by low-threshold T-type Ca\textsuperscript{2+} channels (9). Of the three subtypes of T-type channels, Ca\textsubscript{2+}\textsuperscript{3.1} is expressed exclusively in TC regions, whereas Ca\textsubscript{2+}\textsuperscript{3.2 and 3.3} are abundant in TRN regions (10). T-type channels in TC neurons have been proposed to be a critical component in the generation of physiological and pathological TC oscillations, such as sleep rhythms (1, 11, 12) and the spike-wave discharges (SWDs) of absence seizures (11, 13, 14). One generally accepted hypothesis proposes that inhibitory inputs from TRN neurons de-inactivate T-type Ca\textsuperscript{2+} channels in TC neurons, leading to induction of burst firings in TC neurons, which in turn excite reciprocally connected TRN neurons. These thalamic oscillations are then transmitted from TC neurons to cortical neurons. This model proposes that T-type channel-mediated burst firing in TC neurons underlies sleep spindles and other sleep rhythms within TC circuits (1, 11).

There have long been doubts regarding the extent to which TC T-type Ca\textsuperscript{2+} channels contribute to the heterogeneity of TC oscillations during NREM sleep, which consists of multiple EEG components including slow waves (<1 Hz), delta waves (1–4 Hz), and sleep spindles (7–14 Hz). T-type channels have received particular attention in the genesis of spindles and delta waves, both of which are thought to originate from thalamic neurons (8), although cortically generated delta waves also have been found in cats with thalamic lesions (15). A role for TC T-type channels in sleep has been demonstrated in two studies, one using mice with a global deletion (16) and the other using mice with a thalamus-restricted deletion (17) of Ca\textsubscript{2+}\textsuperscript{3.1} T-type channels. Both mice exhibited reduced delta waves with intact slow waves (16, 17). Fragmented sleep was observed in both mice, indicating that this sleep phenotype in the global Ca\textsubscript{2+}\textsuperscript{3.1}−/− mice

Significance

This study addresses one of the most fundamental issues in sleep rhythm generation. The theory that low-threshold burst firing mediated by T-type calcium channels in thalamocortical neurons is the key component for sleep spindles, has been accepted as dogma and appears throughout the literature. In this study, however, in vivo and in vitro evidence shows that sleep spindles are generated normally in the absence of T-type channels and burst firing in thalamocortical neurons. Furthermore, our data indicate a potentially important role of tonic firing in this rhythm generation. This study advances the knowledge of sleep and vigilance control to another level of understanding.
Results

Sleep Spindles Are Not Altered in Ca$_{3.1}^{−/−}$ Mice During Natural NREM Sleep. First, in an attempt to elucidate the role of low-threshold Ca$^2+$ channels in sleep spindles, we examined the characteristics of spindle oscillations induced by barbiturate injection. In agreement with the literature (20), we found that barbiturate-induced spindles were characterized by large-amplitude oscillations of increased duration compared with sleep spindles (Fig. 2A). The mean duration of barbiturate-induced spindle episodes was almost doubled compared with the spindle episode duration during NREM sleep in the Ca$_{3.1}^{+/+}$ mice, but not in the Ca$_{3.1}^{−/−}$ mice. Compared with the Ca$_{3.1}^{+/+}$ mice, the Ca$_{3.1}^{−/−}$ mice exhibited shorter barbiturate-induced spindle episodes ($^*P < 0.05$; Fig. 2B), as well as smaller peak-to-peak amplitudes ($^*P < 0.05$), whereas there was no difference in the peak frequency of spindles between the two genotypes (Fig. 2B). Of note, the spindle episodes in the Ca$_{3.1}^{−/−}$ mice were composed of shorter episodes compared with those seen in the WT mice (Fig. 2C).

Taken together, these results suggest that low-threshold spikes (LTSs) mediated by Ca$_{3.1}$ T-type Ca$^2+$ channels in TC neurons are not essential for spindle generation, but may play a role in increasing the duration of spindles episodes induced by barbiturates.

Spike Discharges in TC Neurons During in Vivo Spindle Oscillations. Our findings regarding natural sleep spindles and barbiturate-induced sleep spindles appear to indicate that sleep spindles are not altered in Ca$_{3.1}^{−/−}$ mice.

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induced spindles in Ca$_{v}3.1^{-/-}$ mice prompted us to investigate the properties of TC neuron spike discharges during spindle episodes in the absence of low-threshold burst firing. To do this, we performed in vivo extracellular single-unit recordings in TC neurons while simultaneously monitoring the cortical EEG under barbiturate anesthesia.

Both tonic and burst spikes were observed in Ca$_{v}3.1^{+/+}$ TC cells, whereas only tonic spikes were detected in Ca$_{v}3.1^{-/-}$ TC cells (Fig. 3A–C), confirming previous in vitro data (13). Bursts were identified according to the conventional criteria (21), with a burst defined as a cluster of two or more spikes with interspike intervals of ≤4 ms, in which the first spike in the burst has a preceding interspike interval of >100 ms (Fig. S1). Spikes recorded over 15 consecutive spindle sequences (SP1–SP15) were aligned with the initiation time of each cortical EEG spindle event. The perievent histograms showed a large number of TC spikes co-occurring within cortical spindle events (Fig. 3B), as reported previously (22, 23). Of note, we found the spike frequency was significantly higher during spindle periods compared with nonspindle periods in both WT mice (1.71 ± 0.23 Hz vs. 0.73 ± 0.16 Hz) and KO mice (1.94 ± 0.43 Hz vs. 0.18 ± 0.04 Hz) (Fig. 3C, a). Of particular importance, there was no difference in the total spiking rate (across tonic and burst activity) within spindle events between two genotype TC neurons (P = 0.42, Fig. 3C, a) with event correlation (Fig. S2), whereas the nonspike spike frequency was significantly lower in Ca$_{v}3.1^{-/-}$ neurons compared with Ca$_{v}3.1^{+/+}$ neurons (Fig. 3C, a).

Further analysis of the firing mode of TC neurons revealed a significantly higher tonic spike frequency in both WT and mutant mice during spindle periods than during nonspindle periods (Fig. 3C, b). During spindle episodes, the tonic spike frequency was significantly higher in mutant KO mice compared with WT mice, perhaps explaining why the total spiking rate of mutant TC neurons within cortical spindle events was as high as that in WT neurons, even though the mutants are devoid of burst spikes. In contrast, there was no significant difference in burst firing between spindle and nonspindle periods (Fig. 3C, b), either in the frequency of burst spikes (0.58 ± 0.15 Hz vs. 0.35 ± 0.09 Hz) or in the occurrence of burst events (0.28 ± 0.09 Hz vs. 0.16 ± 0.06 Hz; P = 0.27). In other words, the majority of Ca$_{v}3.1^{+/+}$ TC cells discharged burst spikes without preference for spindle episodes; just 2 of 13 fired bursts preferentially during the spindle period.

Also of interest is our finding of no difference in the overall spike frequency of TC neurons during spindle events between the two genotypes. Of note, WT TC neurons exhibited a higher tonic spike frequency than burst spike frequency during spindle episodes. More importantly, the tonic spike frequency in WT mice was significantly (~3.5-fold) higher during spindle periods than during nonspindle periods, whereas the burst spike frequency did not differ in the two periods. The tonic spike frequency observed in Ca$_{v}3.1^{+/+}$ TC cells coincided even more strongly with spindles (~10-fold higher). In addition, spike discharges recorded in both Ca$_{v}3.1^{+/+}$ and Ca$_{v}3.1^{-/-}$ TRN neurons were more frequent during spindle episodes (Fig. S3), in parallel with the findings in TC neurons.

Spindle-Like Oscillations Elicited in Intrathalamic Circuits in Ca$_{v}3.1^{-/-}$ Mice. Our finding that robust generation of cortical spindles is accompanied by increased TC spike discharges in Ca$_{v}3.1^{-/-}$ mice prompted us to further investigate whether spindle oscillations can be generated within an intrathalamic circuit composed only of TRN and TC neurons in the presence or absence of LTSs in TC neurons. For this purpose, we assessed thalamic network...
activity in TC slices with and without attached cortex (24). Previous studies have demonstrated that oscillations within the range observed for sleep spindles in vivo (22) can be evoked in thalamic slices by electrical stimulation of the internal capsule (IC) (25). These are thus referred to as “spindle-like oscillations” (SLOs) (26). We adopted this paradigm using the multielectrode array system as described in SI Materials and Methods.

Thalamic oscillations were evoked by a single 20- to 100-μV, 60- to 80-μs electrical stimulation in the IC in TC slices. Clusters of spikes composed of multiple units with various amplitudes were detected simultaneously in TRN and TC nuclei (Fig. 4A). SLOs were readily generated in these thalamic slices from both Ca\textsubscript{v}3.1\textsuperscript{I-/+} and Ca\textsubscript{v}3.1\textsuperscript{I-/-} mice in the presence and absence of cortex (Fig. 4A). There were no significant differences in the durations of evoked oscillatory activities between Ca\textsubscript{v}3.1\textsuperscript{I-/+} and Ca\textsubscript{v}3.1\textsuperscript{I-/-} thalamic slices with cortex (7 slices from 7 WT mice vs. 8 slices from 7 mutant mice) or without cortex (13 slices from 8 WT mice vs. 12 slice from 9 mutant mice) (Fig. 4B), although the total TC spike numbers in Ca\textsubscript{v}3.1\textsuperscript{I-/-} thalamic slices were lower than those in Ca\textsubscript{v}3.1\textsuperscript{I-/+} slices in the absence of cortex (P = 0.047; Fig. 4C).

The foregoing results provide direct evidence contrary to the classical view that postinhibitory low-threshold burst firing mediated by T-type Ca\textsuperscript{2+} channels in TC neurons is essential for the generation of oscillations within the TRN-TC circuit. They suggest that instead, sleep spindles observed in Ca\textsubscript{v}3.1\textsuperscript{I-/-} mice may be generated within the TRN-TC circuit in the absence of TC bursts.

Excitability of Ca\textsubscript{v}3.1\textsuperscript{I-/-} TC Neurons In Vitro. Our next step was to examine whether generation of spindle oscillations within the Ca\textsubscript{v}3.1\textsuperscript{I-/-} intrathalamic circuit might be related to changes in the intrinsic excitability of TC neurons rather than to a loss of low-threshold burst firing by deletion of Ca\textsubscript{v}3.1 T-type channels. To do this, we used the whole-cell patch-clamp technique to measure the excitability of TC neurons. We found no differences between Ca\textsubscript{v}3.1\textsuperscript{I-/+} (n = 12) and Ca\textsubscript{v}3.1\textsuperscript{I-/-} (n = 14) TC neurons in membrane capacitance (141.2 ± 15.6 pF vs. 137.4 ± 10.5 pF), resting membrane potential (−59.11 ± 5.34 mV vs. −57.54 ± 6.31 mV), or membrane resistance (219.5 ± 48.2 Ω vs. 175.4 ± 38.2 Ω).

We then measured hyperpolarization-activated cyclic nucleotide-gated channel-mediated current (I\textsubscript{h}) in TC neurons at the tail current at 0.7 s after a test voltage pulse (Fig. 5A) to minimize the capacitative contribution (27) or non-Ih-active current (28). We observed an increase in the current density of I\textsubscript{h} in Ca\textsubscript{v}3.1\textsuperscript{I-/-} TC neurons compared with Ca\textsubscript{v}3.1\textsuperscript{I-/+} TC neurons only when membrane potential was hyperpolarized to ≥−100 mV (Fig. 5A and B). We then measured the firing pattern of TC neurons. Low-threshold burst firing was abolished in Ca\textsubscript{v}3.1\textsuperscript{I-/-} TC neurons, whereas vigorous burst firing was induced in Ca\textsubscript{v}3.1\textsuperscript{I-/+} TC neurons, with a membrane potential rebound to −60 mV after hyperpolarization to −70 to −120 mV (Fig. 5C and D). Some Ca\textsubscript{v}3.1\textsuperscript{I-/-} TC neurons (6 of 10) produced a single spike when the membrane potential was returned to −60 mV after hyperpolarization to ≥−100 mV (Fig. 5 C and D), possibly related to the enhanced I\textsubscript{h} current (Fig. 5 A and B). However, the enhanced I\textsubscript{h} current did not affect either the tonic or burst firing rate elicited by depolarizing current injections (Fig. 5 E–E), and did not alter any other intrinsic membrane properties of Ca\textsubscript{v}3.1\textsuperscript{I-/-} TC neurons. Thus, we conclude that deletion of Ca\textsubscript{v}3.1 T-type channel did not significantly alter the intrinsic excitability of TC neurons. The firing pattern of TRN neurons was not changed either (Fig. S4).

Discussion

We have demonstrated here that sleep spindles and intrathalamic SLOs are generated in the absence of Ca\textsubscript{v}3.1 T-type Ca\textsuperscript{2+} channels in TC neurons. This finding calls into question the validity of the long-standing hypothesis that LTSSs mediated by T-type channels in TC neurons are essential for spindle oscillations.

Ca\textsubscript{v}3.1 T-Type Channels in Sleep Rhythms. Sleep spindles have been proposed to arise from the intrathalamic circuit composed of TRN and TC neurons (8, 29). Postinhibitory LTSSs in TC neurons has been considered essential for the generation of intrathalamic oscillations and their propagation to the cortex (1). According to this hypothesis, spindle oscillations should be diminished, if not abolished, in cortical EEGs of Ca\textsubscript{v}3.1\textsuperscript{I-/-} mice lacking LTSSs in TC neurons.

In contrast to this expectation, we clearly observed sleep spindle waves in cortical EEG recordings from both Ca\textsubscript{v}3.1\textsuperscript{I-/+} and Ca\textsubscript{v}3.1\textsuperscript{I-/-} mice during NREM sleep. We chose a method suitable for analyzing spindle waves in mice (19) used in previous studies (1–3). In a previous study on the sleep phenotype of Ca\textsubscript{v}3.1\textsuperscript{I-/-} mice, we filtered EEG data with a high-pass cutoff at 0.1 Hz to include and analyze all sleep rhythms. This did not exclude the possibility that a significant reduction in delta power in Ca\textsubscript{v}3.1\textsuperscript{I-/-} mice might exaggerate the difference in the EEG power density of the spindle component (16). In the present study, we used a 6- to 15-Hz bandpass filter to clearly reveal spindle oscillations and exclude delta and slow waves (Figs. 1A and 2A and Fig. S5), and found no difference in the number of sleep spindle episodes between Ca\textsubscript{v}3.1\textsuperscript{I-/+} and Ca\textsubscript{v}3.1\textsuperscript{I-/-} mice. Neither the duration nor the amplitude of the spindle episodes was reduced in Ca\textsubscript{v}3.1\textsuperscript{I-/-} mice during natural NREM sleep, raising questions about the essential role of TC neuron T-type
channels in the generation of sleep spindles. However, the mean episode duration and amplitude of barbiturate-induced spindles were substantially reduced in Ca$_{v}3.1^{-/-}$ mice, in accordance with our previous findings of a reduced EEG power density at delta and spindle frequencies in Ca$_{v}3.1^{-/-}$ mice under high-dose urethane (2 mg/kg), another spindle-inducing anesthesia (16).

These results suggest that T-type Ca$^{2+}$ channels and the resulting LTSs in TC neurons are not required for the generation of spindle oscillations in TC circuits, but may play a role in enhancing the strength of spindles, as observed in barbiturate-induced spindles. This view is quite different from the classical theory that the generation of sleep spindles requires low-threshold burst firing of TC neurons.

A recent study in mice lacking the Ca$_{v}3.3$ T-type channels abundantly expressed in TRN found a selective reduction in spindle power before the onset of REM sleep (30). The same study also found that delta power was unaffected by the absence of Ca$_{v}3.3$ channels (30), whereas Ca$_{v}3.1^{-/-}$ mice demonstrated a significantly reduced power density of the delta range (1–4 Hz) during NREM sleep (16). These findings suggest that Ca$_{v}3.3$ T-type Ca$^{2+}$ channels in TRN neurons contribute to spindles, whereas channels encoded by Ca$_{v}3.1$ genes in TC neurons contribute to the delta waves during NREM sleep.

Firing Patterns During Spindle Oscillations in TC Neurons. We examined an intriguing question regarding the firing pattern of TC neurons that lack T-type channels in response to inhibitory inputs from the TRN neurons during spindle episodes. Only tonic spikes were observed in TC nuclei in Ca$_{v}3.1^{-/-}$ mice, as expected (Fig. 3). Tonic spikes were more abundant than burst spikes during spindles in Ca$_{v}3.1^{-/-}$ TC neurons, whereas no difference in tonic and burst spike frequency was seen during nonspindle periods.

With regard to the critical role of tonic firing in spindles, our most compelling observation may be that in WT TC neurons, the tonic spike frequency increases significantly during cortical spindle events compared with nonspindle periods. This pattern is clearly different from that seen for burst spike frequency in WT TC neurons, which is almost equal during spindle and nonspindle periods. Of interest, the overall spike frequency during spindle episodes in mutant TC neurons with no burst spikes did not differ from that in WT TC neurons, because the degree of increase in tonic spike frequency during spindle periods was much larger in mutant TC neurons. Nevertheless, the duration of barbiturate-induced spindles was shorter in the mutant mice compared with the WT mice (Fig. 2) even though the total spike frequency was similar in the two mice during barbiturate-induced spindles (Fig. 3), implying that TC burst firing may play a role in increasing the duration of barbiturate-induced spindle oscillations. The total spike frequency during nonspindle periods was lower in mutant mice than in the WT mice, a finding that remains to be explained.

The explicit induction of oscillations within an isolated Ca$_{v}3.1^{-/-}$ intrathalamic circuit convinced us that spindle oscillations may be generated within the TRN-TC network as suggested previously (31), but in the absence of LTSs in TC neurons. We also confirmed that deletion of Ca$_{v}3.1$ T-type channels had no effect on the intrinsic firing properties of TC neurons (Fig. 5). Taken together, these findings suggest that spindle oscillations can be generated and propagated via any type of TC spike activity as long as TC neurons discharge spikes at an appropriate frequency. This conclusion differs from previous observations on the critical role of TC T-type channels in sleep delta waves (16) and paroxysmal SWDs (13, 32, 33), and leads us to suggest that the various TC oscillations may be generated through diverse mechanisms.
CaV3.1 T-Type Channels in Spindles vs. SWDs. It has been proposed that TC oscillations occurring during sleep spindles and SWDs of absence seizures share common elements and mechanisms (11). Clinical observations of the temporal coincidence between sleep spindles and SWDs and experimental findings showing that penicillin, a weak GABA_B receptor antagonist, induces the transformation of sleep spindles into SWDs gave rise to the view that SWDs may be a perverted form of sleep spindles (34).

Previous studies have found that CaV3.1−/− mice are resistant to GABA_A receptor agonist-induced SWDs (13), and that deletion of CaV3.1 completely suppresses or markedly reduces SWDs in genetic mouse models of absence seizures (35), implying that CaV3.1 T-type Ca_2+ channels play a crucial role in the genesis of SWDs. In contrast, the increased T-type Ca_2+ currents in TC neurons occur alongside the generation of SWDs and absence seizure phenotypes in transgenic or mutant mice (32, 33). Our present findings support the hypothesis that CaV3.1 T-type Ca_2+ channels are not an essential component in the generation of sleep spindles. Overall, these results suggest that paroxysmal TC oscillations during SWDs require the post-inhibitory low-threshold burst firing mediated by T-type channels in TC neurons, whereas physiological TC oscillations during sleep spindles do not.

In conclusion, our findings suggest that T-type Ca_2+ channels in TC neurons might not be essential for the generation and propagation of spindle oscillations. This study provides valuable insights into the realm of T-type channels as a possible target for controlling the TC oscillations generated under pathological conditions may be controlled by modulating T-type channels in TC neurons without affecting natural physiological sleep spindles. Our findings raise further questions, including what drives TC neurons to fire during spindles and what determines the spindle frequency. Future studies should be aimed at investigating the detailed mechanism through which each type of TC oscillation is generated.

Materials and Methods

Treatment of animals and all experiments were conducted in accordance with the ethical guidelines of the Institutional Animal Care and Use Committee of Yonsei University, Institute of Basic Science and the Korea Institute of Science and Technology.

The mice and experiments for this study are described in detail in SI Materials and Methods.

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