Introduction

A wealth of data indicate a major role of GABAergic cells in pacing various rhythms via GABA-A receptors (GABA-ARs) in the cortex (Sanchez-Vives and McCormick, 2000; Blatow et al., 2003), hippocampus (Cobb et al., 1995; Wulff et al., 2009), and thalamus (von Krosigk et al., 1993). GABA-ARs are classified into two fundamentally distinct types of receptors, synaptic and extrasynaptic receptors that mediate phasic and tonic inhibition, respectively. Exactly what type of inhibition governs oscillatory activity is presently unclear. The pentameric GABA-ARs can be assembled in theoretically limitless combinations with various types of subunits (Farrant and Nusser, 2005). For the synaptic clustering of GABA-ARs, the γ2 subunit is essential (Essrich et al., 1998). Receptors containing the γ2 subunit typically have rapid kinetics, low sensitivity to GABA, and prompt desensitization, enabling them to conduct fast inhibitory postsynaptic events, typical for phasic inhibition (Brickley et al., 1999). GABA-ARs containing δ instead of γ subunits are abundant in extrasynaptic membranes (Nusser et al., 1998; Wei et al., 2003; Farrant and Nusser, 2005; Belelli et al., 2009), have higher sensitivity to GABA, and are activated by ambient GABA. These extrasynaptic receptors are primarily responsible for a slow, persistent (i.e., tonic) chloride current (Hamann et al., 2002; Wei et al., 2003; Cope et al., 2005; Bright et al., 2007, 2011).

More recently, evidence is accumulating for a third type of GABA-AR-triggered inhibition. This spillover-mediated transmission is caused by diffusion of GABA from the synaptic cleft and allows activation of GABA-ARs in extrasynaptic locations (Capogna and Pearce, 2011), resulting in phasic inhibition with slowed-down kinetics. The time course and amplitude of synaptic inhibition are an important determinant of the oscillation frequency; therefore, it is critical to determine the contribution of these three types of GABA-AR-mediated inhibition to naturally occurring brain rhythms.

Thalamus is one of the brain regions abundantly expressing both synaptic and extrasynaptic receptors and it is known to be involved in large-scale brain oscillations important for states of vigilance, notably sleep. Thalamocortical (TC) cells express only a few of the potential combinations of GABA-AR subunits (Pirker et al., 2000). Synaptic receptors mainly incorporate α1,
β2/3, and γ2 subunits, whereas nonsynaptic receptors contain α4, β2/3, and 6 subunits. Major pacemakers for rhythms in thalamic circuits are the GABAergic cells of the nucleus reticularis thalami (nRT). Recent in vitro data indicate that extrasynaptic receptors can be recruited during nRT-mediated phasic IPSCs (Herd et al., 2013). This raises the possibility that sleep-related oscillations are partially driven by a population of molecularly distinct GABA-ARs located outside the synapse. However, whether and to what extent phasic activation of extrasynaptic GABA-ARs is capable of driving normal rhythms in vivo are not clear. As many sedative–hypnotic agents target synaptic GABA-ARs (Winsky-Sommerer, 2009), elucidating the contribution of nonsynaptic GABA-Rs to sleep oscillations is of great interest for further drug development. In this study, we unraveled a significant role for extrasynaptic receptors in sleep oscillations by using localized, virus-mediated gene-knock-out of synaptic γ2-GABA-ARs in the thalamus, leaving only extrasynaptic inhibition intact.

**Materials and Methods**

*Mice and stereotactic surgery.* Adult GABAα2β2γ1lox/lox and C57BL/6J mice from both genders were used for the experiments. Mice were entrained to a 12 h light/dark cycle with food and water available *ad libitum*. All experimental procedures were performed according to the ethical guidelines of the Institute of Experimental Medicine of the Hungarian Academy of Sciences and the University of Lausanne and approved by the local Ethical Committees and Veterinary Offices of the Canton de Vaud. Surgeries and experiments were done under ketamine/xylazine anesthesia and experiments were done under ketamine/xylazine anesthesia and experiments were done under ketamine/xylazine anesthesia and experiments were done under ketamine/xylazine anesthesia and experiments were done under ketamine/xylazine anesthesia and experiments were done under ketamine/xylazine anesthesia and experiments were done under ketamine/xylazine anesthesia and experiments were done under ketamine/xylazine anesthesia and experiments were done under ketamine/xylazine anesthesia and experiments were done under ketamine/xylazine anesthesia and experiments were done under ketamine/xylazine anesthesia and experiments were done under ketamine/xylazine anesthesia and experiments were done under ketamine/xylazine anesthesia and experiments were done under ketamine/xylazine anesthesia. The primer antibiotics were visualized with corresponding fluorescent secondary antibodies (Alexa-488-, -Cy3-, or-Cy5-conjugated IGs). Finally, sections were mounted on slides and covered with Vectashield (Vector Laboratories).

For the visualization of GABA-AR subunits after the treatment with one of the anti-Cre antibodies, the sections were treated with biotinylated goat anti-mouse or rabbit secondary antibody (1:300, at room temperature for 2 h) and amplified with Tyramide (Invitrogen, 1:50, 15 min). Next, the sections were digested in pepsin (0.2–0.3 mg/ml in 20% HCl solution made with 0.1 M PB; 7 min at 37°C), then incubated in rabbit anti-γ2 (1:4000, Synaptic System), rabbit anti-α1 (1:500, W. Sieghart, Brain Research Institute, Vienna, Austria), or guinea-pig anti-α1 (1:500, Frontier Institute) or rabbit anti-α4 (1:500, W. Sieghart, Brain Research Institute, Vienna, Austria) overnight at room temperature. The Cre immunostaining was visualized with streptavidin-conjugated Alexa-488, while the other antibodies with corresponding Cy3 or Cy5-conjugated secondary IGs (at room temperature for 2 h). Finally, the sections were mounted.

Photographs were taken using either Zeiss Axioplan 2 equipped with DP70 digital camera (Olympus), Olympus Fluoview FV1000 or Nikon ECLIPSE Ni-E confocal laser scanning systems. For the quantitative receptor localization, AAV-Cre (n = 4) and AAV-GFP (n = 4) injected animals were used. In each animal, 3–33 regions of interest (ROIs, area 23,000–30,000 μm²) were analyzed. To determine the abundance of GABA-AR subunits, the images were thresholded and gray values of pixels were registered in ROIs defined in appropriate optical sections of image stacks in Fiji (Schindelin et al., 2012); then the data were exported to a spreadsheet. For the preparation of figures, when necessary, brightness and contrast were adjusted using Adobe Photoshop CS4 (Adobe Systems) applied to whole images only.

For data acquisition of Stochastic Optical Reconstruction Microscopy (STORM) images, the method described by Dani et al., 2010 was used; 20 μm sections (without freeze thawing) were treated similarly as described above with the following exceptions. Normal horse serum (10%) was used for blocking, the pepsin-digestion step was shorter (3 min), and fluorescent secondary antibodies, specific for STORM, were used: horse anti-rabbit Alexa-405/Alexa-647 (1:50; generous gift from B. Dudok, Institute of Experimental Medicine, Budapest, Hungary) visualizing the GABAergic axonal varicosities labeled via guinea-pig antivascular GABA transporter (vGAT, 1:6000, Synaptic System) and horse anti-guinea-pig-Cy3/A647 (1:50; generous gift from B. Dudok) marking the γ2 content of the postsynaptic elements. Finally, the sections were placed on coverslips, uncovered, and stored at 4°C. Directly before image acquisition, sections were mounted with a freshly prepared STORM-specific medium containing 80 μl of 1 x DPBS (Sigma-Aldrich), 10 μl 0.5 x mercaptoethanolamine, pH 8.5, 1 μl of anti-recombinant oxygen scavenger that consists of 10 mg glucose oxidase, 32 μl catalase, and 93 μl DPBS, and finally 20 μl of 30% glucose in distilled water. For STORM image acquisition, a similar protocol was used as described previously (Dani et al., 2010) with a 100× objective (CFI Apo 100× oil NA 1.49 TIRF WD 0.12) and imaging cycles containing one frame of activation laser illumination (405 or 561 nm) followed by three frames of imaging laser illumination (647 nm) at 30 frames/s. To ensure the z-axis stability of data acquisition, Nikon Perfect Focus System was used. Image acquisition was controlled via a STORM Andor camera (iXon 897 back-illuminated EMCCD). The STORM images comprised of 1000–3000 cycles were taken using Nikon-STORM-C2 system using Nikon NIS-Elements 4.01 STORM 2.0 software. An automated algorithm was used to subtract channel crosstalk and nonspecific activation of the A647 reporter in multicolor images.

In *vitro* biocytin-labeled cells were visualized following one of the above-described protocols. The fixative solution contained 2–4% PFA in 0.1 x phosphate buffer (PB). Tissue blocks were cut on a Vibratome (Leica) into 50 μm coronal sections. After extensive washes and incubation in blocking solution containing 10% of normal goat serum or donkey serum and 0.5% Triton-X, recorded and labeled TC cells were visualized using streptavidin-conjugated fluorescent immunoglobulin tagged with a fluorescent protein (Cy3 or Alexa-488; in 1:2000 for 2 h at room temperature) or avidin-biotin complex (Vector Laboratories; 1:300, 2 h) developed by nickel-intensified diaminobenzidine, as a chromogen. Identifying the phenotype of the recorded cells, anti-Cre recombinase fluorescent staining was performed using monoclonal mouse anti-Cre antibody (Millipore; 1:3000, overnight at room temperature) and with Cy3- (Jackson ImmunoResearch Laboratories) or Alexa-488-conjugated anti-mouse secondary antibody (Invitrogen) (1:500; 2 h at room temperature). For precise localization of the recorded cells or the silicon probe tracks, vesicular glutamate transporter type 2 (vGlut2; guinea pig or mouse anti-vGlut2, 1:3000, Millipore) or parvalbumin (rabbit anti-PV; 1:3000, Baimbridge) counterstaining was used and labeled with Cy5or A647-conjugated secondary antibodies. In control animals, the counterstaining was performed using ImmPRESS anti-mouse or rabbit IG Polymer detection kit.

In *vitro* biocytin-labeled cells were visualized following one of the above-described protocols. The fixative solution contained 2–4% PFA (150 μl per animal); 50 μm coronal sections were cut, washed, cryo-protected in 30% sucrose in 0.1 x PB overnight, and freeze thawed over liquid nitrogen. For simple fluorescent double stainings, after incubating the sections in blocking solution (10% normal goat serum for 30 min), sections were treated with monoclonal anti-Cre antibody raised in mouse or rabbit (1:30,000 and 1:12,000; Millipore and Covance, respectively; overnight at room temperature) and one of the following antibodies: mouse anti-NeuN (1:3000, Millipore Bioscience Research Reagents), mouse anti-gephyrin (1:1000, Synaptic System), anti-GABA transporter type 1 (GAT1, Millipore) and type 3 (GAT3, Millipore) overnight at room temperature. The primer antibodies were visualized with corresponding fluorescent secondary antibodies (Alexa-488-, -Cy3-, or-Cy5-conjugated IGs). Finally, sections were mounted on slides and covered with Vectashield (Vector Laboratories).
Na-pyruvate (2 mM). Slices were kept at 35°C for 30 min and then at room temperature for subsequent incubation. For recordings, slices were transferred to a submerged recording chamber held at room temperature, except for recording evoked IPSCs (eIPSCs), which was done at 34°C. The bath solution contained the following (in mM): 120 NaCl, 2.5 KCl, 1.25 NaH2PO4, 25.6 NaHCO3, 2 CaCl2, 1.2 MgCl2, 18 glucose, 1.7 L(+)-ascorbic acid, osmolarity 290 –300 mOsm, continuously bubbled with 2.5% CO2 in a humidified gas mixture of 95% O2/5% CO2, and to measure the cell's input resistance, the liquid junction potential of 10 mV was taken into account. A liquid junction potential of 20 mV was selected for the recording solution. A liquid junction potential of −10 mV was taken into account. Resting membrane potential was measured after gaining whole-cell access. Depolarizing and hyperpolarizing current square pulses were injected into the recorded cells held at different membrane potentials (−60 mV, −70 mV) through DC injections.

For sIPSC and eIPSC recordings in TC cells and to measure the cell's input resistance, the intracellular pipette solution contained the following (in mM): 120 CsCl, 10 HEPES, 2 EGTA, 8 NaCl, 0.2 MgCl2, 5 QX-314-Cl, 0.2 Na-GTP, and 10 phosphocreatine (295 mOsm, pH 7.35). QX-314-Cl and Na-GTP were added freshly daily. For sIPSCs and eIPSCs. Under these recording conditions, differences in holding currents and input resistance between AAV-GFP and AAV-Cre cells resulted entirely from altered tonic Cl− conductances. The holding current shift was measured as the difference in the holding current before and during drug application. For tonic current measurements, only male animals were used to avoid the disturbing effect of periodic changes in neurosteroid levels.

For measuring cellular discharge properties in current-clamp conditions, the intracellular pipette solution contained the following (in mM): 140 KMeSO4, 10 KCl, 10 HEPES, 0.1 EGTA, 2 MgCl2, 4 Mg-ATP, 0.2 Na-GTP, and 10 phosphocreatine (306 –309 mOsm, pH 7.23). Na-GTP was freshly added daily from concentrated stock solutions, and the solution was filtered and maintained on ice. A liquid junction potential of −10 mV was taken into account.

For eIPSCs, a monopolar stimulation electrode filled with ACSF was placed in the nRT in a position radial to the recorded TC cell and the amplitude and kinetics were measured throughout the experiment. For monitoring evoked burst-induced IPSCs (burst IPSCs), the patch pipette solutions used for sIPSCs and eIPSCs. Under these recording conditions, differences in holding currents and input resistance between AAV-GFP and AAV-Cre cells resulted entirely from altered tonic Cl− conductances. The holding current shift was measured as the difference in the holding current before and during drug application. For tonic current measurements, only male animals were used to avoid the disturbing effect of periodic changes in neurosteroid levels.

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For sIPSC and eIPSC recordings in TC cells, the GABA-AR blocker gabazine (SR 55531 hydrobromide/6-imino-3-(4-methoxyphenyl)-1(6H)-pyridazinobutanoic acid hydrobromide; 10 μM; Tocris Bioscience) was added for at least 3 min to cells recorded with the patch pipette solutions used for sIPSCs and eIPSCs. Under these recording conditions, differences in holding currents and input resistance between AAV-GFP and AAV-Cre cells resulted entirely from altered tonic Cl− conductances. The holding current shift was measured as the difference in the holding current before and during drug application. For tonic current measurements, only male animals were used to avoid the disturbing effect of periodic changes in neurosteroid levels.

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pipette was carefully withdrawn at the end of the recordings. Slices were immediately transferred to a fixative solution (4% PFA, at 4°C), fixed for 1–2 h, and intensively washed in PB. Then, slices were embedded into agar (2%) and sectioned with Vibratome.

In vitro analysis. Action potential (AP) characteristics in tonic and burst firing mode were analyzed in Clampfit version 10.2. The peak amplitudes from AP threshold, the half-widths, and the AP afterhyperpolarizations were always determined for the first AP in a burst discharge or in tonic firing trains.

Analysis of sIPSCs was performed in MiniAnalysis 6.0 (Synaptosoft). Root-mean-square noise was determined in an event-free 60 ms time window. A minimal peak threshold of at least twice the determined root-mean-square noise was set as detection criterion. When the detection parameters were applied to traces recorded in gabazine, no events could be detected. To estimate the weighted decay time constant ($\tau_w$) in control cells, monophasic events with a rise time <2 ms were averaged and $\tau_w$ calculated from a weighted average of decay time constants in a biexponential fit, thereby accounting for occasional slow spontaneous events (see Results).

For tonic currents, all-point histograms of amplitude values were generated from 20 s segments before and during drug application and fitted by single Gaussian distribution.

Off-line analysis of eIPSCs and burst IPSCs was performed using Clampfit version 10.2, and IgorPro 5. Means of 4 successive responses were used for analysis and amplitudes calculated with respect to the baseline before stimulation. Instead of 10%–90% rise time constant, time to peak was determined, as the burst IPSC is a multiphasic event. For comparison of eIPSCs and burst IPSCs, two parameters were determined. Conductances were calculated through dividing peak event amplitudes by the driving force. Decay time courses were quantified using monoexponential fits for eIPSCs in IgorPro. To determine burst IPSC decay, responses were fit with biexponential or monoexponential curves in Igor Pro. Biexponential fits were used for burst IPSCs of AAV-GFP cells to account for the involvement of both synaptic and nonsynaptic current components, whereas AAV-Cre burst IPSCs were fit with a monoexponential time constant. Fits were applied only to traces showing smooth decay without contamination by additional spontaneous synaptic events.

In vivo electrophysiology. In vivo recordings were performed 3–8 weeks after the viral injections. For LFP recordings, we used 16-channel silicon probes lowered in the primary somatosensory cortex (S1, anterior 1.2 and mediolateral 3.2 mm from bregma). Thalamic multiunit activity was monitored via 32-channel silicon probes using the same techniques as previously described ($\gamma$2 subunits; $\Delta$1-2 and $\sigma$3-5 μm diameter) for ~30 min.

In vivo analysis. All data analysis was performed in MATLAB (MathWorks). Instead of using arbitrary intraburst and interburst parameters for burst detection, a custom-built semiautomatic burst detection method was used as described previously (Slezak et al., 2011). After the identification of bursts, the following firing parameters were determined: mean firing frequency, frequency of the first APs of the bursts (burst frequency), the firing frequency for the intraburst APs (intraburst frequency), the average number of spikes in bursts (intraburst spike number), the average duration of bursts measured in milliseconds (burst length), and the number of all APs in bursts relative to the number of all spikes fired (burstiness index).

The slow oscillation frequency for the LFP recordings was calculated via wavelet spectra, Fourier spectra and autocorrelations using a custom-built MATLAB routine.

Spindle detection and analysis were performed using a custom-built MATLAB routine. Spindles were detected semiautomatically from the thalamic multiunit activity (MUA). After automatic detection, spindles were verified visually, and false detections were deleted.

Polysomnographic recordings and analysis. Surgeries for combined EEG/electromyographic (EMG) recordings and data scoring were performed on adult GABAAR$\gamma$2$^{T717}$lox mice as previously described (Franken et al., 1998). Two weeks before surgery, 4 animals were bilaterally injected with AAV-GFP (as controls) or AAV-Cre. After 10 d of recovery, the cable was plugged to the connector and mice were allowed an additional week to habituate. Undisturbed sleep–wake behavior was recorded for 48 h. The analog signals were first amplified (2000×), and the EEG and EMG signals were high-pass filtered at 0.7 Hz and 10 Hz, respectively. Signals were digitized at 2 kHz and down-sampled to 200 Hz through Somnologica 3.1.3 software (Embla System). Data were visually scored with a resolution of 4 s to determine the vigilance stage (wake, NREM sleep, REM sleep). The NREM sleep power spectrum was generated from NREM sleep during both light and dark periods, but NREM sleep epochs were included only if they were both preceded and followed by other NREM sleep epochs (Wimmer et al., 2012). For analysis on spectral characteristics of the NREM-to-REM sleep transitions, only the two light phases were taken into account, as these contain the majority of transitions. Data analysis was performed as described previously (Astorì et al., 2011; Wimmer et al., 2012).

Statistics. Statistical significance was assessed using Student’s t test or Mann–Whitney U test based on the normality level of the dataset, and a p < 0.05 was considered statistically significant. For quantitative receptor localization, after normalization for area of ROIs and averaging, samples from Cre- and GFP-expressing animals were compared with Student’s t test using Mystat (Systat Software). For analysis of EEG power spectra, univariate repeated-measures ANOVA with factor “group” and “bin” was used. In both text and figures, data are presented as mean ± SD or as whisker plots.

Results
Viral intervention successfully eliminates synaptic $\gamma$2 subunits
To selectively and focally delete the $\gamma$2 subunits from TC cells, we unilaterally injected Cre-recombinase-containing or, as control, GFP-containing, adeno-associated viral vectors (AAV-Cre or AAV-GFP, respectively) into the somatosensory thalamus of GABAAR$\gamma$2$^{T717}$lox mice (Fig. 1A, B). After 2–3 weeks, we com-
pared ipsilateral and contralateral expression of γ2 subunits with confocal microscopy. In noninfected as well as the AAV-GFP-infected thalami, γ2 immunostaining resulted in dense punctate labeling, reflecting the predominantly synaptic localization of this subunit (Fig. 1C) (Nusser et al., 1998). In contrast, within the AAV-Cre-infected region, punctate γ2 immunoreactivity was not apparent (Fig. 1D). Next, we examined the expression of α1 subunits, which coassemble with γ2 to form functional receptors (Tretter et al., 1997) but, in addition, are also present extrasynaptically. Compared with controls, α1 labeling was reduced in the AAV-Cre-infected regions, but a weak staining, probably reflecting the extrasynaptic pool, persisted (Fig. 1E,F). Quantitative receptor localization demonstrated a nearly complete loss of γ2 immunoreactivity (relative intensity, 6.68 ± 2.11%) and a less dramatic but highly significant decrease in α1 immunostaining (relative intensity, 18.27 ± 20.18%) compared with AAV-GFP-infected regions (Fig. 1F). In contrast, in the case of the extrasynaptic, α4-containing GABA-ARs, which are responsible for tonic inhibition (Porcelli et al., 2003; Chandra et al., 2006), no significant difference was observed between the AAV-Cre- and AAV-GFP-infected thalamic regions (relative intensity, 68.24 ± 50.82%; Fig. 1G–I). Based on these anatomical findings, we conclude that deletion of γ2 and the associated α1 protein was successful and selective in the infected regions.

To analyze the changes of γ2 staining at higher spatial resolution, STORM was applied (Dani et al., 2010). In these experiments, we used vesicular GABA transporter (vGAT) labeling to visualize GABAergic terminals. On the contralateral side, vGAT immunostaining labeled vesicle aggregations in the size range of nRT terminals (1–2 μm). Adjacent to these terminals, vGAT immunostaining formed dense clusters of punctate (Fig. 1J–L). The size of these clusters (400–500 nm) was in the range of synapses established by nRT terminals measured in serial electron microscopic images (Wanaverbeq et al., 2008). Single vGAT-positive terminals contacted one or up to four receptor clusters. In case of multiple contacts, the synapses were on different sides of the terminals, similar to earlier results (Wanaverbeq et al., 2008) (Fig. 1J). In contrast to the contralateral side, in the ipsilateral, AAV-Cre-infected region, γ2-labeled receptor clusters were absent around the vGAT-positive boutons (Fig. 1K–M). These data confirm successful viral infection and the elimination of synaptic GABA-ARs.

Figure 3. Deletion of synaptic γ2 subunits reduces tonic GABAergic inhibition. A. Outward shift in the holding current of a TC cell held at −50 mV before and during gabazine (10 μM, gray shadow) application in a control (top, black) and an AAV-Cre (bottom, red) VB neuron. Gaussian curves indicate a fit to the all-points histograms of current traces before (left) and after (right) gabazine. Dotted horizontal lines indicate mean holding current level before gabazine. The large spontaneous inhibitory events recorded from the AAV-Cre VB cell are discussed in Figure 4. B. Bar graph of gabazine-induced increase in the holding current (ΔI) (control, Ctrl, n = 7; AAV-Cre, Cre, n = 9). Bars indicate mean ± SD. *p < 0.05 (Student’s t test). C. Box plot of cellular input resistance (R) of control (n = 12) and AAV-Cre VB cells (n = 27). The midline indicates the median. Bottom and top of the boxes represent the first and third quartiles, respectively. Whiskers represent full range. *p < 0.05 (Student’s t test).

Removal of γ2 subunits eliminates fast IPSCs

To test the functional consequences of deletion of synaptic γ2-GABA-ARs, we performed whole-cell recordings from VB thalamic neurons in acute horizontal brain slices with intact nRT-TC connections forming the sole inhibitory input. When patched with pipettes containing a K+-based pipette solution (Fig. 2A), these cells showed normal AP discharge properties both in tonic (AP amplitudes 63.1 ± 11.4 mV and 60.9 ± 6.1 mV; half-widths 1.04 ± 0.14 ms and 1.00 ± 0.18 ms; AP afterhyperpolarization −21.4 ± 5.9 mV and −21.3 ± 4.2 mV, p > 0.05 for all values; AAV-GFP, n = 5; AAV-Cre, n = 9) and burst mode (AP amplitudes 68.2 ± 9.8 mV and 63.8 ± 8.1 mV; half-widths 0.83 ± 0.15 ms and 0.84 ± 0.23 ms; AP afterhyperpolarization −13.4 ± 6.5 mV and −12.9 ± 3.0 mV; AP number/burst 1.7 ± 0.4 and 1.6 ± 0.4, p > 0.05 for all values; control, n = 7; AAV-Cre, n = 8). Moreover, cellular resting membrane potentials were unaltered (−76.6 ± 9.6 mV vs −75.0 ± 11.4 mV, p > 0.05; control, n = 22; AAV-Cre, n = 21 cells).

In the presence of glutamate receptor antagonists (DNQX 40 μM, D-APV 100 μM), all VB cells recorded in the control slices displayed fast spontaneous IPSCs (sIPSCs; rise time < 2 ms, weighted decay time constant < 11 ms) at a frequency of 1.33 ± 0.89 Hz that were blocked by the GABA-AR antagonist gabazine (Fig. 2B). In contrast, fast rising sIPSCs were almost completely abolished from VB cells recorded in the AAV-Cre-infected regions (0.006 ± 0.018 Hz, p < 0.001; Fig. 2A–C). In addition to fast IPSCs, slower (rise time > 30 ms) IPSCs were occasionally detected in both groups (see below).

We next examined eIPSCs elicited via activation of GABAergic inputs by extracellular stimulation in the nRT. Fast eIPSCs (rise time < 1 ms, decay time < 4 ms) were observed in 4 of 5 control cells tested with a mean amplitude of −82 ± 27 pA and 0.19 ± 0.15 pC of charge transfer. In contrast, in the AAV-Cre-infected regions, eIPSCs were absent in 5 of 6 TC cells recorded after both single and 10 Hz stimulation (Fig. 2D,E). The one AAV-Cre TC cell that responded to the stimulation showed slow kinetics (rise time 5 ms; decay time constant 28 ms) and large amplitude (~120 pA). In summary, these data demonstrate the absence of fast synaptic inhibition mediated by γ2-GABA-ARs in AAV-Cre-infected VB cells.

To determine the effect of deleting synaptic receptors on tonic GABA-AR-mediated currents, we applied gabazine (10 μM) to cells patched with Cl−-based intracellular solutions. In AAV-GFP-infected cells, gabazine induced an outward shift in the holding current (ΔI = 53 ± 19 pA), reflecting the presence of a tonic current (Fig. 3A). In AAV-Cre VB cells, ΔI was smaller (30 ± 14 pA, p < 0.05; Fig. 3A,B) and cellular input resistance was higher compared with controls (255 ± 110 mΩ vs 177 ± 84 mΩ, p < 0.05; Fig. 3C). These results suggest that either γ2-GABA-ARs account for a portion of the tonic current or existing extrasynaptic receptors have reduced functionality. More importantly, the fact that tonic currents are reduced in γ2-lacking VB cells indicates that a compensatory upregulation of nonsynaptic receptors did not occur.

Burst-mediated IPSCs persist in the absence of synaptic GABA-ARs

Surprisingly, despite the absence of fast sIPSCs, we observed large and slow spontaneous inhibitory events in slices prepared from both control and AAV-Cre TC cells (Fig. 4A). These typically occurred at low frequencies (1.9 ± 3.9/min vs 1.7 ± 2.0/min, p > 0.05; control, n = 16; AAV-Cre, n = 13), singly or in groups of 2–4 events with interburst intervals of hundreds of milliseconds,
and are caused by spontaneous repetitive nRT burst discharge (Wimmer et al., 2012). In control cells, these events showed the typical waveform of burst IPSCs consisting of multicomponent fast events riding on a slower current envelope. In contrast, in AAV-Cre TC cells, burst IPSCs consisted of only the slow current envelope and lacked the fast phasic inhibitory currents (Fig. 4B), showing distinctly slower rising phases.

To quantitatively compare the burst IPSC properties in the two cell groups, we electrically stimulated the nRT to produce evoked burst IPSCs (Fig. 4C,D). Similar to spontaneous burst IPSCs, evoked burst IPSCs were abolished by gabazine (Fig. 4C). Compared with single AP-evoked IPSCs, the evoked burst IPSCs in control cells had a ~9.5-fold greater conductance (15.3 ± 10.0 nS for burst IPSCs, n = 13, vs 1.6 ± 0.60 nS for eIPSCs, n = 4) and slower decay kinetics (biexponential decay time constants (τ1, τ2) of 19 ± 10 ms and 90 ± 44 ms in n = 8 cells for burst IPSCs, Fig. 4D, third panel, vs monoexponential decay time constant (τ) 3.0 ± 1.7 ms for eIPSCs, n = 4).

Similar to spontaneous burst IPSCs, evoked burst IPSCs were crowded by fast IPSCs in control, but not in AAV-Cre VB cells (Fig. 4C). Moreover, the average amplitude of the evoked burst IPSCs in AAV-Cre TC cells was diminished by ~60% compared with control (control, 425 ± 291 pA, n = 13; AAV-Cre, 161 ± 93 pA, n = 20; p < 0.01; Fig. 4D) and burst IPSCs generated in AAV-Cre cells showed decelerated time-to-peak (33.1 ± 10.0 ms in control vs 55.1 ± 25.9 ms in AAV-Cre TC cells, p < 0.01, Fig. 4D). To compare decay time courses, the slow time constant of AAV-GFP burst IPSCs was compared with a monoexponential fit of the AAV-Cre burst IPSCs, thereby taking into account the lack of a dominant fast current component resulting from the elimination of synaptic receptors. The monoexponential decay time constant of the AAV-Cre burst IPSC was comparable with the slow current component of control cells (AAV-Cre, τ = 69 ± 23 ms, n = 15, vs control, τ = 90 ± 44 ms, n = 8; p > 0.05; Fig. 4D). The total charge transfer, calculated as the integral of the current trace, was diminished by 48% (control, 32.8 ± 26.5 pC, n = 13; AAV-Cre, 15.7 ± 8.3 pC, n = 20; p < 0.05; Fig. 4D). Moreover, burst IPSCs from both control and AAV-Cre TC cells had a reversal potential corresponding to the Cl− gradient imposed by the patch pipette. These data indicate that, in the absence of synaptic GABA-ARs, significant GABA-AR-mediated, slow, phasic events persist when the presynaptic neurons fire in burst mode. These events are independent of synaptic GABA-ARs and involve phasic recruitment of nonsynaptic GABA-ARs.

The efficiency of burst IPSCs in AAV-Cre VB cells to generate rebound low-threshold spikes could be decreased because of the missing synaptic component and the slowed-down rising phase. To test this, in current-clamped noninfected VB neurons, representative spontaneous burst IPSCs from control or AAV-Cre cells were injected as a command current (Fig. 4E). Injected waveforms were scaled in their amplitude (maximal peak = −420 pA) to mimic a variety of endogenous inputs of different efficacy. Cells injected with current commands obtained from control or AAV-Cre cells displayed a comparable input–output curve when considering the latency to the first AP in the rebound burst, the amplitude of the low-threshold Ca2+-spike, and similar numbers of APs (H) to control protocol (black). F, G, Insets, Representative response with dashed lines indicating the analyzed response property. H, Red and black dashed lines indicate the number of APs corresponding to the mean charge transfer of burst IPSCs across all analyzed cells (seen in D). Data are mean ± SD. *p < 0.05 (Student’s t-test). **p < 0.01 (Student’s t-test).

Deletion of synaptic GABA-ARs does not produce aberrant oscillation

Our in vitro data indicate that, after the removal of γ2/GABA-ARs, a sizable portion of the slow burst IPSCs remained intact. To test the role of this slow nonsynaptic current in controlling network rhythms, we used in vivo assays.

We first iontophoretically injected the selective GABA-AAR blocker gabazine (1 mM) into the somatosensory thalamus of control mice (Fig. 5A, B) to block both synaptic and extrasynaptic receptors. In accordance with previous data (Castro-Alamancos, 1999), gabazine application into the thalamus of mice (n = 3) disrupted the normal slow cortical oscillation and induced aberrant, large-amplitude regular activity in the cortex with a power spectrum peaking at 2.6 Hz (Fig. 5A, B). This indicates that, when all thalamic GABA-ARs are focally blocked, normal TC oscillations are disrupted.
Figure 5. Removal of γ2-GABA-ARs in the thalamus does not cause aberrant rhythmic cortical activity. A, Cortical (primary somatosensory cortex, S1) LFP recordings before (control, black) and after unilateral administration of gabazine (blue) into somatosensory thalamus under ketamine/xylazine anesthesia. There are epileptiform bouts of activity (curved arrows) in the cortical LFP recordings after gabazine. B, Scheme of the unilateral, thalamic (Thal), gabazine (GBZ) injection site. C, Low-power fluorescent image depicting Cre immunostaining (red) after thalamic injection of AAV-Cre. Scale bar, 500 μm. D, The cortical LFP recorded from the ipsilateral S1 (red) after AAV-Cre injection to the thalamus is comparable with control (A). E, Power spectra (left) and autocorrelations (right) of the cortical LFPs in control condition (black), after gabazine (blue) and AAV-Cre injection (red). There is a large central peak at 2.6 Hz (marked by arrow) and the side lobes (arrowheads) in the GBZ treated animal but not in Ctrl and Cre cases. F, In freely moving animals, slow oscillations are comparable in control (top black) and AAV-Cre-injected animals (top red) during NREM sleep. Bottom, Black and red traces represent the simultaneous EMG activity. G, Scheme of the bilateral thalamic injection of AAV-Cre in the freely moving animals. H, Power spectrum of NREM sleep over 24 h from 4 animals each of the AAV-GFP and AAV-Cre groups. Data were binned into 1 Hz bins and analyzed statistically using univariate repeated-measures ANOVA with factor “group” and “bin,” yielding $p > 0.05$ for the group $\times$ bin interaction.

Next, we performed similar experiments using focal delivery of AAV-Cre (Fig. 5C) vectors instead of gabazine injection to examine whether deletion of synaptic receptors alone results in disturbed cortical activity. However, 3–8 weeks after viral delivery, selective deletion of only the synaptic GABA-ARs did not result in aberrant oscillations (Fig. 5D; $n = 9$). As a consequence, none of the cortical LFP autocorrelograms recorded from AAV-Cre-treated animals displayed the large central peaks, flanked by side lobes, which characterize the large amplitude, regular, rhythmic activity we observed after the injection of gabazine (Fig. 5E).

We also investigated the consequence of γ2-GABA-AR deletion on oscillatory activity during natural sleep. After bilateral thalamic injection of AAV-Cre ($n = 4$) or AAV-GFP (as control, $n = 4$), polysomnographic recordings yielded no sign of overt large-amplitude, aberrant activity during NREM sleep (Fig. 5F,G) after 3 weeks of recovery in freely moving mice. Mean power spectra across 24 h of NREM sleep were indistinguishable for AAV-GFP and AAV-Cre animals (repeated-measures ANOVA, $p > 0.05$, Fig. 5H). Thus, in contrast to the pharmacological blockade of all GABA-ARs, the TC system produces normal oscillatory activity even when largely devoid of thalamic γ2-GABA-ARs.

Altered firing properties of TC cells in the absence of synaptic GABA-ARs

Complementing the EEG and LFP recordings, we also examined the firing pattern of individual TC cells in ketamine/xylazine-anesthetized mice through juxtacellular recordings (20–30 min) in two somatosensory thalamic nuclei known to have different inhibitory inputs (Pinault, 2004), the VB and the posterior nucleus (Po) (Barthó et al., 2002; Bokor et al., 2005; Wanaverbecq et al., 2008) (Fig. 6). Nuclear localizations of the juxtacellularly neurobiotin-labeled cells were post hoc identified using immunostaining against vGluT2 (Fig. 6D,I), whereas their phenotype was determined with Cre immunostaining (Fig. 6E,F).

The firing patterns of control, AAV-Cre-negative VB ($n = 11$) and Po ($n = 7$) cells in mice were similar to those of rats under the same recording conditions (Slézia et al., 2011). More than 90% of spikes occurred in bursts. In accordance with our previous rat data, the average intraburst frequency of spikes in VB cells was significantly faster than in Po cells (264.6 ± 47.2 Hz vs 204.3 ± 29.4 Hz, $p < 0.05$; Fig. 6 A,F).

Similar to controls, all AAV-Cre VB ($n = 9$) and Po ($n = 8$) cells were able to fire bursts of APs coupled to the rhythmic LFP activity (Fig. 6 B,G). However, the properties of bursts were significantly altered in AAV-Cre cells and the alterations displayed nucleus-specific patterns (Fig. 6C,H).

In the absence of synaptic inhibition, the mean firing rate significantly decreased in case of AAV-Cre VB cells (control, 3 ± 1.3 Hz; AAV-Cre, 1.6 ± 1.1 Hz, $p < 0.05$; Fig. 6C). This was largely because of the reduced number of burst events (burst incidence, 0.96 ± 0.4 Hz in control vs 0.52 ± 0.36 burst/s in AAV-Cre, $p < 0.05$). In addition, the intraburst frequencies of spikes in AAV-Cre VB cells were lowered by 42 Hz (16%) (control, 264.6 ± 47.2; AAV-Cre, 222.5 ± 20.5 Hz, $p < 0.05$); however, the intraburst spike number did not change. As a consequence, the length of the AAV-Cre VB bursts increased significantly (burst length; 7.2 ± 1 ms in control vs 8.7 ± 1.7 ms, in AAV-Cre, $p < 0.05$; Fig. 6C).

In contrast to AAV-Cre VB cells, the mean firing rate of AAV-Cre Po cells was not altered (Fig. 6H). In these cells, the intraburst spike number displayed a slight decrease (3.1 ± 0.4 spike/burst in control vs 2.7 ± 0.5 spike/burst in AAV-Cre Po cells, $p < 0.05$; Fig. 6H). Because the intraburst frequency did not change, this resulted in a decrease of the overall burst length (control, 10.7 ± 1.6 ms; AAV-Cre, 9.0 ± 1.9 ms, p < 0.05; Fig. 6H).

These data show that TC cells retain low threshold bursts of APs in the absence of synaptic GABA-ARs, although the lack of synaptic inhibition slowed down the affected neurons, especially in VB.

Spindle oscillations persist in the absence of synaptic GABA-ARs

Next, we examined the effect of synaptic GABA-AR removal on spindle oscillations in the thalamus of urethane-anesthetized mice through multichannel silicone probes. The probes were labeled with Dil, and their depths were registered to determine the exact position of the electrode shanks relative to the infected regions (Fig. 7A,D). For these experiments, we chose urethane anesthesia because it provides a network activity generating spindles similar to natural sleep (Steriade et al., 1993) unlike ket-
amine/xylazine anesthesia, during which short spindles are more prevalent (Slézia et al., 2011).

Spindles were detected in VB as transient rhythmic elevations of thalamic MUA, in the 7–15 Hz range, and could be quantitatively characterized (frequency, length, number of cycles, and occurrence; Fig. 7). We analyzed spindles in four experimental groups: noninjected wild-type, noninjected GABAARγ2^77Ilox mice; AAV-Cre injected, contralateral to the thalamus injection (AAV-Cre, contralateral, Cre; Fig. 7A–C) and AAV-Cre injected, ipsilateral, within the injected zone (AAV-Cre, ipsilateral, Ctrl; Fig. 7D,E). The two noninjected groups did not differ in their quantitative measures and thus were grouped together (NonI, n = 7). The occurrence of spindles was variable during the recordings because of the spontaneous fluctuation in the depth of anesthesia. To overcome this variability, we performed long recordings (129 ± 60 min/animal), during which we detected together 109,762 spindles (on average 5777 ± 5509 spindle/animal, 294 ± 191 spindle/channel).

Spindles in the noninjected groups occurred at 3.36 ± 2.16 event/min (Fig. 7F). On average, the spindles consisted of 6.84 ± 0.78 cycles. The spindles lasted for 654 ± 74 ms and the average intraspindle frequency was 10.46 ± 0.35 Hz, which is consistent with earlier data (Steriade et al., 1993).

In all experiments of the Cre group (8 penetrations in 6 animals), normal spindle activity was detected (Fig. 7D,E) in the AAV-Cre-injected zone. The quantitative analysis demonstrated that the spindles had similar rates of occurrence (3.02 ± 0.5 event/min), similar numbers of spindle cycles (6.30 ± 0.39 ms), and lengths (659 ± 52 ms) to control. The only consistent change due to the deletion of synaptic inhibition was a slight decrease (10.5 ± 0.5 Hz in Ctrl vs 9.6 ± 0.4 Hz in Cre, p < 0.01) in intraspindle frequency (Fig. 7F).

We also followed on naturally occurring spindle activity through EEG recordings of NREM sleep in drug-free freely moving animals that previously received bilateral injections of either AAV-Cre or AAV-GFP (as control) vectors. NREM sleep stages were analyzed with respect to the time course of power spectral densities at transitions to REM sleep. At these transitions, the power of frequencies between 5 and 25 Hz increases rapidly, notably at spindle-related EEG frequencies (Astori et al., 2011) (Fig. 8A). No change in the time course and power of this surge, averaged here for frequencies between 7 and 15 Hz, was found in the AAV-Cre animals (Fig. 8B). These data demonstrate that oscillations in the spindle frequency range are preserved after the removal of synaptic inhibition from the thalamus.

**Discussion**

We show here that oscillatory activity in thalamic circuits is under the control of inhibition that involves γ2 subunit-independent GABA-ARs. These receptors are activated as nRt cells switch from tonic to burst discharge and suggest a “firing pattern-specific” form of inhibitory communication. Contrary to long-standing, previous views, conventional synaptic GABA-ARs are not absolutely required for TC rhythms. Phasic, nonsynaptic inhibition can efficiently drive TC cell rebound discharge and entrain slow oscillations and spindle waves during both sleep and anesthesia.

Our data demonstrate that, during burst firing of nRt cells, IPSCs in TC cells have both synaptic and nonsynaptic components in approximately equal proportions because 50% of the charge transfer remained intact after complete removal of the synaptic receptor pool in case of burst IPSCs. The remaining current had slow rise times and lacked fast-rising individual IPSCs, suggesting the contribution of extrasynaptic receptors. Most recently, an involvement of δ4-containing GABA-ARs has been recognized during burst discharge at nRt-TC synapses in a manner that required GABA spillover, confirming our conclusions (Herd et al., 2013).

In the case of burst IPSCs, extrasynaptic receptors were recruited in a phasic, not in a tonic, manner. AP-mediated activation of extrasynaptic receptors, termed “GABA-A-slow” events, has already been demonstrated in several brain regions (Rossi and Hamann, 1998; Szabadi et al., 2007; Mánko et al., 2012). However, studying the extrasynaptic component of GABA-A-slow in isolation has not been possible with conventional pharmacological or genetic knock-out methods. Using virus-mediated gene knock-out of synaptic receptors, here we could quantitatively characterize the extrasynaptic component of GABA-AR responses and demonstrate that phasic extrasynaptic inhibition is mediated by burst discharge and powerful enough to drive global brain rhythms. Our data disclosed a striking difference between eIPSCs when the presynaptic neuron fired in tonic as opposed to the burst
mode. The complete disappearance of sIPSCs and eIPSCs after the removal of synaptic receptors in TC cells indicates that inhibition caused by single or tonic APs was mediated entirely by synaptic receptors. In particular, up to 10 Hz, tonic discharge did not recruit slow persistent currents (Wanaverbecq et al., 2008, Herd et al., 2013). In sharp contrast, during burst firing mode, a significant extrasynaptic receptor pool was additionally recruited that resulted in a strong increase of peak Cl⁻ conductance (up to ~10-fold). According to Herd et al. (2013), extrasynaptic receptors contribute not only to burst IPSCs, but also prolong the decay time course of IPSCs after single APs. Small differences in the actual role of α4-GABA-ARs could vary according to the preparation used. Herd et al. (2013) studied a complete α4 KO, whereas in this study virally mediated gene knock-out was used in adult animals. Full KO techniques are more prone to induce compensatory changes, such as alterations in properties and exact location of remaining receptors. Intriguingly, however, this study mirrors our finding by showing that reduction of only extrasynaptic, but not nonconventional GABA-ARs, also produces a 50% reduction in charge transfer.

Phasic, extrasynaptic inhibition was sufficient to promote rebound burst firing in TC cells, as demonstrated through command waveforms derived from burst IPSCs. This conclusion was also supported by the in vivo data showing burst firing in all TC neurons lacking synaptic GABA-ARs and preserved TC oscillations.

The burst IPSCs did not result from a temporal summation of phasic, conventional GABA-AR-mediated fast IPSCs. Instead, burst IPSCs reflect a rapid accumulation of GABA release that evokes phasic, nonsynaptic receptor activation in a manner not reached via repetitive tonic discharge up to tens of Hz (Wanaverbecq et al., 2008). As nRT-TC synapses are depressing, specific presynaptic release mechanisms activated during high-frequency discharge could contribute to the rapid increases in synaptic GABA levels, which remain the subject of further molecular characterization of nRT-TC synapses.

An unexpected change in our studies was a pronounced decrease in tonic inhibition and the associated increase in membrane resistance, without an apparent change in the expression of α4-subunits. One explanation for the decreased tonic conductance is the disappearance of phasic inhibition in the distal dendritic region of TC cells, which is indistinguishable from the tonic conductance at the somatic level (Farrant and Nusser, 2005). Other possibilities, such as homeostatic downregulation of ambient GABA levels or involvement of nonconventional GABA-AR pools, remain to be determined. Based on our data, we currently cannot exclude the contribution of synaptic receptors to tonic inhibition.

Our experimental approach provided a unique opportunity to study the firing pattern of individual neurons lacking synaptic receptors embedded into a neuronal network oscillating normally. We performed the experiments in two thalamic nuclei, VB and Po, known to have distinct excitatory and inhibitory connectivity (Barthó et al., 2002; Bokor et al., 2005; Wanaverbecq et al., 2008; Groh et al., 2013). We found significant internuclear differences in response to the removal of synaptic GABAergic inhibition. Major changes were found in the VB, including a
significant drop in firing frequency, whereas changes in the Po were less prevalent. We attribute these differences to the distinct connectivity and activity pattern of the two nuclei (Hoogland et al., 1991; Groh et al., 2013). In case of VB cells, during the Up states of ketamine/xylazine induced slow oscillation, the intracellular activity is dominated by nRT-mediated rhythmic inhibition, whereas in case of Po cells fast rising EPSPs arising from layer 5 pyramidal cells determine Po activity under similar conditions (Groh et al., 2013). In agreement with these data, removal of synaptic inhibition affected VB cells to a much greater extent than Po cells. Thus, we conclude that the exact role of synaptic inhibition in controlling firing activity depends on the neuronal network the cell is embedded.

Thalamic application of GABA-AR antagonists, which block both synaptic and extrasynaptic receptors, disrupts the major TC oscillation and evokes aberrant, large-amplitude activity in ferrets and rats (Bal et al., 1995a; Castro-Alamancos, 1999), an observation that we replicated now in mice. These studies demonstrate that GABA-ARs in the thalamus have a critical role in supporting TC oscillations and GABA-B receptors alone cannot maintain normal rhythmic activity. In this study, we compared the consequences of full pharmacological blockade of GABA-ARs with those induced by virally mediated knock-out of synaptic GABA-ARs. Whereas total pharmacological blockade of thalamic GABA-ARs disrupts the major TC oscillations (Kim et al., 1995; Castro-Alamancos, 1999), the selective virus-mediated deletion of γ2-GABA-ARs did not affect the normal brain rhythms, and removal of only the synaptic GABA-ARs did not produce aberrant cortical activity. These experiments were performed under two different kinds of anesthesia (ketamine/xylazine and urethane) and in drug-free conditions as well, with identical outcomes, indicating that particular experimental conditions did not affect the results. Our data thus show that activation of extrasynaptic receptors by GABA spillover during nRT bursts alone generates sufficient amount of inhibition to maintain normal slow cortical oscillations and prevent the development of aberrant rhythmic activity.

A surprising outcome of our experiments was that spindle activity and EEG hallmarks of sleep spindles were also preserved in the brain of AAV-Cre-infected mice, indicating that relatively fast oscillations (7–15 Hz) can persist in the absence of synaptic receptors. The data demonstrate that slow GABA-A currents alone are able to maintain normal spindles with statistically similar occurrence and duration as in the intact brain. Despite the reduced charge transfer of burst IPSCs, after the removal of synaptic receptors, the phasic, extrasynaptic component of the burst IPSCs provided enough hyperpolarization to allow rebound burst discharges in TC cells. Indeed, in TC cells, a relatively small proportion of T-type Ca2+ channels’ deactivation is sufficient for rebound burst generation (Dreyfus et al., 2010). As the frequency of rebound burst depends on the integrated membrane hyperpolarization imposed by nRT-IPSPs (Bal et al., 1995b), we found a slightly but significantly slower mean spindle frequency (~1 Hz decrease). In our experiments, the time course of thalamic GABA-A slow/burst IPSCs was 100–120 ms, which is well suited to pace a rhythm in the spindle frequency range (7–15 Hz) via the rebound mechanism. Thus, we propose here that this peculiar form of inhibition displaying dual synaptic and extrasynaptic nature was evolved to entrain a stable oscillation in the 7–15 Hz frequency range.

In conclusion, in the present study, we identified a discharge mode-specific mechanism of inhibition at a long-studied synapse implicated in a classical pacemaker activity. The synapse’s main function is to synchronize TC cells to recruit them into oscillatory activity. Our data imply a complex organization of the synapse and a fine-tuned cooperation between different GABA-ARs according to the discharge mode. The molecular constitution of thalamic GABA-ARs is central to our understanding of TC synchronization during sleep, as synaptic GABA-ARs are the targets of many hypnotic and sedative drugs (Winsky-Sommerer, 2009). Here we propose that a nonsynaptic receptor entrains major sleep-associated TC oscillations in a phasic manner. Our findings provide a compelling explanation for the specific roles of different GABA-ARs in distinct behavioral states and thus open novel avenues to the development of drugs that act specifically on receptor subtypes contributing to burst IPSCs. These would have the unique potential to boost sleep yet protect phasic inhibition mediated by tonic discharge and associated wake-related functions.

References


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