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Citation for final published version:


Publishers page: http://dx.doi.org/10.1523/JNEUROSCI.3169-13.2013

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Essential Thalamic Contribution to Slow Waves of Natural Sleep

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Slow waves represent one of the prominent EEG signatures of non-rapid eye movement (non-REM) sleep and are thought to play an important role in the cellular and network plasticity that occurs during this behavioral state. These slow waves of natural sleep are currently considered to be exclusively generated by intrinsic and synaptic mechanisms within neocortical territories, although a role for the thalamus in this key physiological rhythm has been suggested but never demonstrated. Combining neuronal ensemble recordings, microdialysis, and optogenetics, here we show that the block of the thalamic output to the neocortex markedly (up to 50%) decreases the frequency of slow waves recorded during non-REM sleep in freely moving, naturally sleeping-waking rats. A smaller volume of thalamic inactivation than during sleep is required for observing similar effects on EEG slow waves recorded during anesthesia, a condition in which both bursts and single action potentials of thalamocortical neurons are almost exclusively dependent on T-type calcium channels. Thalamic inactivation more strongly reduces spindles than slow waves during both anesthesia and natural sleep. Moreover, selective excitation of thalamocortical neurons strongly entrains EEG slow waves in a narrow frequency band (0.75–1.5 Hz) only when thalamic T-type calcium channels are functionally active. These results demonstrate that the thalamus finely tunes the frequency of slow waves during non-REM sleep and anesthesia, and thus provide the first conclusive evidence that a dynamic interplay of the neocortical and thalamic oscillators of slow waves is required for the full expression of this key physiological EEG rhythm.

Introduction

Slow waves and their neuronal counterpart, the cortical and thalamic oscillations between depolarized UP states and hyperpolarized DOWN states (Steriade et al., 1993a; Contreras and Steriade, 1995; Petersen et al., 2003; Sirotta and Buzsáki, 2005; Crunelli et al., 2012), are the main EEG hallmark of non-rapid eye movement (non-REM) sleep (Crunelli and Hughes, 2010; Brown et al., 2012) and are also observed during anesthesia (Chauvette et al., 2011). The physiological importance of these waves of natural sleep is emphasized by their ability to group together other EEG rhythms of non-REM sleep (Steriade, 1997) and by their putative role in the consolidation of recently acquired memories (Tononi and Cirelli, 2001; Marshall et al., 2006; Ji and Wilson, 2007).
animals. Moreover, our current mechanistic knowledge of slow waves of natural sleep continues to be clouded by the speculative extrapolations of findings obtained in anesthetized conditions.

Using a combination of neuronal ensemble recordings, thalamic-selective pharmacological inactivation, and optogenetic activation of thalamic cortical (TC) neurons in naturally sleeping or anesthetized rats, here we show, for the first time, that the thalamus is required for finely tuning the frequency of slow waves during non-REM sleep and anesthesia. Moreover, we demonstrate that the entrainment of EEG slow waves by selective thalamic activation is dependent on T-type calcium channels. Together, these results provide the first conclusive evidence that cortical and thalamic oscillators are necessary for the full expression of slow waves of non-REM sleep.

Materials and Methods

All experimental procedures were performed in accordance with the United Kingdom Animals (Scientific Procedure) Act 1986 and local ethics committee guidelines. All efforts were made to minimize animal suffering and the number of animals used. Experiments were performed on male adult Wistar rats (260 – 400 g, Harlan Laboratories), maintained on a normal diet and under an 8:00 A.M. to 8:00 P.M. light-on regimen. Experiments in anesthetized rats

**Surgery.** After anesthesia induction with 5% isoflurane, rats received an intraperitoneal injection of ketamine (120 mg/kg) and xylazine (20 mg/kg). Anesthesia was then maintained with a constant flow of ketamine (42 mg/kg/h) and xylazine (7 mg/kg/h) delivered via an intraperitoneal catheter connected to a pump (NewEra NE-300 syringe pump). Body temperature was maintained at 37°C with a heating pad and rectal probe. Rats were implanted with gold-plated skull screws (diameter 1 mm, length 3 mm) for EEG recordings in S1: anteroposterior (AP) = −2.2 mm, mediolateral (ML) = ±5.5 mm from bregma (Paxinos and Watson, 2007). An additional screw (ground electrode) was placed anterior to the bregma, and two other screws above the cerebellum were used as reference electrodes for EEG and thalamic recordings.

For measuring the spatial extent of the action of 3,5-dichloro-N-[1-(2,2-dimethyl-tetrahydro-pyran-4-ylmethyl)-4-fluoro-piperidin-4-ylmethyl]-benzamide (TTA-P2), a potent and selective T-type channel antagonist (Uebele et al., 2009; Dreyfus et al., 2010), or of the sodium channel blocker TTX in the thalamus (i.e., see experiments illustrated in Fig. 3), a 1 mm hole was drilled unilaterally above the ventrobasal thalamic nucleus (VB), the dura was carefully removed, and a 32-channel silicone probe (10 mm length, 0.6 mm width, 1–4 μm) with four shanks (200 μm recording point distance, 15 μm thickness) (NeuroNexus Technologies) was slowly lowered in the VB (dorsoventral [DV] = −4.5 to −5.5 mm). A second hole in the same hemisphere was used to slowly lower (200 μm every 5 min) a microdialysis probe (CMA 12 Elite, 2 mm dialysis membrane length, 20 kDa cutoff, with a 16° angle with respect to the vertical axis; see Fig. 3A) to a final position that was between 0.05 and 1 mm away from the silicone probe. The 16° angle positioning of the dialysis probe was dictated by the space constraints of the silicone and dialysis probes connecting devices on the animal skull.

For measuring the effect of TTX and TTA-P2 (applied by microdialysis in the VB) on slow and spindle waves during anesthesia (i.e., see experiments illustrated in Fig. 4), EEG electrodes were implanted as above, and two microdialysis probes (one in each VB) were slowly lowered fully vertically until their tips rested 6.5 mm below the skull (i.e., in the most ventral part of the VB). In some rats, a silicone probe was also inserted unilaterally in the VB with a 16° angle with respect to the vertical axis.

For measuring the effect of systemic TTA-P2 injection (i.e., see experiments illustrated in Fig. 5), rats were implanted with EEG electrodes and a unilateral silicone probe in the VB (as described above).

**Systemic and microdialysis solutions.** For intraperitoneal injection, TTA-P2 was dissolved in saline containing 4% DMSO and the pH adjusted with potassium hydroxide (1 m sys). Control intraperitoneal injections contained 4% DMSO in saline. For reverse microdialysis injection, TTA-P2 was dissolved in aCSF with 4% DMSO. Tetrodotoxin citrate (TTX) was dissolved in aCSF. Flow rate of the microdialysis injection was set at 1 μl/min.

**Injections.** For the systemic injections, once stable EEG slow waves were recorded for at least 30 min and high amplitude well isolated units were present in some of the silicone probe channels, a control period of at least 40 min was recorded before injecting intraperitoneally either saline/DMSO or TTA-P2 while continuing recording for at least another 2 h. For intrathalamic drug application, reverse microdialysis injection of aCSF was initiated as soon as the microdialysis probes were in position, and continued for at least 1 h before electrical recordings commenced. Once stable, high-amplitude, well-isolated units could be recorded from
some of the silicone probe channels, recording of a control period of at least 40 min was initiated. This was followed either by continuation of aCSF application or the inlet tubing was switched to a solution containing either TTX or TTA-P2. Recordings and dialysis application then continued for at least 1 h. At the end of the recording session, rats were transcardially perfused.

Recordings. Thalamic signals were amplified with an Omnetics preamplifier (gain 20, bandwidth 0.8 Hz to 54 kHz) and a Plexon recorder/64 channel amplifier (gain 7500–12,500, bandwidth 1–6000 Hz, Plexon). The EEG signal was amplified with a combination of SuperTech BioAmp (Pecs) pre- (bandwidth 0.1–500 Hz) and main-amplifiers (bandwidth DC to 500 Hz). When combined unit and EEG recordings were made, signals were digitized with a Plexon recorder/64 system at 20 kHz with 16-bit resolution. EEG recordings were digitized using the Plexon recorder input via the IP16 event input break-out panel. The digitized data were converted to Spike2 format (version 5.13, CED). For all further analyses, data were converted to a raw binary format using tools of the freely available Klusters, Neuroscope, and NDManager software suite (Hazan et al., 2006). EEG data were low-pass filtered with a windowed sinc filter at 100 Hz and downsampled to 200 Hz.

Experiments in freely moving rats

Surgery. Rats were anesthetized and implanted with EEG electrodes, as described above, and an EMG electrode was positioned in the neck muscle. Two or four microdialysis guide cannulae (one or two in each VB, respectively) (i.e., see experiments illustrated in Fig. 6 for two probes in each VB) were slowly lowered fully vertically so their tip was just above the VB (DV = −4.4 mm) (Paxinos and Watson, 2007). When two guide cannulae were inserted in each VB, their AP coordinates were as follows: −2.6 and −3.8 mm (Paxinos and Watson, 2007). Both EEG electrodes and guide cannulae were fixed to the skull with dental acrylic cement. Rats were allowed at least 7 d to recover from surgery (single housed) and to habituate to the recording cage (4 h/d). At the end of the recording sessions, rats were transcardially perfused.

Injections. For the systemic injections, on the day of the experiment rats were transferred to the recording cage, connected to the electrical recording and microdialysis apparatus (as appropriate), and allowed to move freely in their cage for at least 1 h before any recording commenced. A control period of 1.5 h was then recorded before an intraperitoneal injection of either saline or TTA-P2 was made while continuing recording for at least another 2 h. Four days were allowed between two consecutive recording days in each rat. For intrathalamic drug application, 24 h before recording, microdialysis probes were slowly inserted into the brain to replace the dummy probe in the guide cannulae. On the day of the experiments, rats were transferred to the recording cage and connected to the dialysis probes and electrical recording apparatus. aCSF dialysis was initiated immediately while the rats were allowed to habituate for 1 h. A control period of 2.5 h was then recorded while administering aCSF (same conditions as described for the anesthetized condition). Animals were then recorded for an additional 2 h while receiving either aCSF or drug-containing aCSF (in a random order on consecutive recording days, each separated by at least 4 d). Video recording was performed simultaneously with electrical recordings in all experiments in freely moving rats.

TTA-P2 levels in the VB. The concentration of TTA-P2 in samples taken from the inlet and outlet dialysis tubes was measured, following protein precipitation with acetonitrile, by liquid chromatography-mass spectrometry under a validated analytical protocol (Shipe et al., 2008; Uebel et al., 2009). TTA-P2 concentration in the brain tissue outside the dialysis membrane was estimated according to the equilibrium equation described previously (Chan and Chan, 1999).

Optogenetics

Viral injection. pAAV-CaMKIIα-bChR2(H134R)-mCherry plasmids (K. Deisseroth laboratory, Addgene plasmid 26975) were packaged into recom-
binant AAV2 vectors and serotyped with AAV1 coat proteins. Viral suspensions were titered to 1.14 × 10^10 genome copies/ml (GC) (University of Pennsylvania Vector Core). Concentrated stock virus was diluted with 0.1 M PBS tinted with Fast Green FCF (Sigma), giving a final viral concentration of 5.70 × 10^8 to 2.28 × 10^9 GC/μl for injection. The dorsal surface of the skull was exposed, a small craniotomy made over one VB (AP = −3.34 mm, ML = 2.80 mm from bregma) (Paxinos and Watson, 2007), and the dura reflected. A 10 μl Gastight Hamilton syringe and 34 Ga needle were front filled with mineral oil and diluted virus. Needles were then lowered slowly into the thalamus (DV = −5.75 mm from the pia) and left in place for 10 min. A 1 μl virus (5.70 × 10^8 to 2.28 × 10^9 infectious units) was injected at a rate of 100 nl/min using a programmable micro-pump (UMP3-1, WPI) and allowed to disperse for a further 10 min before the needle was slowly retracted. Rats were then individually housed and allowed to recover with ad libitum access to food and water for a minimum of 3 weeks to allow viral gene expression.

**Optical stimulation and recording.** Rats previously injected with AAV-CaMKII-ex-hChR2 (H134R)-mCherry were anesthetized and EEG screws and two microdialysis probes (one in each VB) were implanted (as above) with a 16° angle with respect to the vertical axis. A 200 μm multimode (0.39 NA) optic fiber (CFM121L20; Thorlabs) was custom-glued to a 32-channel silicone probe, with the fiber tip −400 μm above the top recording sites. The fiber ferrule was connected to a compatible patch-cord and 473 nm laser diode (70 mW Stradus; Vortran Laser Technology), and the silicon probe and EEG wires connected to a Digital Lynx 10SX recording system (with Hybrid Input Boards; Neuralynx) via HS-36 unity gain preamplifiers. One optrode was slowly lowered to just above one VB, 300 μm posterior and medial to the virus injection site (AP = −3.44 mm, ML = 2.50 mm from bregma) (Paxinos and Watson, 2007), and DV = −4.60 mm from the pia, and then moved in small steps to locate light-responsive ChR²⁺ cells exhibiting characteristic TC neuron bursts (see experiments illustrated in Figs. 9 and 10). Extracellular action potentials (sampling frequency: 32 kHz per channel, filtered from 600 Hz to 9 kHz), continuous extracellular signal (sampling frequency: 32 kHz per channel and broadband filtered from 0.1 Hz to 9 kHz), EEG signals (sampling frequency: 4 kHz, filtered from 0.1 Hz to 1 kHz), and light stimulation events were simultaneously recorded using Cerebex 5 Data Acquisition software (Neuralynx). Digital laser modulation was controlled with pclamp software and a 1322A Digitida (Molecular Devices), synchronized with the Digital Lynx 10SX. Laser output power for 5, 20, and 100 ms pulses was 40 mW, which equated to <10 mW at the fiber tip (dependent on stimulation frequency), quantified with a digital power meter and photodiode sensor (PM120D; Thorlabs).

**Data analysis.** Spike sorting and data preprocessing were performed with the Klusters, Neuroscope, NDManager software suite (Hazen et al., 2006). All other analyses were performed with routines based on the free toolboxes Scipy 0.8 (Jones et al., 2001), OpenElectrophy 0.2 (Garcia and Fourcaud-Trocmé, 2009), running under Python 2.6.6 and MATLAB (R2010b, MathWorks) on a 64-bit Linux computer.

Data preprocessing and spike sorting. To extract spikes from the extracellular field potential, the signal was high-pass filtered with median filter (0.5 ms window half-length). Spikes were detected by thresholding at 1.2 SD and clustered by an expectation maximization algorithm (Klustakwik; Harris et al., 2000) on the basis of their first three principal components. All results of the automatic clustering were verified post hoc by visual inspection. Units were excluded from further analyses if more than one high amplitude cell was present on a single channel, if their autocorrelogram did not show a refractory period of at least 2 ms, and if their spike amplitude markedly changed during the experimental session.

**Burst analysis.** Bursts were defined as two or more spikes that were preceded by at least 100 ms of silence and had interspike intervals (ISIs) <10 ms, and were visually examined post hoc. For each burst, the following parameters were calculated (Fig. 1): (1) ISI, (2) interburst interval, (3) number of spikes per second in 5 min windows (spike rate), (4) number of bursts per second in 5 min windows (burst rate), (5) ratio of all spikes taking part in a burst, (6) number of spikes in each burst, and (7) ISI as a function of the ISI number within a burst (i.e., burst signature). Units with a decelerando burst signature (Fig. 1C,D) were classified as TC neurons, whereas those with a burst signature and spike autocorrelogram typical of thalamic reticular neurons (Huguenard and Prince, 1992) were discarded.

**Detection of slow waves and spindles under anesthesia.** To quantify EEG slow waves beyond power spectral analysis, a slow wave detection algorithm similar to those described previously (Mölle et al., 2009; Nir et al., 2011) was implemented (Fig. 2A–D). On the 0.2 to 4.5 Hz bandpass-filtered signal, all negative to positive zero-crossings were detected as slow waves. To discard spurious slow waves, the local minimum and maximum around a crossing were determined. If the difference between these was <60% of the mean maximum-to-minimum distance, the slow wave was discarded.
wave was discarded. For the analysis of UP and DOWN state duration, the periods above and below the midpoint of the DOWN to UP transitions in the bandpass-filtered signal were labeled as UP and DOWN states, respectively. States with a magnitude of <25% of the average magnitude during the control period were discarded and the neighboring states of equal type (UP or DOWN) merged. Spindle waves were detected in the frequency range 5–12.5 Hz (Fig. 2E, F).

Spectral analysis. Power spectral densities were estimated with Welch’s average periodogram method using a Hanning window with length of 4096 data points. Total power in a frequency band was determined as the integral over that frequency band.

Detection of non-REM sleep epochs. The power of the EEG signal between 0.5 and 4 Hz was computed for every 10 s of recording. If the calculated power was higher than twice the power calculated during a period of active wakefulness, the 10 s epoch was classified as non-REM sleep. To ensure detection reliability and behavioral state stability over the selected periods, only consecutive epochs of sleep lasting at least 60 s were included in further analysis.

Detection of slow waves and spindles during natural sleep. The EEG signal was convolved with complex Morlet wavelets of 2.5 cycles for sleep. Power spectral densities were estimated with Welch’s method, each oscillatory epoch of the EEG was extracted with an energy threshold to detect its beginning and end (see Figs. 2G, 6A, and 7A) (Roux et al., 2007; García and Fourcaud-Trocme, 2009). The boundary frequencies of wave detection were chosen as from 0.5 to 4 Hz for slow waves (see Fig. 6B) and from 6 to 14 Hz for spindle oscillations (see Fig. 2G,H). The threshold was defined as 3 times the average energy during a non-REM sleep period during the control session. Slow and spindle waves with <2 and 3 cycles, respectively, were discarded. When overlapping oscillations were detected, the wave with the highest energy was selected.

Statistical analysis. Group comparisons were performed using the Mann–Whitney U test. Paired data were tested with Wilcoxon’s signed rank test. All quantitative data in figures and text are given as mean ± SEM.

Histology
Electrode and microdialysis probe tracking. Before insertion, silicone and microdialysis probes were immersed in 1/2 h and 5 min, respectively, in a 1% Vybrant Dil (Invitrogen) dye solution. At the end of the experiment, rats were injected with a lethal dose of urethane (40%), and the brains were removed and placed in a 4% PFA solution for 48 h. The brains were then transferred and stored in a 0.1 M PBS. Sections (100 μm thick) containing the VB were cut with a vibratome (Leica VT1000S) and mounted on coverslips to measure the relative position of microdialysis and silicone probes, which were visualized using a fluorescent microscope (Leica).

Immunofluorescence. Rats were given an overdose of ketamine-xylazine and transcardially perfused with 4% PFA. Brains were fixed in 4% PFA and then cryoprotected in 20% (w/v) sucrose in 0.1 M PBS. Each brain was blocked to give coronal sections, mounted onto a freezing microtome (Leica), and cut into 50 μm sections. Free-floating sections were processed for NeuN and RFP immunofluorescence. Briefly, sections were washed 3 × 10 min with fresh 0.1 M PBS and blocked for 1.5 h in 1 × PGT (0.1 M PBS, 3% NGS, and 0.2% Triton X-100 (Sigma-Aldrich)). Sections were gently shaken at room temperature for 2 h, then overnight at 4°C with primary antibodies against both NeuN (1:500 mouse monoclonal, Millipore), and RFP (1:1000 Living Colors DsRed rabbit polyclonal, Clontech) in 1 × PGT. Sections were rinsed 3 × 10 min with fresh 0.1 M PBS and incubated for 2 to 3 h at room temperature with 1:200 goat anti-mouse AlexaFluor-488 (Invitrogen) and 1:200 goat anti-rabbit AlexaFluor-594 (Invitrogen) secondary antibodies in 1 × PGT. Sections were washed 3 × 10 min in fresh 0.1 M PBS and mounted onto gelatin-subbed Superfrost Plus microscope slides (Thermo Scientific). Slides were coverslipped with VectaShield fluorescent mounting medium (Vector Laboratories) and visualized using a fluorescence microscope (Leica).

Results
Effect of thalamic inactivation on slow waves during anesthesia
We first abolished the somatosensory thalamic output to the cortex by bilateral reverse microdialysis of TTX in the ventrobasal thalamus (VB) of ketamine-xylazine anesthetized rats, while
simultaneously recording slow waves from primary somatosensory cortex (S1) and the firing of different single TC neurons in the VB. TTX abolished action potentials in TC neurons around the VB injection site (predrug: 2.51 ± 0.11 spikes/s, TTX: 0.001 ± 0.002, n = 11 rats, n = 33 neurons, p < 10⁻⁷, Wilcoxon signed-rank test) (Fig. 3B1,B2, blue lines, C, bottom plot; see also Fig. 4A, unit recordings). This effect led to a concomitant marked reduction (45 ± 1.4%) in the number of EEG slow waves after 1 h of drug dialysis (Fig. 4A,B) and thus to a frequency decrease from 1.19 ± 0.02 to 0.63 ± 0.03 slow waves/s (n = 5, p = 0.0017, Mann–Whitney U test compared with aCSF, n = 6) (Fig. 4A–D). The duration of both UP and DOWN states was increased by TTX from 0.40 ± 0.01 (UP) and 0.42 ± 0.03 s (DOWN) to 0.68 ± 0.07 and 0.58 ± 0.06 s, respectively (p < 0.006, Mann–Whitney U test compared with aCSF) (Fig. 4B). In the same rats, the simultaneously recorded spindle waves, which are a thalamically generated rhythm (Morison and Bassett, 1945; Steriade et al., 1985; De Gennaro and Ferrara, 2003; Astori et al., 2011), were abolished by TTX (predrug: 0.13 ± 0.02 spindles/s, TTX: 0.009 ± 0.008, p = 0.0016) (Fig. 4E), indicating the effectiveness of our thalamic inactivation by microdialysis administration of TTX.

Because T-type calcium channels play a key role in the thalamic output to cortex (Llinás and Jahnsen, 1982; Crunelli et al., 1989; Deleuze et al., 2012) and underlie TC neuron UP states (Hughes et al., 2002), we next investigated the effect on slow waves of the potent and selective T-type calcium channel antagonist, TTA-P2 (Uebele et al., 2009; Dreyfus et al., 2010), directly applied in the VB by reverse microdialysis. This drug produced a block of high-frequency bursts of TC neurons, which was dependent on its concentration in the dialysis inlet tube and on the distance between the recorded neuron and the dialysis probe (Fig. 3B1; see also unit recordings in Fig. 4A). Interestingly, single action potential firing (which accounted for ~18.4 ± 1.2% of the total firing) was also markedly decreased by TTA-P2 (Fig. 3B2) (see Fig. 4A, unit recordings) so that, at a distance of 500 μm from the dialysis probe, only 3.3 ± 2.1% of total spikes (i.e., in bursts and as single action potentials) remained after 1 h of TTA-P2 application. We chose a microdialysis inlet tube concentration of 300 μM TTA-P2 because: (1) it virtually abolished the total TC neuron firing in a region that almost fully covered the mediolateral extent of the VB (Fig. 3C,D), and (2) it resulted in a tissue concentration around the probe of ~42 μM (based on a 14% recovery rate) (for explanation, see Fig. 3E), which is well in the range of concentrations that we previously showed to be required to abolish intrinsic and synaptically driven TC neuron firing in the VB in vitro (Dreyfus et al., 2010, their Fig. 1C). Thus, TTA-P2 applied in the VB at 300 μM elicited a clear reduction (25 ± 3%, n = 5, p = 0.02 compared with aCSF, n = 6) of slow waves (Fig. 4A,B), leading to a frequency decrease from 1.00 ± 0.02 to 0.69 ± 0.03 slow waves/s (p = 0.020) (Fig. 4A,C,D). The durations of UP and DOWN states were both prolonged from 0.59 ± 0.04 (UP) and 0.58 ± 0.02 (DOWN) to 0.77 ± 0.06 and 0.70 ± 0.03 s, respectively (p < 0.05, Mann–Whitney U test compared with aCSF) (Fig. 4B). Moreover, in the same experiments, spindle waves were markedly suppressed (88 ± 5%) by microdialysis injection of TTA-P2 in the VB (aCSF: 0.15 ± 0.01 spindles/s; TTA-P2: 0.019 ± 0.008; p = 0.02) (Fig. 4E), indicating the effectiveness of our thalamic inactivation by microdialysis administration of TTA-P2.

TTA-P2 also elicited a dose-dependent decrease in slow waves when applied systemically (Fig. 5A–D), with a similar ED₅₀ on burst (0.18 ± 0.05 mg/kg) (Fig. 5E) and total firing (0.26 ± 0.06 mg/kg) (Fig. 5F). In particular, at a dose (3 mg/kg) that abolished
burst firing in the VB (Fig. 5E), TTA-P2 produced a stronger decrease (63 ± 1%, n = 3, p = 0.018) of slow waves (Fig. 5B) than an equipotent (on burst firing) intra-VB application (compare with Fig. 3B1), indicating that T-type calcium channels in other thalamic nuclei and brain regions effectively contribute to the full expression of EEG slow waves in S1 during anesthesia.

Effect of thalamic inactivation on slow waves during natural sleep
Because anesthesia does not fully reproduce the spatiotemporal dynamics of slow waves during non-REM sleep (Chauvette et al., 2011; Nir et al., 2011; Vyazovskiy et al., 2011), we then applied TTX and TTA-P2 by reverse microdialysis in the thalamus of naturally sleeping-waking rats. The same inlet dialysis concentration of aCSF, 3 mM TTA-P2, and 50 μM TTX was used. Transient slow waves (white lines) were detected as ridgets in the wavelet spectra (see Materials and Methods) (Fig. 2B). Slow wave frequency density distribution during aCSF, TTA-P2, and TTX dialysis, 1 h after the start of drug dialysis (arrows indicate the measured peaks). C, Time dependence of TTA-P2 (n = 7 rats) and TTX (n = 5 rats) effects on the normalized peak of the slow wave frequency distribution. D, Raw non-REM EEG power spectra show TTA-P2- and TTX-elicited decrease of power in sleep spindle frequency range and increase of power in slow wave frequency range. E, Time dependence of TTA-P2 and TTX effects on sleep spindles normalized count. B, D, Solid lines indicate the mean; color shading indicates SEM. C, E, Error bars indicate SEM.

Figure 6. Block of thalamic firing decreases slow wave frequency during natural sleep. A, Non-REM sleep EEG (bottom) and corresponding wavelet spectra (top) during VB microdialysis of aCSF, 3 mM TTA-P2, and 50 μM TTX. Transient slow waves (white lines) were detected as ridgets in the wavelet spectra (see Materials and Methods). B, Slow wave frequency density distribution during aCSF, TTA-P2, and TTX dialysis, 1 h after the start of drug dialysis (arrows indicate the measured peaks). C, Time dependence of TTA-P2 (n = 7 rats) and TTX (n = 5 rats) effects on the normalized peak of the slow wave frequency distribution. D, Raw non-REM EEG power spectra show TTA-P2- and TTX-elicited decrease of power in sleep spindle frequency range and increase of power in slow wave frequency range. E, Time dependence of TTA-P2 and TTX effects on sleep spindles normalized count. B, D, Solid lines indicate the mean; color shading indicates SEM. C, E, Error bars indicate SEM.

Figure 7. Systemic injection of TTA-P2 markedly decreases the frequency of slow waves and abolishes spindles during natural sleep. A, Non-REM sleep EEG (middle), corresponding wavelet spectra (top), and EMG (bottom) after intraperitoneal injection of saline (left) and 10 mg/kg TTA-P2 (right). Transient slow waves (white lines) were detected as ridgets in the wavelet spectra as in Figure 2. B, Slow wave frequency distribution after saline and TTA-P2 injection. There is a shift of the peak (arrows) from ~2 Hz to ~0.7 Hz. C, Time dependence of TTA-P2 (n = 4 rats) and saline (n = 6 rats) effects on the normalized peak of the slow wave frequency distribution. D, Raw non-REM sleep EEG power spectra show TTA-P2-elicited decrease of power in sleep spindle frequency range and increase of power in slow wave frequency range compared with saline injection. E, Time dependence of TTA-P2 effects on sleep spindles normalized count. F, G, Dose–response curve of burst (F) and total spike (G) rate measured 40 min after systemic intraperitoneal injection of TTA-P2 (logistic regression fits, p < 0.05) (ED<sub>50</sub> for bursts: 0.55 ± 0.03 mg/kg; ED<sub>50</sub> for total spikes: 1.71 ± 0.11 mg/kg). The 3 and 10 mg/kg TTA-P2 abolish bursts recorded during natural sleep (p < 10<sup>-6</sup> compared with saline injection, Mann–Whitney U test, n = 42 TC neurons). B, D, Solid lines indicate the mean; color shading indicates SEM. C, E–G, Error bars indicate SEM.

Produced a significant decrease (23 ± 7.8%, n = 7, p = 0.0095, Mann–Whitney U test compared with aCSF, n = 12) in the peak of the instantaneous frequency distribution of slow waves during non-REM sleep (predrug: 2.19 ± 0.09 Hz; TTA-P2: 1.72 ± 0.13 Hz), as did TTX (50 μM) (TTX: 0.92 ± 0.13 Hz, 50 ± 13% reduction, n = 5, p = 0.0061) (Fig. 6A–D). As expected, in the same animals, sleep spindles were more potently decreased by TTX and TTA-P2 (94 ± 7% and 48 ± 6%, respectively) than slow waves (predrug: 0.36 ± 0.02 spindles/s; TTX: 0.05 ± 0.03, TTA-P2: 0.19 ± 0.02; p = 0.001 for both compared with aCSF) (Fig. 6D, E).
As it was observed during anesthesia, also in naturally sleeping rats systemic injections of TTA-P2 dose-dependently blocked burst and total firing in VB TC neurons during non-REM sleep (with an ED$_{50}$ of 0.55 ± 0.03 and 1.71 ± 0.11 mg/kg, respectively) (Fig. 7F,G), as well as slow waves and sleep spindles (Fig. 7A–E). In particular, a dose of 10 mg/kg of TTA-P2 markedly reduced slow wave (saline: 1.94 ± 0.13 Hz, n = 6; TTA-P2: 0.8 ± 0.01 Hz, n = 4 rats) and sleep spindle frequencies (saline: 0.32 ± 0.13 spindles/s, TTA-P2: 0.05 ± 0.01) by 73 ± 5% (p = 0.0095) and 86 ± 4% (p = 0.00031), respectively (Fig. 7E).

**Entrainment of EEG slow waves by optogenetic stimulation of TC neurons during anesthesia**

Because thalamic inactivation by either TTX or by selective block of thalamic T-type calcium channels with TTA-P2 decreased the frequency of slow waves during anesthesia and natural sleep, the thalamic input to the neocortex should be able to entrain these EEG waves. Because electrical stimulation of the thalamus leads to antidromic excitation of the somatotopic cortical region, which may in turn affect the cortical slow wave oscillator, we addressed this question using selective optogenetic activation of channelrhodopsin2-expressing TC neurons (Fig. 8) with short (5, 20, or 100 ms) 473 nm light pulses at stimulation frequencies that ranged from 0.75 to 4 Hz (n = 7 rats) (Fig. 9). At the cellular level, each light pulse elicited a single high-frequency burst of action potentials, which was invariably followed by a 100–250 ms period of electrical silence before firing resumed (Fig. 9C). For stimulation frequencies from 0.75 to 1.5 Hz, this pattern of TC neuron activation elicited a clear peak in the EEG wavelet (Fig. 9A) and FFT power spectrum (Fig. 9D) at the respective stimulation frequency, which had higher amplitude than the peak of the control EEG (i.e., with no light stimulation) (Fig. 9D, black line). As seen from the light-pulse triggered averages (Fig. 9B), light stimulation for frequencies >1.5 Hz not only failed to entrain EEG slow waves but also markedly flattened the power spectra eliminating the peak present at ~1 Hz during the control condition (i.e., without light stimulation; Fig. 9D, black line). When T-type calcium channels in VB were blocked by microdialysis of 300 μM TTA-P2 (n = 3 rats) (Fig. 10A), the cellular burst response upon stimulation was virtually abolished (Fig. 10B), and no entrainment of slow waves occurred for stimulation frequencies between 0.75 and 1.5 Hz (Fig. 10C,D). Moreover, a similar block of EEG slow wave entrainment was observed after systemic injection of TTA-P2 (3 mg/kg) at these stimulation frequencies (data not shown). Finally, no significant effect of TTA-P2 injected locally (Fig. 10C2,D) or systemically (data not shown) was observed at stimulation frequencies ≥ 1.75 Hz.

**Figure 8.** Channelrhodopsin-2 expression in VB TC neurons. A, Immunostaining of channelrhodopsin-2-mCherry (red) showing that the expression of the channelrhodopsin-2 protein is restricted to VB TC neurons and some TC axons passing through the NRT (top). Arrow indicates the putative site of the virus injection. NeuN staining (green) is evident in both VB and NRT somata (middle). Merged images (bottom) demonstrate colocalization of channelrhodopsin-2 and NeuN in TC, but not NRT, neurons. B, Higher magnification of a portion of the respective panels in A. Scale bars, A, B: 300 μm.

**Discussion**

Our results conclusively demonstrate that the full manifestation of EEG slow waves during non-REM sleep in freely moving, naturally waking-sleeping rats requires a dynamic interplay of cortical and thalamic neuronal ensembles because inactivation of the thalamic output to the neocortex brings about a marked deceleration of slow waves and selective excitation of TC neurons entrains EEG slow waves.

**Contribution of thalamic oscillators to sleep rhythms**

There is a general consensus that an isolated neocortex can generate and maintain the UP and DOWN state dynamics that underlie EEG slow waves of natural sleep via an intricate balance of excitation and inhibition that is mostly generated by synaptically driven cortical slow wave oscillators (Sanchez-Vives and McCormick, 2000; Timofeev et al., 2000; Bazhenov et al., 2002; Cossart et al., 2003; Shu et al., 2003; Le Bon-Jego and Yuste, 2007; Beltramo et al., 2013). In line with this interpretation, the cortical firing (during UP states) and the electrical silence (during DOWN states) impose similar UP/DOWN state transitions on thalamic neurons, making the thalamus fully subservient to corticofugal activity. Over the last 10 years, however, this view has been challenged by solid experimental evidence demonstrating that an isolated thalamus as well as single TC and nucleus reticularis thalami (NRT) neurons in vitro can sustain slow waves and an intrinsic UP/DOWN states dynamics, respectively, which are similar to those observed during natural sleep (Crunelli...
The full physiological rhythm might, indeed, be the result of two oscillators (Crunelli and Hughes, 2010) because the intrinsic slow wave thalamic oscillator increases its frequency with the progressive hyperpolarization that accompanies non-REM sleep deepening (Hughes et al., 2002; Crunelli and Hughes, 2010). Support to the thalamic modulation of EEG slow waves also comes from data showing that an acceleration of these waves is accompanied by a phase shift of the TC neuron firing toward the start of an UP state (Slézia et al., 2011).

In the same animals where slow waves of non-REM sleep were decreased in frequency, the simultaneously recorded sleep spindles, which are a thalamically generated rhythm (Morison and Bassett, 1945; Steriade et al., 1985; De Gennaro and Ferrara, 2003; Astori et al., 2011), were abolished by inactivation of the same thalamic region. This demonstrates that, whereas both cortex and thalamus are required for the full expression of slow waves and spindles of natural sleep, the relative contribution of these brain areas to these oscillations is markedly different, reflecting the diverse cellular/network generators that underlie these two EEG rhythms.

By comparing, for the first time, slow waves during natural sleep and anesthesia under the same laboratory conditions, we could identify that a larger area of thalamic inactivation was necessary to obtain a significant effect on slow waves during non-REM sleep compared with anesthesia. This indicates that the thalamic modulation of slow waves during anesthesia is very different from that during natural sleep, stressing the diverse nature of the cortical and thalamic neuronal dynamics underlying these behavioral states, the limitations associated with extrapolating results from one experimental condition to the other, and a potentially different involvement of the “core” and “matrix” thalamic projection systems (Jones, 2001).

**Figure 9.** Thalamic entrainment of EEG slow waves during anesthesia. A, EEG trace (bottom) and wavelet transform (top) showing the effect of 20 ms, 473 nm light pulses at 1, 1.5, and 2 Hz. Dashed white line indicates the start of the first pulse. B, Event-triggered EEG averages centered on the 20 ms light pulses (blue vertical bars) for the illustrated stimulation frequencies. Gray areas indicate SEM. A total of 10 s of EEG was used for each average. C, Raster plots of firing of 4 VB TC neurons in response to two consecutive 20 ms pulses (at 1 Hz) (blue vertical lines). *Bursts. The corresponding EEG trace is superimposed as a black line. D, EEG power spectra in response to 20 ms light pulses at the illustrated frequencies (indicated by arrows). There are frequency-dependent amplification and the shift of the peak of the power spectra compared with control (i.e., without light stimulation, black line).

and Hughes, 2010). This thalamic activity only occurs when the metabotropic glutamate receptors (which are present postsynaptically to the corticofugal terminals) of TC and NRT neurons (Godwin et al., 1996) are active, making these neurons “conditional oscillators” for slow waves (Crunelli and Hughes, 2010).

The present findings strongly support this hypothesis: removal of the thalamic input to cortex reduces the frequency of slow waves of non-REM sleep, demonstrating, for the first time, a necessary role for the thalamus in this physiological rhythm. The generation of slow waves during natural sleep, therefore, derives from cortical networks as primary oscillators with the thalamic oscillator providing a fine-tuning by imposing its faster frequency and phase properties to the cortical slow wave rhythm. The full physiological rhythm might, indeed, be the result of two competing oscillators (Gutierrez et al., 2013), with the thalamus being able to reset the phase of the slower cortical networks, as indicated by the ability of selective optogenetic excitation of thalamofugal fibers to entrain EEG slow waves (this work) and to induce isolated UP states in awake, head-restrained rats (Poulet et al., 2012). Importantly, the ability of the thalamus to modulate cortical UP states depends on thalamic T-type calcium channels, and in particular on the high-frequency burst of action potentials that invariably marks the start of TC neuron UP states (Contreras andSteriade, 1995, their Fig. 9; Slézia et al., 2011; Ushimaru et al., 2012). The effect of this thalamic modulation will also be important in determining the slight acceleration of slow waves observed from sleep Stage 2 to 4 (Crunelli and Hughes, 2010; Brown et al., 2012) because the intrinsic slow wave thalamic oscillator increases its frequency with the progressive hyperpolarization that accompanies non-REM sleep deepening (Hughes et al., 2002; Crunelli and Hughes, 2010). Support to the thalamic modulation of EEG slow waves also comes from data showing that an acceleration of these waves is accompanied by a phase shift of the TC neuron firing toward the start of an UP state (Slézia et al., 2011).

**Thalamic T-type calcium channels and the slow rhythm**

Our investigation is the first to provide direct evidence that the T-type calcium channels of thalamic (i.e., TC and NRT) neurons are required for the full expression of slow waves during natural sleep, although we could not distinguish between TC and NRT neuron contribution because of TTA-P2 spread to the latter neuronal population at the highest concentrations. A previous study reported an increased number of awakenings in mice carrying a supposed thalamic-selective deletion of CaV3.1 T-type channels expression (Anderson et al., 2005). However, the presence of recombination in piriform cortex, some hypothalamic nuclei, and other brain areas questions the selectivity of this genetic approach and weakens its conclusions. Another study in anesthe-
tized CaV3.1 T-type calcium channel KO mice (Lee et al., 2004) suggested a contribution of these channels to δ waves (defined as power in the 1–4 Hz frequency band) but not to slow waves (defined as power in the <1 Hz frequency band). Our results in anesthetized rats instead indicate that slow waves peaking at ~1 Hz are decreased in number by TTA-P2, leading to a slowing down in frequency to ~0.6 Hz. Although these contradictory data may be the result of the use of different species or anesthetics, they may also result from the different wave classifications. Indeed, it is debatable (Luczak and Barthö, 2012) whether EEG waves occurring at the upper end of the δ frequency range (2–4 Hz) in naturally sleeping rats do represent a separate entity from slow (<2 Hz) waves because the underlying UP and DOWN state firing dynamics of slow waves in the 2–4 Hz band is similar to that in 0.5–2 Hz band (Ji and Wilson, 2007, their Fig. 2).

It is not surprising that the vast majority of the total TC neuron firing (i.e., high-frequency bursts plus single action potentials) depends on T-type calcium channels (i.e., it is blocked by TTA-P2). Indeed, although these channels are classically viewed as underlying only high-frequency bursts of TC neurons, recent evidence has indicated that the continuous opening of a very small number of T-type channels that occurs at ~−60 mV generates a depolarizing window current (Dreyfus et al., 2010, their Fig. 2) that is crucial for the maintenance of the UP state (Hughes et al., 2002; Crunelli and Hughes, 2010). This strong reliance of the thalamic output on T-type channels also indicates that during slow waves synaptic activity, and in particular cortical inputs to the thalamus, are not sufficient to drive a TC neuron output in the absence of these channels.

Resetting and entrainment of EEG slow waves can be achieved by whisker stimulation (Civitico and Contreras, 2012) or by optogenetic activation of cortical neurons in anesthetized rats (Kuki et al., 2013), whereas long (>4 s) optogenetic stimulation of TC neurons in head-restrained mice can induce UP states in neocortical neurons that do not outlast the light stimulus (Poulet et al., 2012). Our results show that short (5 ms) optogenetic stimuli, which evoke high-frequency bursts in TC neurons, strongly entrain EEG slow waves, and that this response is only present when thalamic T-type calcium channels are functionally active. This strengthens the suggestion that thalamic T-type channel-mediated bursts may critically control UP states in related cortical territories during non-REM sleep by finely tuning the frequency of the EEG slow waves that occur during this behavioral state (Crunelli and Hughes, 2010). In contrast, a recent imaging study has reported that thalamic slow waves always follow cortical slow waves during anesthesia (Stroh et al., 2013). However, these results shed little light on the relative contribution of thalamic and cortical neuronal ensembles to slow waves of natural sleep, because, as indicated by Wester and Contreras (2013), they may be confounded by the experimental conditions, in particular the use of an anesthetic agent that is known to markedly block T-type calcium channels (Jokovic and Todorovic, 2010; Eckle et al., 2012).

In conclusion, these findings provide conclusive demonstration of our hypothesis (Crunelli and Hughes, 2010) that both cortical and thalamic population activities are required for the full manifestation of EEG slow waves of natural sleep, and suggest that a decreased thalamic output to the neocortex resulting from thalamic pathologies (Schmahmann, 2003; Koppelmann et al., 2009; Parneau et al., 2013) will bring about a slowing down of slow waves during non-REM sleep with deleterious consequences for memory processes.

References


