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Variable Action Potential Backpropagation during Tonic Firing and Low-Threshold Spike Bursts in Thalamocortical But Not Thalamic Reticular Nucleus Neurons

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Backpropagating action potentials (bAPs) are indispensable in dendritic signaling. Conflicting Ca2⁺-imaging data and an absence of dendritic recording data means that the extent of backpropagation in thalamocortical (TC) and thalamic reticular nucleus (TRN) neurons remains unknown. Because TRN neurons signal electrically through dendrodendritic gap junctions and possibly via chemical dendritic GABAergic synapses, as well as classical axonal GABA release, this lack of knowledge is problematic. To address this issue, we made two-photon targeted patch-clamp recordings from rat TC and TRN neuron dendrites to measure bAPs directly. These recordings reveal that "tonic" and low-threshold-spike (LTS) "burst" APs in both cell types are always recorded first at the soma before backpropagating into the dendrites while undergoing substantial distance-dependent dendritic amplitude attenuation. In TC neurons, bAP attenuation strength varies according to firing mode. During LTS bursts, somatic AP half-width increases progressively with increasing spike number, allowing late-burst spikes to propagate more efficiently into the dendritic tree compared with spikes occurring at burst onset. Tonic spikes have similar somatic half-widths to late burst spikes and undergo similar dendritic attenuation. In contrast, in TRN neurons, AP properties are unchanged between LTS bursts and tonic firing and, as a result, distance-dependent dendritic attenuation remains consistent across different firing modes. Therefore, unlike LTS-associated global electrical and calcium signals, the spatial influence of bAP signaling in TC and TRN neurons is more restricted, with potentially important behavioral-state-dependent consequences for synaptic integration and plasticity in thalamic neurons.

Key words: action potential; backpropagation; dendrite; low-threshold spike; thalamic reticular nucleus; thalamocortical

Introduction

Dendritic backpropagating action potentials (bAPs) have critical roles including induction of spike-timing-dependent synaptic plasticity, dendritic Ca2⁺ spike generation, and triggering dendritic neurotransmitter release (Larkum et al., 1999; Kampa et al., 2006, 2007; Acuña-Goycolea et al., 2008). In thalamocortical (TC) and thalamic reticular nucleus (TRN) neurons, despite im-
aging and computational investigations, the site of AP origin and the true extent of AP backpropagation throughout the dendritic tree remains unknown. In TC neurons, Ca\(^{2+}\)-imaging studies (Crandall et al., 2010; Errington et al., 2010, 2012; Sieber et al., 2013) and a solitary dendritic recording study (limited for technical reasons to only ~60 μm from the soma; Williams and Stuart, 2000) suggest that bAPs are strongly attenuated, with even spike trains unable to evoke significant distal dendritic Ca\(^{2+}\) influx. In TRN neurons, there is conflicting data, with some studies suggesting bAPs produce distance-dependent dendritic calcium transients (Δ(Ca\(^{2+}\))) similar to those in TC neurons (Gueni et al., 2008; Crandall et al., 2010) and others suggesting that bAPs can produce significant Δ(Ca\(^{2+}\)) throughout the dendritic tree (Chauvon et al., 2013). However, unlike TC neurons, no data from direct patch-clamp recordings on AP backpropagation in TRN dendrites are currently available. This is problematic because notable discrepancies have been reported previously between optical and electrical recording studies. For example, in cortical pyramidal neurons, very different estimates of AP backpropagation into basal dendrites were obtained using patch-clamp recordings (Nevian et al., 2007) and voltage-sensitive dye imaging (Antic, 2003). Although the underlying reasons are unclear, temporal undersampling might confound data from imaging experiments. Conversely, the main problem associated with electrical recording in thin dendrites, series resistance, can be adequately negated even for high-resolution recording electrodes (Nevian et al., 2007; Krueppel et al., 2011; Connelly et al., 2015). Furthermore, whereas Ca\(^{2+}\) imaging can report evoked Δ(Ca\(^{2+}\)), these might not capture the full effects of bAP propagation because they also rely on dendritic Ca\(^{2+}\) channel distribution. For example, in the absence of distal dendritic Ca\(^{2+}\) channels, a bAP may not, on its own, induce Ca\(^{2+}\) entry but may still provide sufficient distal depolarization to relieve magnesium-dependent NMDA receptor block if appropriately timed relative to an EPSP. Therefore, the full impact of AP backpropagation in thalamic neurons can only be understood by investigating the voltage transients that APs produce throughout the dendritic tree.

From a physiological perspective, understanding the extent of backpropagation in TRN neurons is important because they signal, not only via classical axonal synapses, but also through dendritic electrical synapses mediated by Cx36-dependent gap junctions (Landisman et al., 2002) and, on the basis on structural evidence, GABAergic dendrodendritic synapses (Deschenes et al., 1985; Pinault et al., 1997). In other cell types, including interneurons of the dorsal LGN (Acuna-Goycolea et al., 2008), bAPs are effective in evoking dendritic neurotransmitter release; therefore, understanding backpropagation in TRN neurons will shed further light on the spatial extent of electrical and chemical signaling in TRN dendrites. Moreover, in both TC and TRN neurons, cortical and subcortical synaptic inputs have differential distribution patterns across the dendritic tree, with the former targeting more distal dendritic locations. Therefore, understanding the true extent of AP backpropagation in thalamic neurons is necessary to predict how bAPs might interact with specific synaptic inputs. Using two-photon targeted patch-clamp recordings from dendrites of TC and TRN neurons, we have for the first time measured AP backpropagation directly both during “tonic” and low-threshold-spike (LTS) “burst” firing modes. We find that APs are of axosomatic origin and undergo substantial voltage attenuation as they propagate into the dendrites of both cell types. Moreover, we find differences in AP propagation within LTS bursts and between tonic and burst firing in TC neurons that are absent in TRN cells. Therefore, whereas TC and TRN neurons share some dendritic properties (Connelly et al., 2015), there are key differences in AP backpropagation between these cells.

**Materials and Methods**

**Electrophysiology.** Coronal slices (300 μm) containing the dLGN and horizontal slices (250 μm) containing the TRN were prepared from postnatal day 20–25 (dLGN) and 17–21 (TRN) Wistar rats of either sex, deeply anesthetized using isoflurane, as described in Errington et al. (2010) with approval of the Cardiff University Research Ethics Committee and in accordance with the Home Office Animals (Scientific Procedures) Act 1986, United Kingdom. For recording, slices were transferred to a submersion chamber continuously perfused with warmed (35–45°C) ACSF containing the following (in mM): 125 NaCl, 2.5 KCl, 2 CaCl\(_2\), 1.25 NaH\(_2\)PO\(_4\), 25 NaHCO\(_3\), and 25 d-glucose with 305 mOsm at a flow rate of 2.5–3 ml/min. Somatic whole-cell patch-clamp recordings were made from TC and TRN neurons (visually identified by infrared gradient contrast video microscopy) using a Multiclamp 700B amplifier (Molecular Devices) and pipettes with resistances of 4–6 MΩ when filled with internal solution containing the following (in mM): 130 K-gluconate, 20 KCl, 10 HEPES, 0.16 EGTA, 2 Mg-ATP, 2 Na\(_2\)-ATP, and 0.3 Na\(_2\)-GTP at pH 7.3 and 295 mOsm and supplemented with 50 μM Alexa Fluor 594 (Invitrogen). Recording solutions did not routinely include any synaptic blockers or other blocking toxins unless specifically indicated.

**Electrophysiological data were sampled at 20–50 kHz and filtered at 6 kHz.** Somatic series resistance at the start of experiments was between 9 and 15 MΩ and varied ±20% during recordings. Two-photon fluorescence microscopy using a Prairie Ultima (Prairie Technologies) microscope and titanium:sapphire pulsed laser (Chameleon Ultra II; Coherent Technologies) tuned to λ = 810 nm was combined with IR-scanning gradient contrast to make targeted dendritic patch-clamp recordings from thin (~0.7–2 μm) dendrites of TC and TRN neurons (Connelly et al., 2015). Some data included in this study was obtained from neurons recorded previously in a separate study (Connelly et al., 2015). Dendritic recording pipettes were made from borosilicate glass capillaries (BF200-100-10; Sutter Instruments) and had resistances of 25–40 MΩ when filled with the internal recording solution described above. Although this resulted in dendritic recordings with high series resistance (40–110 MΩ), the ability of high-resistance recording electrodes to faithfully record dendritic membrane potentials has been demonstrated previously by others (Nevian et al., 2007; Bathellier et al., 2009; Larkum et al., 2009; Krueppel et al., 2011) and in our own laboratory (Connelly et al., 2015). As described previously (Connelly et al., 2015), somatic and dendritic bridge balance and pipette capacitance neutralization were carefully monitored and adjusted throughout experiments by application of low-frequency (50 Hz), low amplitude (10–40 pA) current steps. Although nonbursting TRN neurons have been identified previously (Contreras et al., 1992; Brunton and Charpak, 1997; Lee et al., 2007), for the purposes of this study, all TRN cells recorded from were capable of producing LTS bursts in response to both depolarizing and hyperpolarizing current injections.

**Measurement of AP properties.** Tonic APs and LTS burst-associated APs (LTS-burst) were evoked using three different stimulation protocols and simultaneously recorded in the soma (depicted by blue traces in figures) and dendrites (depicted by red traces in figures) of TC and TRN neurons. First, to produce single tonic APs, TC and TRN neurons were depolarized to approximately ~55 mV using direct current injection to inactivate the majority of T-type Ca\(^{2+}\) channels and to prevent triggering LTS bursts and brief current injection steps of between 1 and 2 nA and 2 ms duration were injected via the somatic recording electrode (Fig. 1A). Hereafter, when recorded at the soma, these will be referred to as evoked APs (EAPs) and when recorded in the dendrites as evoked bAPs (EbAPs). The average of a minimum of 10 EAPs/EbAPs were used to measure AP properties at each different dendritic location. To investigate frequency-dependent effects on AP backpropagation, EAPs were also evoked as trains of 3 spikes at 10, 30, and 50 Hz. Second, trains of APs typical of the prolonged tonic firing observed in TC and TRN neurons in behaving animals were analyzed. These tonic APs will hereafter be referred to as TAPs when recorded at the soma and TbAPs when recorded in the dendrites. TAPs were evoked from the resting membrane potential...
of TC and TRN neurons by injecting long (0.5–1 s) depolarizing current steps (200 pA) via the somatic recording electrode. Spikes that were not associated with the initial LTS burst were considered TAPs (dashed boxes in Figs. 4C, 5C) and included in the analysis. TAPs and TbAPs were detected using a threshold-crossing detection method and aligned to threshold before averaging. A minimum of 10 TAPs/TbAPs were averaged for measurement of spike properties at each different recording location. Finally, backpropagation of LTS-burst-associated APs was assessed for spike bursts evoked by a 1-s-long hyperpolarizing current injection. For all spikes, including EAPs/EbAPs, TAPs/TbAPs, and BXAPs/BXbAPs, the first temporal derivative (δV/δt) of the somatic and dendritic Na\(^+\) spike was calculated by multiplying the voltage response of the smallest EPSP by a scaling factor (i.e., measured EPSP1 was multiplied by three to estimate expected EPSPs). To compare the size of measured dendritic EPSPs recorded in response to EPSC-like current injection versus the size of the expected EPSPs generated by EPSCs with linearly increasing amplitude, expected EPSPs were calculated by multiplying the voltage response of the smallest EPSP (EPSP\(_{\text{c}}\)) evoked by the minimal EPSC (EPSC\(_{\text{c}}\)) by the necessary scaling factor (i.e., measured EPSP\(_{\text{c}}\)) was multiplied by three to estimate the expected EPSP evoked by a 30 pA EPSC. 

Data analysis and statistics. Throughout this study, a previously used schematic illustration scheme (Connelly et al., 2015) is used to distinguish experiments performed in TC and TRN neurons. For clarity, these schematic symbols are redefined in Figures 4 (TC) and 5 (TRN). Distances between the dendritic and somatic recording electrode were measured post hoc from 2D maximum intensity projections of 3D Z-series image stacks (120–150 images at 1 μm step size) collected at the end of each experiment as described in Errington et al. (2010). The dendrites recorded from were constrained to a narrow optical plane (≤20 μm Z variance) parallel to the surface of the slice. Data analysis was performed using Clampfit 10 (Molecular Devices), Excel (Microsoft), ImageJ, and Prism (GraphPad) software. Statistical testing was by unpaired t test or repeated-measures ANOVA where appropriate and all data values are presented as mean ± SEM. Quoted values for n are the number of neurons in each group.

Results

AP and dendritic sodium spike initiation in TC and TRN neurons

Both TC and TRN neurons characteristically fire APs in two distinct patterns, namely tonic and burst mode, depending upon their membrane potential, which in vivo is dependent on behavioral state (Sherman, 2001). In burst mode, which most typically,
but not exclusively, occurs during low vigilance states and sleep. APs in both TC and TRN neurons are fired on the crest of a Ca$^{2+}$-LTS at frequencies of several hundred hertz with interspike intervals as short as 2 ms. In contrast, during periods of wakefulness, APs are fired in tonic mode at frequencies typically <50 Hz. This behavioral-state-dependent change in firing pattern is thought to primarily allow LTS burst spikes to transmit different signals to postsynaptic cortical targets compared with tonic spikes (Swadlow and Gusev, 2001). Nonetheless, these firing modes might also facilitate, via AP backpropagation, transmission of variable information about thalamic neuronal output into the dendritic tree and selective activation of specific dendritic signaling mechanisms. Therefore, we performed two-photon targeted paired somatodendritic recordings (Connelly et al., 2015) to assess AP backpropagation definitively in TC and TRN neurons during tonic and burst firing.

First, the ability to record the dendritic membrane potential directly afforded us the opportunity to investigate experimentally the subcellular AP initiation site in TC and TRN neurons. Consistent with the only previous study to investigate AP backpropagation directly (Williams and Stuart, 2000), our data confirm that, in TC neurons, APs originate in the axosomatic region before backpropagating into dendrites with a tangible delay regardless of whether APs are fired in the tonic mode (n = 25) or are associated with LTS bursts (n = 36). In every TC neuron from which we recorded, APs were always recorded first at the somatic recording electrode, followed by the dendritic recording electrode (data not shown). However, for TRN neurons, no equivalent direct measure of the AP initiation site exists. Therefore, we examined AP initiation during rebound LTS bursts by injecting hyperpolarizing current via the somatic or dendritic recording electrode. APs associated with the first rebound LTS, evoked by either somatic (Fig. 2A,B) or dendritic (Fig. 2C,D) hyperpolarization, were always recorded first at the somatic electrode and subsequently the dendritic electrode (n = 16). In the majority of TRN neurons, after strong membrane potential hyperpolarization, rhythmic sequences of LTS bursts occur (Fig. 2A,C). It has been suggested that these secondary LTS bursts differ from the primary burst in respect of the relative contribution of R-type (Ca$^{2+}$,2.3) versus T-type Ca$^{2+}$ (Ca$^{2+}$,3.2 and 3.3) channels (Zaman et al., 2011). However, our recordings show that APs associated with secondary LTS bursts follow the same pattern as observed for the initial LTS burst, being observed first in the soma and then dendrite with the same latency between AP peaks for both bursts (Fig. 2B,D). Similarly to LTS burst-associated APs, tonic APs were also always recorded first at the somatic recording electrode, followed by the dendritic electrode (Fig. 2E,F), even when evoked by injecting depolarizing current into the dendrites (Fig. 2G,H). Therefore, by recording directly from dendrites, these data demonstrate for the first time that APs in TRN neurons are initiated in the axosomatic region and not in dendrites.

Nonetheless, in thin dendrites of other neurons, notably cortical and hippocampal pyramidal neurons (Golding and Spruston, 1998, Bathellier et al., 2009), voltage-dependent dendritic Na$^+$-spikes can be initiated locally. To test whether TC and/or TRN dendrites support dendritic Na$^+$ spike initiation, we injected EPSC-like currents into distal dendrites (TC: 100–156 μm, n = 9; TRN: 129–200 μm, n = 3; Bathellier et al., 2009; Ledergerber and Larkum 2010) and compared the recorded dendritic EPSPs with the expected EPSP, estimated using the amplitude of the smallest recorded subthreshold EPSP (see Materials and Methods). In TRN neurons, fast dendritically injected EPSCs produced local EPSPs with measured amplitudes that were linearly related to the expected EPSP (n = 3; Fig. 3A). Moreover, we observed no indication of a nonlinear step in EPSP amplitude or maximum δV/δt (Fig. 3A) indicative of Na$^+$ spikes as described in dendrites of other neurons (Bathellier et al., 2009; Ledergerber and Larkum 2010). Injection of slower dendritic EPSCs, although producing larger local EPSPs, was also incapable of evoking nonlinear responses (Fig. 3B). Despite the small number of observations for EPSP-like current injections into the distal dendrites of TRN neurons, the highly linear relationship between the measured local EPSP and the expected EPSP support the conclusion that Na$^+$ spikes are not generated in TRN dendrites. This is supported by the fact that we did not observe the occurrence of spikes in dendrites before the soma even with larger and longer depolarizing steps (Fig. 2G). In TC neurons (n = 9), we observed a weakly sublinear relationship between measured and expected dendritic EPSP amplitude for both fast (Fig. 3C) and slow (Fig. 3D) injected EPSCs. Finally, for TC neurons, maximum EPSP δV/δt was linearly related to the size of the injected EPSC (Fig. 3C,D). Therefore, distal dendrites of both TC and TRN neurons are incapable of initiating local dendritic voltage-dependent Na$^+$-spikes.

**Backpropagation of tonic APs in TC and TRN neuron dendrites**

Having established their axosomatic origin, we performed a detailed examination of AP backpropagation in TC and TRN neuron dendrites. First, we evoked APs in the tonic mode, which is typified by a depolarized resting membrane potential and commonly associated in vivo with wakefulness. To do this, we recorded membrane potential simultaneously at the soma and dendrites (Figs. 4A, 5A) while evoking spikes using two distinct current injection protocols. First, to measure backpropagation of EAPs, neurons were depolarized to −55 mV using d.c. injection and brief somatic current injections (2 ms, 1–2 nA) were used to trigger spikes (Figs. 4B, 5B). The mean amplitude of EAPS was 50.7 ± 1.5 mV for TC neurons (n = 25; Fig. 4B,F) and 59.8 ± 1.3 mV for TRN neurons (n = 7; p < 0.01, unpaired t-test; Fig. 5B,F). Consistent with differences observed between other excitatory and inhibitory neurons, EAPS of GABAergic TRN neurons were markedly faster than those of TC neurons having significantly shorter half-widths (TRN: 0.27 ± 0.02 ms, n = 7; TC: 0.57 ± 0.01 ms, n = 25; p < 0.0001 unpaired t-test; Figs. 4B, E, H, 5B, E, H) and greater maximum δV/δt (TRN: 369.0 ± 21.3 mV · ms$^{-1}$, n = 7; TC: 202.0 ± 8.4 mV · ms$^{-1}$, n = 25; p < 0.0001, unpaired t-test; Figs. 4B, E, J, 5B, E, J). In both TC and TRN neurons, EAPS showed considerable distance-dependent voltage attenuation as they backpropagated into dendrites (Figs. 4B, E, F, 5B, E, F). To quantify the spatial extent of AP backpropagation, we fitted plots of EbAP amplitude, normalized to EAP amplitude, against distance of the dendritic recording location from the soma with single exponential functions. These fits revealed λ$_{eff}$ for EAP attenuation of 64 μm for TC neurons (Fig. 4G) and 57 μm for TRN neurons (Fig. 5G). Therefore, for EAPS, attenuation of spike amplitude is greater per micrometer in TRN neurons than in TC neurons and, for both cells, bAP attenuation is steeper than that found in other neurons, for example, cortical pyramidal cells (Nevian et al., 2007). Consistent with AP backpropagation in other neurons (Nevian et al., 2007, Larkum et al., 2009, Bathellier et al., 2009; Ledergerber and Larkum 2010), as spikes propagated into the dendritic tree of both TC and TRN neurons, they became markedly broader and slower, as shown by distance-dependent increases in AP half-width and reduction in maximum δV/δt.
Finally, as described above, in TC and TRN neurons, the EAP peak was always observed before the peak of the EbAP. In TC neurons, a linear regression fit to the peak spike latency versus distance from the soma revealed an average conduction velocity for EbAPs of 243.1 μm/ms (n = 25; Fig. 4I). For TRN neurons, due to the relatively small number of recordings and strong bAP attenuation, it was difficult to estimate conduction velocity of EbAPs accurately. Nonetheless, by excluding a single outlying data point from our analysis (Fig. 5I), we were able to estimate a conduction velocity of 295.7 μm/ms (n = 6) for EbAPs in TRN neurons (Fig. 5I). This measurement is similar to that for recorded for TbAPs (described below) in TRN dendrites, for which we have a larger dataset, and demonstrates that bAPs propagate more rapidly in TRN versus TC dendrites.

Because, in vivo, both TC and TRN neurons often respond to stimuli by firing trains of TAPs, we next investigated spike backpropagation during such tonic firing by injecting prolonged somatic depolarizing current steps. APs associated with the initial LTS burst were excluded from the analysis (Figs. 4C, 5C). In both TC and TRN neurons, we found no apparent activity-dependent...
Figure 3. Absence of local dendritic sodium spike initiation in thalamocortical and thalamic reticular nucleus neurons. A, Voltage at the soma (blue) and dendrite (red) resulting from fast EPSC-like current injections of increasing size into a distal dendrite (200 μm) of a TRN neuron. Even large (>40 mV) dendritic depolarizations cannot produce local dendritic Na⁺ spikes. Plots demonstrate the linear relationship between the measured dendritic EPSP and the expected EPSP and between the maximum V/\partial t of dendritic EPSPs and the size of the injected dendritic EPSC.

B, As in A, but for slow dendritic EPSC injections. Slow EPSCs induce larger dendritic EPSPs but still reveal no evidence for the presence of dendritic Na⁺ spikes.

C, Voltage at the soma (blue) and dendrite (red) resulting from fast EPSC-like current injections of increasing size into a distal dendrite (104 μm) of a TC neuron. Plots demonstrating the weakly sublinear relationship between measured and expected dendritic EPSPs in TC neurons and the linear increase in maximum dendritic EPSP V/\partial t with increasing EPSC size.

D, As in C, but for slow dendritic EPSC injections.
Changes in individual spike properties of TAPs or TbAPs throughout the spike train. Figures 4D and 5D show the average TAP (blue) and TbAP (red) overlaid onto the individual APs (gray) occurring during the trains shown for TC (Fig. 4C) and TRN (Fig. 5C) neurons, respectively. Interestingly, in TC neurons, we found notable differences between TAPs and EAPs. Both the amplitudes $(52.6 \pm 1.2 \text{ mV}, n = 18, p > 0.05, \text{unpaired } t\text{-test}; \text{Fig. 4E,F})$ and maximum $\delta V/\delta t$ $(198.0 \pm 9.1 \text{ mV} \cdot \text{ms}^{-1}, n = 18, p < 0.05,$
unpaired $t$ test; Fig. 4E,F) of TAPs were not significantly different from those of EAPs. However, the half-widths of TAPs (0.76 ± 0.04 ms, $n = 18$) were significantly greater than those of EAPs (0.57 ± 0.01 ms, $n = 25$, $p < 0.001$, unpaired $t$ test; Fig. 4D,E,H).

Coincident with this, TbAPs, on average, had significantly larger amplitudes than EbAPs (TbAP: 23.1 ± 3.0 mV, $n = 25$, EbAP: 15.7 ± 2.1 mV, $n = 18$, $p < 0.05$, unpaired $t$ test; Fig. 4E,F), having an $\lambda_{\text{eff}}$ of 91 μm (Fig. 4G). Moreover, the peak latency

Figure 5. Tonic AP backpropagation in TRN neurons. A, Overlay of two-photon fluorescence maximum intensity projection and scanning gradient contrast image of a somatosensory TRN neuron showing the location of the somatic (blue) and dendritic (red) recording pipettes. Inset, Schematic of the recording configuration illustrating the neuron recorded from (TRN) and the placement of electrodes. This schematic is used throughout to indicate that the illustrated data are drawn from TRN neuron recordings. B, $V_m$, $\delta V/\delta t$ for APs evoked by brief somatic current injections recorded in the dendrites (EbAP, red) of four different TRN neurons at increasing distances from the soma (EAP, blue). C, Typical train of tonic APs recorded in the soma (TAP, blue) and dendrites (TbAP, red) of a TRN neuron evoked by a prolonged somatic current injection step. Dashed boxes enclose those spikes included for analysis. D, Gray traces show overlaid the individual TAPs and TbAPs measured from the period indicated by the dashed box in C. The blue and red traces are the average TAP and TbAP, respectively. E, EAP (blue) and EbAP (red) recorded at the same dendritic location in the same TRN neuron compared with the average TAP and TbAP (gray traces). F, Amplitude of EAPs (open blue squares) and EbAPs (open red circles) and TAPs (filled blue squares) and TbAPs (filled red circles) versus recording distance from soma. For clarity, somatic values are mean ± SEM. Inset bar chart shows significant difference in mean somatic EAPs and TAP amplitude. G, Normalized dendritic EbAP (EbAP/EAP) and TbAP (TbAP/TAP) amplitude versus recording distance from soma. H, EAP and EbAP and TAP and TbAP half-width versus recording distance from soma. I, Peak latency between EAPs and EbAPs and TAPs and TbAPs versus distance from the soma. J, Maximum $\delta V/\delta t$ versus distance from the soma for EAPs and EbAPs and TAPs and TbAPs. Inset shows no significant difference in the somatic maximum $\delta V/\delta t$ for TAPs versus EAPs.
between TAPs and TbAPs in TC neurons was greater than the latency between EAPs and EbAPs, with an average conduction velocity for TbAPs of 141.9 μm/ms (Fig. 4I). Therefore, it appears that slower TAPs, backpropagate more efficiently into the dendrites of TC neurons than EAPs, in which somatic half-widths are shorter. The reason for this difference in TAP and EAP half-width is due to the membrane potential from which they are evoked. On average, the resting membrane potential before spike onset and that this, in turn, determines the efficiency of spike backpropagation into dendrites.

In comparison, in TRN neurons, we did not see differences between EAPs and TbAPs. Although TAPs in TRN neurons had significantly smaller amplitudes than EAPs (TAP: 49.6 ± 1.4 mV, n = 16, p < 0.001, unpaired t test; Fig. 5E,F), their half-widths were not significantly different (TAP: 0.23 ± 0.01 ms, n = 16, EAP: 0.27 ± 0.02 ms, n = 7, p > 0.05, unpaired t test; Fig. 5D,E,F) despite being evoked from different membrane potentials (TAP ~ −45 to −40 mV, Fig. 5D, EAP ~ −60 to −55 mV, Fig. 5E). Therefore, in TRN cells, the mean amplitudes of TbAPs were not significantly different from those of EbAPs (TbAP: 9.2 ± 1.6 mV, n = 16, EbAP: 14.0 ± 3.4 mV, n = 7, p < 0.05, unpaired t test; Fig. 5D–F), with TbAPs having an λ_eff of 40 μm, almost identical to that of EbAPs (Fig. 5G). The conduction velocity of TbAPs in TRN dendrites was 424 μm/ms (n = 16; Fig. 5I). Therefore, in TRN neurons, the preceding membrane potential has little effect on the shape and duration of individual APs and, as a result, their backpropagation into the dendritic tree.

Finally, we examined two physiological mechanisms previously associated with bAPs, namely frequency-dependent bAP attenuation (Jung et al., 1997) and frequency-dependent Ca^{2+} electrogenesis (Larkum et al., 1999; Ledergerber and Larkum, 2010). To do this, we evoked trains of five EAPs while performing simultaneous somatodendritic recordings. We restricted our analysis to physiologically relevant tonic firing frequencies (10/30/50 Hz) in these neurons. In TC neurons, our previous work using two-photon imaging indicated that neither frequency-dependent reduction in bAP-associated Ca^{2+} signals nor frequency-dependent initiation of dendritic Ca^{2+} spikes was a feature of these neurons (Errington et al., 2010). Our current work confirms these earlier findings, demonstrating no significant reduction in bAP amplitude (n = 25) or evidence of regenerative Ca^{2+} spikes across all frequencies tested (Fig. 6A,B). By dividing the amplitude of the fifth spike in the evoked train by the amplitude of the first spike for both the somatic and dendritic recordings, we found that frequency-dependent attenuation of spike amplitudes during trains was not observed at any distance from the soma for all spike frequencies in TC neurons (Fig. 6C). In TRN neurons, these properties have not been previously tested either directly by patch...
clamp or by imaging approaches. Similar to TC neurons, we observed no difference in bAP amplitude in TRN neurons \((n = 7)\) throughout spike trains at each tested frequency \((\text{Fig. 6D,E})\). Once more, like TC neurons, frequency-dependent attenuation did not occur at any dendritic recording location at any frequency tested \((\text{Fig. 6F})\). Moreover, no evidence of bAP evoked dendritic \(\text{Ca}^{2+}\) electrogenesis, such as an increased spike afterdepolarization, was observed at frequencies up to 50 Hz \((\text{Fig. 6D})\).

In sum, these data demonstrate directly that, during tonic firing, APs backpropagate into the dendritic tree of both TC and TRN neurons but undergo strong voltage attenuation. In TC, but not TRN, neurons, membrane potential before spike onset can influence AP shape and duration and this produces variability in backpropagation efficiency. Moreover, the strength of AP backpropagation is not affected by either the frequency of firing or duration of the train, APs are always generated in the axosomatic region and TC and TRN dendrites do not support Na\(^+\)-spike electrogenesis, most likely as a consequence of insufficient distal dendritic Na\(^+\) conductance.

**Distinct TC and TRN LTS burst properties determine AP backpropagation**

So far, we have focused on tonic APs fired when thalamic neurons are depolarized. Although previous \(\text{Ca}^{2+}\)-imaging studies \((\text{Cran dall et al., 2010; Errington et al., 2010})\) suggest that LTS-burst APs contribute little to dendritic \(\text{Ca}^{2+}\) signaling in TC and TRN neurons, distal LTS-burst AP voltage responses remain to be described. Therefore, we investigated LTS-burst AP backpropagation in both types of thalamic neurons. To do this, we injected hyperpolarizing current into the soma of TC and TRN neurons and measured somatic (B1AP) and dendritic (B2bAP) rebound LTS-burst APs. Whereas the numbers of APs per burst varied in both TC \((\text{range: 1–6 spikes, 3.6 ± 0.3 spikes, } n = 36)\) and TRN \((\text{range: 2–9 spikes, 5.5 ± 0.7 spikes, } n = 16)\) neurons, all LTS bursts \((3 \text{ or more spikes per burst})\) displayed the decelerating and accelerating–decelerating temporal firing patterns characteristic of TC and TRN neurons \((\text{Figs. 7A, 8A, 9G})\). When recording somatic LTS-burst APs, we found differences in individual spike properties throughout the burst that clearly distinguish TC from TRN neuron bursts. First, in TC neurons, the first spike, B1AP, has a substantially more hyperpolarized threshold than the next spike, B2AP, but the AP threshold varies little between B2bAP and subsequent spikes in the burst \((\text{Fig. 7A})\). This threshold transition can be readily observed in the somatic AP phase plot \((\text{Fig. 7B})\). In comparison, as has been described in vivo \((\text{Munoz and Fuentelaba, 2012})\), in TRN neurons, a much smaller, incremental increase in spike threshold occurs throughout the duration of the burst \((\text{Fig. 8A,B})\). Second, in TC neurons, as has been described previously \((\text{Turner et al., 1997; Williams and Stuart, 2000})\), we saw a significant decrease in AP amplitude between B1AP \((54.3 ± 1.1 \text{ mV, } n = 36)\) and B2AP \((39.8 ± 1.0 \text{ mV, } n = 30, p < 0.0001, \text{paired } t \text{ test})\), followed by a progressive increase in amplitude for subsequent spikes \((\text{B3AP: 38.6 ± 1.2 mV, } n = 23, \text{ B4AP: 41.0 ± 1.0 mV, } n = 22; \text{ B5AP: 44.5 ± 1.4 mV, } n = 12, \text{ B6AP: 46.3 ± 1.8 mV, } n = 5)\). However, late spikes never matched the amplitude of B1AP. Again, this is clearly illustrated by the AP phase plot \((\text{Fig. 7B})\). This is not a feature of APs during LTS bursts in TRN neurons, the amplitudes of which do not show significant variability throughout the burst \((\text{B1AP: 55.5 ± 1.3 mV, } n = 16, \text{ B2AP: 55.0 ± 1.4 mV, } n = 16, \text{ B3AP: 52.7 ± 1.6 mV, } n = 13, \text{ B4AP: 50.7 ± 1.2 mV, } n = 11, \text{ B5AP: 48.9 ± 1.1 mV, } n = 9, \text{ B6AP: 47.7 ± 1.2 mV, } n = 8, \text{ B7AP: 47.2 ± 1.4 mV, } n = 7, \text{ B8AP: 47.0 ± 2.1 mV, } n = 6, \text{ B9AP: 46.3 ± 3.4 mV, } n = 2)\).

This difference between TC and TRN neurons in somatic spike amplitude during bursts is shown clearly in Figure 9H, where somatic AP amplitude, normalized to B1AP, is shown for all bursts comprising three or more spikes. Third, and most critically, in TC neurons, we observed a progressive increase in somatic spike half-width \((\text{B1AP: 0.42 ± 0.01 ms, } n = 36, \text{ B2AP: 0.50 ± 0.01 ms, } n = 30, \text{ B3AP: 0.58 ± 0.01 ms, } n = 23, \text{ B4AP: 0.71 ± 0.02 ms, } n = 22, \text{ B5AP: 0.84 ± 0.03 ms, } n = 12, \text{ B6AP: 0.92 ± 0.06, } n = 5)\). In this respect, APs at the end of LTS bursts more closely resembled TAPs in terms of their amplitude \((\text{Fig. 9A–C})\) and half-width \((\text{Fig. 9A–C})\). In comparison, in TRN neurons, spike half-width \((\text{B1AP: 0.23 ± 0.01 ms, } n = 16, \text{ B2AP: 0.24 ± 0.1 ms, } n = 16, \text{ B3AP: 0.24 ± 0.01 ms, } n = 13, \text{ B4AP: 0.23 ± 0.01 ms, } n = 11, \text{ B5AP: 0.23 ± 0.01 ms, } n = 9, \text{ B6AP: 0.23 ± 0.01 mV, } n = 8, \text{ B7AP: 0.23 ± 0.01 ms, } n = 7, \text{ B8AP: 0.22 ± 0.01 ms } n = 6, \text{ B9AP: 0.24 ± 0.01 ms, } n = 2)\). This difference is further illustrated by the maximum AP \(B5/\delta t\) of B1AP \((\text{Fig. 8B, A})\) and maximum AP \(B5/\delta t\) of B1AP \((\text{Fig. 8A, B})\) remained unaltered throughout the LTS burst, as demonstrated clearly by the somatic AP phase plot \((\text{Fig. 8B})\). Comparing B1AP \((\text{Fig. 9E})\) of a TRN LTS burst with B1AP \((\text{Fig. 9D})\) illustrates the lack of change in spike amplitude \((\text{Fig. 9H})\) and half-width \((\text{Fig. 9K})\) during the burst. Moreover, APs during LTS bursts in TRN neurons were not different from TAPs in any of the measured parameters \((\text{Fig. 9F, H, K, L})\). Therefore, unlike TC neurons, TRN neuron APs are invariable and display fast-spiking properties in both LTS burst and tonic mode.

In TC neurons, the increase in somatic half-widths of LTS-burst APs were coincident with an increase in the efficiency of propagation into the dendrites. Similarly to their EAP and TAP counterparts, all LTS-burst APs in TC neurons from B1AP to B5AP showed a significant distance-dependent reduction in amplitude \((\text{Fig. 7A–F})\) and maximum \(B5/\delta t\) \((\text{Fig. 7A–D})\) as they propagated into dendrites. However, in contrast to somatic APs, in which the amplitude of spikes up to B6AP were reduced compared with B1AP \((\text{Fig. 7A})\), dendritic AP amplitudes were initially reduced, but subsequently increased such that, on average, B3bAP and B4bAP were in fact larger than B1bAP \((\text{Fig. 7C})\). Figure 9H illustrates clearly how B3bAP has a greater amplitude than B2bAP despite the fact that B3bAP is smaller than B2bAP. When considering these differences in somatic and dendritic AP amplitude during the LTS burst together, it is apparent that APs later in the LTS burst undergo weaker attenuation as they propagate into the dendrites \((\text{Fig. 7F})\). Comparing AP attenuation for subsequent LTS burst spikes against the attenuation experienced by B3bAP illustrates the increased backpropagation efficiency of late-burst spikes compared with their earlier counterparts \((\text{Fig. 9A, B, J})\). Indeed, \(\lambda_{\text{att}}\) of B3bAP was 72 μm, whereas for B2bAP, it was 156 μm. Furthermore, the decrease in AP attenuation during the LTS burst was accompanied by an increase in the peak latency between somatic and dendritic APs \((\text{Fig. 7A, C, G})\). When the latency for each spike was normalized to the latency of the first LTS burst spike, we observed a greater than twofold increase in
the latency between B6AP and B6bAP versus B1AP and B1bAP and
clear similarity in the latency between late-burst spikes and
TbAPs (Fig. 9L). Therefore, during LTS bursts in TC neurons, AP
backpropagation strength is variable. B1AP, despite having a
greater somatic amplitude than all subsequent APs in the burst,
undergoes the strongest attenuation such that B1bAP is actually
smaller than bAPs later in the burst despite their somatic ampli-
tudes being less. The degree of attenuation is coincident with a
marked broadening of the somatic APs during the burst.

In comparison, TRN neurons did not show variability in spike
backpropagation during LTS bursts. Like EAPs and TAPs, all LTS
burst spikes from B1AP through to B9AP showed significant
distance-dependent attenuation as they propagated into the den-
drites (Fig. 8C, E, F). As can be seen in Figures 8, A and C, and 9F,
somatic and dendritic LTS-burst APs showed only a small reduc-
tion in amplitude compared with B1AP and B1bAP respectively as
the burst progressed. Therefore, the distance-dependent AP am-
plitude attenuation across successive spikes remained virtually
unaltered compared with the attenuation of B1AP and was nearly
identical to the attenuation observed for TbAPs in TRN dendrites
(Fig. 9I). In fact, λeff of B1bAP was 36 μm and, by B7bAP, it was
unchanged at 39 μm. This is clearly illustrated by comparing
B1bAP of a typical TRN burst with B7bAP of the same burst and
the averaged TbAP from the same neuron (Fig. 9D–F). Further-
more, the lack of variability in attenuation strength between LTS-
burst APs in TRN neuron dendrites was not related to the
distance from the soma. Plotting both the mean and range of
attenuation for LTS burst spikes against distance of the dendritic
recording from the soma reveals that variability in AP attenua-
tion during bursts is observed throughout the dendritic tree in

Figure 7. Backpropagation of APs during LTS bursts in TC neurons. A, Somatic Vm and ∂V/∂t of a TC neuron LTS burst. Individual APs within the burst (B1AP–B6AP) are color coded red to purple. B, AP phase plot for the somatic LTS-burst APs shown in A. Colors match the code in A. Gray boxes highlight spike threshold (i), maximum amplitude (ii), and maximum ∂V/∂t (iii). C, Dendritic (B1bAP–B6bAP) Vm and ∂V/∂t of the LTS burst shown in A. D, AP phase plot for the dendritic LTS burst shown in C. Colors match the code in A. E, Amplitude of somatic and dendritic LTS-burst APs versus recording distance from soma. Somatic values are mean ± SEM. Inset shows the color coding for each spike in the train. F, Normalized dendritic LTS-burst AP amplitude versus recording distance from soma. G, Peak soma to dendrite latency versus distance from the soma for LTS-burst APs. H, Maximum ∂V/∂t versus distance from the soma for LTS-burst APs.
TC neurons (black circles and bars), but not TRN neurons (Fig. 9J, orange circles and bars). Finally, in TRN neurons, the peak latency between somatic and dendritic APs, unlike TC neurons, was not altered for any LTS burst spikes or TbAPs when normalized to B1bAP (Fig. 9L).

**Discussion**

The important outcomes of this study are as follows: (1) tonic and LTS-burst APs are generated in the axosomatic region in both TC and TRN neurons; (2) APs, in both firing modes in both TC and TRN neurons, backpropagate into the dendritic tree undergoing substantial distance-dependent voltage attenuation; (3) differences in somatic spike half-width between APs within LTS bursts and between LTS bursts and TbAPs result in variable AP backpropagation efficiency in TC, but not TRN, neurons; and (4) neither TC or TRN neuron dendrites support generation of local dendritic Na+ spikes. When considered alongside our own previous studies (Errington et al., 2010, 2012; Connelly et al., 2015, 2016) and those of others (Crandall et al., 2010, Cueni et al., 2008, Chausson et al., 2013), these new findings allow us to build a more complete picture of the role that dendrites play in thalamic neuron electrogenesis and signaling.

First, using somatodendritic recording, we established that both tonic and LTS-burst APs are recorded first at the soma and subsequently in the dendrites in both TC and TRN neurons. This
Figure 9. Variability of somatic spike shape influences backpropagation during bursts in TC but not TRN neurons. A, Example traces showing the first spike of a typical LTS burst recorded in the soma (B1AP, blue) and dendrites (B1bAP, red) of a TC neuron. B, B5AP (blue) and B5bAP (red) from the same LTS burst in A overlaid onto B1AP and B1bAP (gray). C, Average TAP (blue) and TbAP (red) from the same neuron as in A overlaid onto B1AP and B1bAP (gray). D, Example traces showing the first spike of a typical LTS burst recorded in the soma (B1AP, blue) and dendrites (B1bAP, red) of a TRN neuron. E, B7AP (blue) and B7bAP (red) from the same LTS burst in D overlaid onto B1AP and B1bAP (gray). F, Average TAP and TbAP from the same neuron (Figure legend continues.)
demonstrates that, consistent with most other mammalian neurons (Clark et al., 2005; Kole et al., 2008; Schmidt-Hieber et al., 2008; Fousst et al., 2010), APs are initiated in the perisomatic region, most likely the axon initial segment. For TC neurons, these results support those previously reported by Williams and Stuart (2000), but this is the first direct demonstration of axosomatic AP origin in TRN neurons. Ultrastructural studies have shown previously that axons occasionally originate from proximal dendrites rather than the soma in TRN neurons (Pinault and Deschênes, 1998). Although our limited number of dendritic recordings cannot exclude the possibility that spike initiation may occur in axons branching from dendrites rather than the soma, the fact that we observed no cases where APs were recorded first in the dendrite indicates that APs are triggered close to the soma and not from some distal part of the dendritic tree. From a physiological and computational perspective, it appears to be of little significance whether APs are generated in an axon that originates at the soma or from a proximal dendrite. Moreover, comparing the peak latency between somatic and dendritic APs confirms that, unsurprisingly, APs fired on the crest of low threshold Ca\(^{2+}\) spikes originate from the same subcellular location as tonic APs.

By recording directly from dendrites, our new findings establish definitively the distance dependence of AP backpropagation in TC and TRN. In TC neurons, dendritic membrane voltage recordings strongly support previous findings from two-photon Ca\(^{2+}\) imaging studies (Crandall et al., 2010; Errington et al., 2010, 2012; Sieber et al., 2013). We found that tonic APs undergo strong voltage attenuation as they backpropagate into the dendritic tree, having amplitudes only 10–20% of those recorded at the soma once they reach the dendritic tips. This is consistent with larger bAP-evoked Ca\(^{2+}\) signals in proximal parts of the dendritic tree and suggests that the absence of bAP-evoked Ca\(^{2+}\) signals in distal branches (Crandall et al., 2010; Errington et al., 2010, 2012; Sieber et al., 2013) results from weak Na\(^+\) conductance-dependent active backpropagation coupled with strong passive dendritic filtering rather than a lack of distal dendritic high-voltage-activated Ca\(^{2+}\) channels. Moreover, because bAPs do not undergo any frequency-dependent reduction in their amplitude, they can reliably signal cellular spike output to the dendritic tree even for high firing frequencies. In TRN neurons, we also observed strong voltage attenuation of APs as they backpropagated into the dendritic tree. In this respect, our data from the first direct patch-clamp recordings of APs in TRN neuron dendrites more closely support the conclusion that bAP-evoked Ca\(^{2+}\) signals are minimal in distal dendrites (Cueni et al., 2008; Crandall et al., 2010) than those of Chausson et al. (2013), who reported bAP-evoked Ca\(^{2+}\) entry throughout much of the dendritic tree. The reasons for these discrepancies remain unclear, but might relate to differences in experimental conditions. For example, the duration of the AP trains in the study by Chausson et al. (2013) were typically much longer, introducing the possibility that the distal dendritic Ca\(^{2+}\) signals that they observed were due to slow axial dendritic diffusion rather than local Ca\(^{2+}\) entry.

APs are efficient triggers for neurotransmitter release at both axonal and dendritic presynaptic terminals (Bischoffberger and Jonas, 1997). In TRN, GABAergic synapses between dendrites have been hypothesized to exist based on electron microscopy studies, although their functional existence remains to be demonstrated. We demonstrate here that APs attenuate significantly as they propagate into TRN neuron dendrites, producing depolarizations of only 10 mV or less at distances >100 µm from the soma. Therefore, if AP-triggered neurotransmitter release occurs at dendrodendritic synapses between TRN neurons, then it is probable that those synapses will be located close to the soma. Conversely, dendrodendritic synapses may be located at more distal locations, but in this case, given their robust attenuation, it is unlikely that bAPs would trigger release at these synapses. In contrast, release at distal dendrodendritic synapses may be triggered by low-threshold Ca\(^{2+}\) spikes, which as we have shown previously, are associated with global dendritic depolarization (Connelly et al., 2015). Importantly, therefore, signaling at dendrodendritic synapses in TRN, if it occurs, may show marked behavioral state dependence.

Functional evidence for the existence of dendrodendritic chemical synapses in TRN remains absent. In comparison, the functional evidence for the presence of electrical synapses formed by Cx36-dependent gap junctions between TRN dendrites is far more convincing (Landisman et al., 2002; Landisman and Connors, 2005; Lee et al., 2014). These electrical synapses transmit both slow membrane potential changes and fast APs between TRN neurons. Based on paired somatic recordings, the efficiency of transmission of fast Na\(^{+}\) spikes through TRN gap junctions is reasonably low. However, dye-coupling experiments also demonstrate the presence of coupled cells with intrasomatic distances of several hundred micrometers indicating coupling between distal dendrites (Lee et al., 2014). Therefore, although gap junctions themselves act as low-pass filters, the attenuation of spike amplitude by TRN dendrites is also likely to have a critical role in electrical synaptic transmission. Considering the strong soma to dendrite attenuation of fast signals such as APs in TRN dendrites alongside the highly efficient soma to dendrite transfer of slow voltage signals and the global mechanism of LTS generation (Connelly et al., 2015), it seems likely that these dendrites are optimized for slower electrical signaling. Conversely, because high local dendritic input impedance (Connelly et al., 2015) produces large local EPSPs, these synaptic potentials may be transmitted between TRN neurons through distal gap junctions. Whereas, as a result of strong dendrite to soma voltage attenuation (Connelly et al., 2015), these potentials may not individually be sufficient to influence somatic membrane potential, they could be integrated with other local chemical and electrical synaptic signals to control cellular output.

We have shown here in TC neurons that somatic AP half-width is influenced by the preceding membrane potential with important consequences for AP backpropagation. When APs were evoked from a resting membrane potential of ~55 mV, their half-width was significantly shorter than when fired from a more depolarized potential close to spike threshold. This pattern is also observed during an LTS burst, in which the first spike in the burst is significantly faster than the fifth or sixth spike. The physiological significance of this difference in AP backpropagation between burst and tonic firing in TC, but not TRN, neurons remains to be determined. It is possible that these differences in spike shape are more significant to the axonal output of thalamic neurons rather than
having a major impact on dendritic signaling. For example, at the mossy fiber-CA3 synapse in the hippocampus, repetitive axonal stimulation leads to broadening of APs in the presynaptic terminal and enhanced neurotransmitter release via inactivation of A-type K⁺ channels (Geiger and Jonas, 2000). Conversely, during TC neuron bursts, the initial spike in the burst is thought to have the greatest efficacy in driving postsynaptic neocortical spikes, whereas subsequent spikes in a burst are no more efficacious than their tonic counterparts (Swadlow and Gusev, 2001).

In this study, we have demonstrated directly for the first time that APs undergo strong voltage attenuation as they propagate into the dendrites of both excitatory glutamatergic TC and inhibitory GABAergic TRN neurons, but that state-dependent variability exists between tonic and LTS-burst dendritic AP signaling in TC neurons that are absent in TRN neurons.

References

Acuna-Goycolea C, Brenowitz SD, Regehr WG (2008) Active dendritic conductances dynamically regulate GABA release from thalamic interneurons. Neuron 57:420–431. CrossRef Medline
Antic SD (2003) Action potentials in basal and oblique dendrites of rat neocortical pyramidal neurons. J Physiol 550:35–50. CrossRef Medline
Bathellier B, Margrie TW, Larkum ME (2009) Properties of piriform cortex pyramidal cell dendrites: implications for olfactory circuit design. J Neurosci 29:12641–12652. CrossRef Medline
Bean BP (2007) The action potential in mammalian central neurons. Nat Rev Neurosci 8:451–465. CrossRef Medline
Bischofberger J, Jonas P (1997) Action potential propagation into the presynaptic dendrites of rat mitral cells. J Physiol 504:359–365. CrossRef Medline
Brunton J, Charpak S (1997) Heterogeneity of cell firing properties and opioid sensitivity in the thalamic reticular nucleus. Neuroscience 78:303–307. CrossRef Medline
Chaussou P, Leresche N, Lambert RC. (2013) Dynamics of intrinsic dendritic calcium signalling during tonic firing of thalamic reticular neurons. PLoS One 8:e72275. CrossRef Medline
Clark BA, Monsivais P, Branco T, London M, Häusser M (2005) The site of axon potential initiation in cerebellar Purkinje neurons. Nat Neurosci 8:137–139. CrossRef Medline
Connelly WM, Cruccu R, Steriade M (1992) Bursting and tonic discharges in thalamic reticular neurons. Neuron 11:6902.

Connelly WM, Crunelli V, Errington AC (2016) Passive synaptic normalization and input synchrony-dependent amplification of cortical feedback in thalamocortical neurons. J Neurosci 36:3753–3754. CrossRef Medline
Costeras D, Curto NA, Díaz R, Steriade M (1992) Bursting and tonic discharges in two classes of reticular thalamic neurons. J Neurophysiol 68:973–977. CrossRef Medline
Crandall SR, Govindiah G, Cox CL (2010) Low-threshold Ca²⁺ current amplifies distal dendritic signalling in thalamic reticular neurons. J Neurosci 30:15419–15429. CrossRef Medline
Cueni L, Canepari M, Luján R, Emmenegger Y, Watanabe M, Bond CT, Franken P, Adelmann JP, Lithüi A (2008) T-type Ca²⁺ channels, SK2 channels and SERCA gate sleep-related oscillations in thalamic dendrites. Nat Neurosci 11:683–692. CrossRef Medline
Deschénes M, Madarász-Domingh A, Steriade M (1985) Dendro-dendritic synapses in the cat reticularis thalami nucleus: a structural basis for thalamic spindle synchronization. Brain Res 334:165–168. CrossRef Medline
Errington AC, Renger JJ, Uebel VN, Crunelli V (2010) State-dependent firing determines intrinsic dendritic Ca²⁺ signalling in thalamic reticular neurons. J Neurosci 30:14843–14853. CrossRef Medline
Errington AC, Hughes SW, Crunelli V (2012) Rhythmic dendritic Ca²⁺ oscillations in thalamocortical neurons during slow non-REM sleep-related activity in vitro. J Physiol 590:3691–3700. CrossRef Medline
Foua A, Popovic M, Zecevic D, McCormick DA (2010) Action potentials initiate in the axon initial segment and propagate through axon collaterals reliably in cerebellar Purkinje neurons. J Neurosci 30:6891–6902. CrossRef Medline
Geiger JR, Jonas P (2000) Dynamic control of presynaptic Ca²⁺ inflow by fast-inactivating K⁺ channels in hippocampal mossy fibre boutons. Neuron 28:927–939. CrossRef Medline
Golding NL, Spruston N (1998) Dendritic sodium spikes are variable triggers of axonal action potentials in hippocampal CA1 pyramidal neurons. Neuron 21:1189–1200. CrossRef Medline
Jung HY, Mickus T, Spruston N (1997) Prolonged sodium channel inactivation contributes to dendritic action potential attenuation in hippocampal CA1 neurons. J Neurosci 17:6639–6646. CrossRef Medline
Kampa BM, Stuart GJ (2006) Calcium spikes in basal dendrites of layer 5 pyramidal neurons during action potential bursts. J Neurosci 26:7424–7432. CrossRef Medline
Kampa BM, Letzkus JJ, Stuart GJ (2007) Dendritic mechanisms controlling spike-time-dependent synaptic plasticity. Trends Neurosci 30:456–463. CrossRef Medline
Kole MH, Ishseru SU, Kampa BM, Williams SR, Ruben PC, Stuart GJ (2008) Action potential generation requires a high sodium channel density in the axon initial segment. Nat Neurosci 11:178–186. CrossRef Medline
Kreuppel R, Remy S, Beck H (2011) Dendritic integration in hippocampal dentate granule cells. Neuron 71:512–528. CrossRef Medline
Landisman CE, Connors BW (2005) Long-term modulation of electrical synapses in the mammalian thalamus. Science 310:1809–1813. CrossRef Medline
Landisman CE, Long MA, Beielein M, Deans MR, Paul DL, Connors BW (2002) Electrical synapses in the thalamic reticular nucleus. J Neurosci 22:1002–1009. CrossRef Medline
Larkum ME, Kaiser KM, Sakmann B (1999) Calcium electrogenesis in distal apical dendrites of layer 5 pyramidal cells at a critical frequency of back-propagating action potentials. Proc Natl Acad Sci U S A 96:14600–14604. CrossRef Medline
Larkum ME, Nevian T, Sandler M, Polsky A, Schiller J (2009) Synaptic integration in tuft dendrites of layer 5 pyramidal neurons: a new unifying principle. Science 325:756–760. CrossRef Medline
Ledgerbergher D, Larkum ME (2010) Properties of layer 6 pyramidal neuron apical dendrites. J Neurosci 30:13031–13044. CrossRef Medline
Lee SC, Patrick SL, Richardson KA, Connors BW (2014) Two functionally distinct networks of gap junction–coupled inhibitory neurons in the thalamic reticular nucleus. J Neurosci 34:13170–13182. CrossRef Medline
Lee SH, Govindiah G, Cox CL (2007) Heterogeneity of firing properties among rat thalamic reticular nucleus neurons. J Physiol 582:195–208. CrossRef Medline
Muñoz F, Fuentes P (2012) Dynamics of action potential initiation in the GABAergic thalamic reticular nucleus in vivo. PLoS One 7:e30015. CrossRef Medline
Nevian T, Larkum ME, Polsky A, Schiller J (2007) Properties of basal dendrites of layer 5 pyramidal neurons: a direct patch-clamp recording study. Nat Neurosci 10:206–214. CrossRef Medline
Pinault D, Deschénes M (1998) Projection and innervation of individual thalamic reticular axons in the thalamus of the adult rat: a three-dimen-sional graphic and morphometric analysis. J Comp Neurol 391:180–203. CrossRef Medline
Pinault D, Smith Y, Deschénes M (1997) Dendrodendritic and axonaxonic synapses in the thalamic reticular nucleus of the adult rat. J Neurosci 17:3215–3233. Medline
Schmidt-Hieber C, Jonas P, Bischofberger J (2008) Action potential initiation and propagation in hippocampal mossy fibre axons. J Physiol 586:1849–1857. CrossRef Medline
Sherman SM (2001) Tonic and burst firing: dual modes of thalamocortical relay. Trends Neurosci 24:122–126. CrossRef Medline
Sieber AR, Min R, Nevian T (2013) Non-Hebbian long-term potentiation of thalamic reticular axons in a neuron. J Neurosci 33:15675–15685. CrossRef Medline
Swadlow HA, Gusev AG (2001) The impact of ‘bursting’ thalamic impulses at a neocortical synapse. Nat Neurosci 4:402–408. CrossRef Medline
Turner JP, Anderson CM, Williams SR, Cruccu V (1997) Morphology and membrane properties of neurons in the cat ventrobasal thalamus in vitro. J Physiol 505:767–772. CrossRef Medline
Williams SR, Stuart GJ (2000) Action potential backpropagation and somato-dendritic distribution of somatic Na⁺ current in thalamic reticular neurons. J Neurosci 20:1307–1317. Medline
Zaman T, Lee K, Park C, Paydar A, Choi JH, Cheong E, Lee CJ, Shin HS (2011) Cav2.3 channels are critical for oscillatory burst discharges in the reticular thalamus and absence epilepsy. Neuron 70:95–108. CrossRef Medline