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Published in:
European Journal of Neuroscience

DOI:
10.1111/ejn.12601

Publication date:
2014

Document Version
Publisher's PDF, also known as Version of record

Link to publication in Discovery Research Portal

Citation for published version (APA):
The general anaesthetic etomidate inhibits the excitability of mouse thalamocortical relay neurons by modulating multiple modes of GABA<sub>A</sub> receptor-mediated inhibition

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Keywords: nucleus reticularis, phasic inhibition, spill-over inhibition, thalamus, tonic inhibition

Abstract

Modulation of thalamocortical (TC) relay neuron function has been implicated in the sedative and hypnotic effects of general anaesthetics. Inhibition of TC neurons is mediated predominantly by a combination of phasic and tonic inhibition, together with a recently described ‘spillover’ mode of inhibition, generated by the dynamic recruitment of extrasynaptic (GABA)<sub>Rs</sub> receptors (GABA<sub>Rs</sub>). Previous studies demonstrated that the intravenous anaesthetic etomidate enhances tonic and phasic inhibition in TC relay neurons, but it is not known how etomidate may influence spillover inhibition. Moreover, it is unclear how etomidate influences the excitability of TC neurons. Thus, to investigate the relative contribution of synaptic (GABA)<sub>x</sub> and extrasynaptic (GABA)<sub>y</sub> GABA<sub>Rs</sub> to the thalamic effects of etomidate, we performed whole-cell recordings from mouse TC neurons lacking synaptic (GABA)<sub>x</sub> or extrasynaptic (GABA)<sub>y</sub> GABA<sub>Rs</sub>. Etomidate (3 μM) significantly inhibited action-potential discharge in a manner that was dependent on facilitation of both synaptic and extrasynaptic GABA<sub>Rs</sub>, although enhanced tonic inhibition was dominant in this respect. Additionally, phasic inhibition evoked by stimulation of the nucleus reticularis exhibited a spillover component mediated by GABA<sub>Rs</sub>, which was significantly prolonged in the presence of etomidate. Thus, etomidate greatly enhanced the transient suppression of TC spike trains by evoked inhibitory postsynaptic potentials. Collectively, these results suggest that the deactivation of thalamus observed during etomidate-induced anaesthesia involves potentiation of tonic and phasic inhibition, and implicates amplification of spillover inhibition as a novel mechanism to regulate the gating of sensory information through the thalamus during anaesthetic states.

Introduction

The ability of general anaesthetics to impair consciousness, and induce analgesia, amnesia and immobility, has been exploited clinically for over a century (Rudolph & Antkowiak, 2004). Yet, despite their routine clinical use, the neuroanatomical substrates of anaesthetic actions remain enigmatic, with several components of the circuitry regulating the sleep–wake cycle implicated to date (Franks, 2008). Nevertheless, electrophysiological, neuroimaging and modelling studies consistently pinpoint the thalamus as an important neural locus for anaesthetic-induced hypnosis (Angel, 1991; Fiset et al., 1999; Alkire et al., 2000; White & Alkire, 2003; Ching et al., 2010; Andrade et al., 2012). Such observations are consistent with the classical role of the thalamus in controlling transitions between conscious states (Steriade et al., 1993).

At the cellular level, γ-aminobutyric acid (GABA)<sub>A</sub> receptors (GABA<sub>Rs</sub>) represent key molecular targets for several general anaesthetics, including the intravenous agents etomidate and propofol (Hales & Lambert, 1991; Orser et al., 1994; Uchida et al., 1995). Importantly, anaesthetic-induced modulation of inhibition may be influenced by the molecular identity of GABA<sub>Rs</sub> (Belelli et al., 1996; Bonin & Orser, 2008), which are composed of five subunits drawn from a pool of 19 isoforms (Olsen & Sieghart, 2009). Etomidate is unique in this respect, displaying selectivity for β2- or β3-containing receptors vs. those incorporating the β1 subunit (Hill-Venning et al., 1997). Subunit composition additionally influences the functional properties and spatiotemporal expression of GABA<sub>Rs</sub>. Thus, synaptically targeted GABA<sub>Rs</sub>, which mediate brief phasic inhibition, are composed of αβγ2 subunits, whereas receptors incorporating a δ rather than a γ2 subunit, are localized to extra- or peri-synaptic locations, and mediate persistent tonic inhibition (Farrant & Nusser, 2005). However, a substantial number of GABA<sub>Rs</sub> with a typically ‘synaptic’ subunit composition also exist extrasynaptically, although the functional role of such receptors remains to be established (Kasugai et al., 2010). δ-GABA<sub>Rs</sub> have been proposed as important general anaesthetic targets, a suggestion supported by their abundant expression in brain regions implicated in some anaesthetic behaviours, including thalamocortical (TC) relay.
nuclei (Belelli et al., 2005; Jia et al., 2005). However, while etomidate enhances both phasic and tonic currents recorded from TC neurons (Belelli et al., 2005), it is unknown how these respective actions influence the excitability of relay neurons.

Recently, we demonstrated that in addition to generating tonic inhibition, γ2-GABAARs of TC neurons may be recruited via a spillover-like mechanism to prolong phasic inhibition generated in response to presynaptic spike bursts (Herd et al., 2013). This observation may have particular relevance to anaesthetic effects in thalamus, given that nucleus reticularis thalami (nRT) neurons, which provide the major source of GABA-ergic innervation to TC neurons, exhibit prominent spike bursts during unconscious states. Whether and how anaesthetics may influence this ‘burst-mediated spillover’ mode is not known.

Here, we investigated the relative contribution of synaptic and extrasynaptic GABAARs to the inhibitory effects of etomidate on relay neuron output. Using transgenic mice in which TC relay neurons lack synaptic (α1100, Peden et al., 2008) or extrasynaptic (δ00, Herd et al., 2009) GABAARs, we report that, although enhancement of tonic inhibition dominates the inhibitory effect of etomidate on TC neuron excitability, the synaptic α1β2GABAAR population also significantly influences the effects of the anaesthetic. Moreover, we demonstrate that etomidate greatly prolongs burst-mediated spillover inhibition, providing a novel mechanism whereby a putatively relevant network may be influenced by anaesthetics.

Materials and methods

Use of α1100 and δ00 mice as models to investigate the thalamic effects of etomidate

Compensatory adaptations represent a significant caveat to the interpretation of experiments utilizing constitutive knockout mice. Indeed, biochemical and immunohistochemical evidence revealed compensatory up-regulation of α2 and/or α3 subunits in the neocortex and cerebellum of α1100 mice (Sur et al., 2001; Kralic et al., 2002a,b). However, in the thalamus of α1100 mice, the ablated α1 subunits are not replaced by α2 or α3 subunits (Kralic et al., 2006) and α3 expression remains unaltered in the presynaptic nRT (Kralic et al., 2006). Thus, at the functional level, the lack of any clear upregulation of the remaining palette of GABAAR subunits is reflected by a complete lack of miniature inhibitory postsynaptic currents (mIPSCs) in mature ventrobasal (VB) neurons, but unaltered mIPSC properties in the nRT (Peden et al., 2008). Moreover, extrasynaptic GABAARs do not appear to compensate for the lost phasic inhibition, as tonic inhibition is not significantly altered in α1100 VB neurons, at least at the limited age range studied here (Herd et al., 2009). In the δ00 mouse (Mihalek et al., 1999), increased expression of γ2 subunits has been reported (Tretter et al., 2001; Peng et al., 2002). Importantly, however, receptors incorporating α1, α2, α3 or α5 subunits do not appear to substitute for the lost δ-containing receptors (Peng et al., 2002). In agreement, we did not observe any clear alterations in the properties of relay neuron mIPSCs that could compensate for the greatly reduced tonic inhibition in δ00 mice (Herd et al., 2009). Thus, the lack of clear adaptive changes in the properties of phasic and tonic inhibition at the thalamic level in the α1100 and δ00 mice validates their utility as models to investigate the influence of etomidate on relay neuron excitability.

The α1100 and δ00 mice were generated on a mixed C57BL/6J-129SvEv (α1100) or single C57BL/6 (δ00) background at the Merck Research Laboratories at the Neuroscience Research Centre in Harlow and at the University of Pittsburgh, respectively, as described previously (Mihalek et al., 1999; Sur et al., 2001). Brain slices were prepared from the first two generations of α1100 and δ00 and their respective wild-type (WT) mice from breeding pairs derived from the corresponding heterozygous mice bred at the University of Dundee. Measures of inhibition (phasic and tonic) and neuronal excitability [input resistance (IR) and action potential rheobase] did not differ between α1100 and δ00 (data not shown). Results obtained from the respective WT strains were thus pooled.

Slice preparation

Animals were killed by cervical dislocation in accordance with Schedule 1 of the UK Government Animals (Scientific Procedures) Act 1986. Thalamic slices were prepared from mice of either sex (postnatal days 16–24) according to standard protocols (Herd et al., 2009). Dissected brains were sliced horizontally (300–350 μm) in an ice-cold sucrose-based cutting solution, using a Leica VT1000S vibratome (Nussloch, Germany). The slices were incubated at room temperature (20–23 °C) for a minimum of 1 h prior to recording in an oxygenated, extracellular solution (ECS) containing (in mM): 126 NaCl, 2.95 KCl, 26 NaHCO3, 1.25 NaH2PO4, 2 CaCl2, 10 glucose and 2 MgCl2 (pH 7.4; 300–310 mOsm).

Electrophysiology

Electrophysiological recordings were performed from thalamic slices maintained in warmed ECS (30 °C), using an Axopatch 200B amplifier (Molecular Devices, Sunnyvale, CA, USA). Thalamocortical relay neurons of the VB complex were visually identified with an Olympus BX51 microscope (Olympus, Southall, UK), equipped with differential interference contrast/infrared optics and a charge-coupled device camera. Whole-cell voltage- and current-clamp recordings from VB neurons were obtained using patch pipettes prepared from thick-walled borosilicate glass (King Precision Glass, Claremont, CA, USA) using a Narishige PC-10 vertical puller, and had open tip resistances of 4–6 MΩ when filled with a solution containing (in mM): 130 K-glucinate, 2 KCl, 10 NaCl, 10 HEPES (4-(2-hydroxyethyl)-1-piperazineethanesulfonic acid), 0.2 EGTA (ethylenediaminetetraacetic acid), 2 Mg adenosine triphosphate, 0.5 Na guanosine-5′-triphosphate, 10 Tris-phosphocreatine, adjusted to pH 7.2–7.3 with KOH, 280–290 mOsm. Series resistance was intermittently monitored, compensated for (up to 80% for voltage-clamp, 100% for current-clamp) and recordings discarded if values changed by more than 25%, or increased beyond 25 MΩ. Voltage-clamp recordings were performed at a holding potential of ~−50 mV (GABA Erev ~−85 mV), and thus responses mediated by GABAARs were outwardly directed. Inhibitory responses were evoked in VB neurons using a bipolar tungsten electrode (World Precision Instruments, Stevenage, UK), placed within the borders of the nRT. Stimuli were delivered (20-μs duration, 0.1 Hz) using a Digitimer DS3 Isolated Stimulator (Digitimer, Welwyn Garden City, UK), with stimulus intensity adjusted to the minimum value that avoided failures, but still generated IPSC bursts of relatively consistent amplitude (i.e. for each recording, the excitatory IPSC (eIPSC) peak amplitude standard deviation (SD) was <15% of the mean peak amplitude obtained from that cell; WT: 8.5 ± 1.4%, n = 7; α1100: 10.5 ± 1.4%, n = 7; δ00: 8.9 ± 0.8%). At the stimulation intensities utilized, the number of peaks observed in response to each stimulus was relatively consistent from trial to trial (SD = 0.94 and 1.30 for WT and δ00, respectively). Although we find GABAAR activation is minimal under these recording conditions/stimulation parameters (data not shown), we performed the experiments in the presence of CGP 55845 (1 μM) to block GABAARs. Holding currents (Ihold) were
continuously monitored throughout voltage-clamp recordings to determine the effect of etomidate on tonic inhibition. Current-clamp recordings were also performed to determine the effect of burst-mediated inhibition on VB neuron spiking. Thus, evoked inhibitory postsynaptic potentials (eIPSPs) were evoked at regular intervals (0.09–0.11 Hz) during tonic trains of action potentials. Tonic spike trains were generated by depolarizing the recorded neuron to a supra-threshold membrane potential ($V_m$) using depolarizing direct current (DC) injection delivered via the recording electrode. The amplitude of injected DC was carefully adjusted to maintain a tonic spike frequency of ~4–8 Hz. In this paradigm, eIPSPs generated a transient cessation of action potential discharge. eIPSPs were also recorded at or near the resting potential, and in some cells a rebound low-threshold Ca$^{2+}$ potential was observed, which was commonly crowned by a high-frequency spike burst. In such recordings, the latency to rebound, relative to eIPSP onset, was determined for each event before and after etomidate application. VB neuron excitability was also examined in current-clamp experiments by monitoring action potential output in response to a family of current steps. The stimulation protocol consisted of a set of hyperpolarizing and depolarizing current pulses (–200 to +300 pA, 400-ms duration, in increments of 50 pA, every 15 s), delivered via the recording electrode before and after the bath application of etomidate. Current stimulations were delivered from a holding potential of ~66 mV to isolate the tonic spike mode during depolarizing stimulations. The number of action potentials generated in response to positive current steps, and within low-threshold Ca$^{2+}$ current ($I_{Ca}$)-mediated rebound bursts following release from hyperpolarization, were counted for each condition. Membrane IR was calculated according to Ohm’s law following delivery of small (~20 pA, 300 ms) hyperpolarizing current steps from the resting membrane potential. A measured liquid junction potential of 11 mV was corrected offline. Recordings were filtered at 2 or 5 kHz for voltage- and current-clamp recordings, respectively, and acquired directly to PC using a NI-DAQmx analog-to-digital interface (National Instruments UK, Newbury, UK) for subsequent offline analysis.

**Data analysis**

All electrophysiological data were analysed offline using the Strathclyde Electrophysiology Software, WinEDR/WinWCP (J. Dempster, University of Strathclyde, UK) software package. For voltage-clamp experiments, eIPSCs were analysed with respect to peak amplitude, 10–90% rise times and charge transfer. The decay time course following the final peak of multi-peak eIPSCs was adequately described by fitting a double exponential function ($\gamma(t) = A_1e^{(-t/\tau_1)} + A_2e^{(-t/\tau_2)}$) using the least squares method, where $A$ is amplitude, $t$ is time and $\tau$ is the decay time constant. Analysis of the SD of residuals and use of the $t$-test was used to compare goodness of fit, confirming an improved fit of the decay was achieved when employing a double- rather than a mono-exponential function. Thus, a weighted decay time constant ($\tau_w$) was also calculated according to the equation, $\tau_w = \tau_1P_1 + \tau_2P_2$, where $\tau_1$ and $\tau_2$ are the decay time constants of the first and second exponential functions and $P_1$ and $P_2$ are the proportions of the synaptic current decay described by each component. To determine the effect of etomidate on VB tonic inhibition, etomidate-induced outward shifts of the holding current ($I_{Hold}$) were quantified by sampling sequential 51.2-ms epochs for a 2-min period before and after application of etomidate (Belelli et al., 2005). Epochs containing evoked spontaneous synaptic currents were excluded. The mean holding current in each condition was determined from Gaussian fits to the distribution of holding current values sampled from each epoch.

For current-clamp experiments investigating the effect of etomidate on VB neuron excitability, input-output curves were constructed by plotting the number of action potentials generated as a function of stimulus amplitude. To quantify etomidate-induced shifts in input–output curves, data were fit with a Boltzmann sigmoidal curve according to the equation $\gamma(x) = N_{max}/[1 + e^{(X - X_{50})/K} + N_{max}]$, where $N_{max}$ is the number of action potentials generated at the maximum stimulus amplitude (in this case 300 pA), $X$ is the stimulus amplitude, $X_{50}$ is the stimulus amplitude producing half-maximal spike output (hereafter $'E_{A50}'$) and $S$ is the slope factor. For experiments investigating the effect of eIPSPs on tonic spike trains, inter-spike intervals (ISIs) were measured throughout the spike train. For baseline measurements, ISIs were determined for a 5-s period preceding each eIPSP (minimum of three stimulations) to ensure spike rates recovered to stable frequencies following inhibition. The ISI during an IPSP was simply the interval between the final spike prior to delivery of an eIPSP and the first spike following recovery from inhibition. The transient suppression of tonic spike activity induced by an eIPSP was observed as an immediate and obvious increase in the ISI (see Figs 2 and 3).

All results are reported as the arithmetic mean ± standard error of the mean (SEM). Statistical significance of the mean data was assessed with a Student’s $t$-test (paired, or unpaired), one-way analysis of variance (ANOVA) or a two-way mixed-design ANOVA as appropriate, using the SPSS (SPSS Inc., Chicago, IL, USA) software package. Sphericity was assessed with Mauchly’s test, and degrees of freedom were adjusted using a Greenhouse–Geisser correction if sphericity was violated. Post-hoc multiple comparisons were conducted using Tukey’s test. Unless explicitly stated, statistical significance was assessed using Student’s $t$-test. Boltzmann curve fits were performed in Origin v7 (OriginLab, Northampton, MA, USA).

**Salts and drugs**

$R$-(-)-etomidate (supplied by Merck, Glasgow, UK) was prepared as a concentrated aqueous stock and diluted 1000-fold in ECS to a final concentration of 3 μM. CGP 55835 (Abcam, Cambridge, UK) was prepared in dimethyl sulphoxide (DMSO) as a 1000-fold concentrated stock solution and diluted in ECS to the final desired concentration. The final DMSO concentration (0.1%) had no effect on any of the measured parameters (data not shown). Etomidate was applied via the perfusion system (2–4 mL/min) and allowed to infiltrate the slice for a minimum of 10 min. All other salts and drugs were obtained either from Sigma-Aldrich (Poole, UK), Tocris Bioscience (Bristol, UK) or VWR (Lutterworth, UK).

**Results**

**Etomidate prolongs eIPSCs by potentiating synaptic and extrasynaptic GABA$\AA$Rs**

In VB neurons, etomidate prolongs mIPSCs and enhances tonic currents mediated by $\alpha$1$\beta$2$\gamma$- and by $\alpha$4$\beta$2 GABA$\AA$Rs, respectively (Belelli et al., 2005). However, our recent demonstration that thalamic δ-GABA$\AA$Rs may influence the kinetics of action potential-dependent phasic inhibition, particularly in response to spike bursts (Herd et al., 2013), provides an additional route whereby etomidate may influence VB inhibition. Thus, we investigated the effects of etomidate on electrically evoked IPSCs generated in response to extracellular stimulation of the nRT of WT, 11° and 18° mouse brain slices.

In voltage-clamp recordings utilizing K$^+$-gluconate-based pipette solutions (GABA$\AA$R $E_{eq} \approx -85$ mV, $V_h = -50$ mV), extracellular stimulation of the nRT in WT brain slices generated outwardly
eIPSCs were recorded from (Fig. 1F, drug overall effect of etomidate on eIPSC charge transfer relative to WT eGABAAR contribution to eIPSC properties, etomidate (3 neurons (Table 1). Nevertheless, despite the absence of the component normally engaged by neurally released GABA acting on WT synaptic GABAARs (Table 1, Fig. 1B). The slow kinetics of the rising and decaying phase of (Table 2, Fig. 1A and D, n = 7, P < 0.001, and τw (Table 2, Fig. 1A and E, P < 0.001), resulting in a significant enhancement of eIPSC charge transfer (Table 2, Fig. 1A and F, P = 0.001). Consistent with our previous report (Herd et al., 2013), robust residual eIPSCs were recorded from z1000 VB neurons, despite the absence of synaptic GABAARs (Table 1, Fig. 1B). The slow kinetics of the rising and decaying phase of z1000 eIPSCs, together with their sensitivity to modulation by the δ-selective compound DS2 (Wafford et al., 2009; Herd et al., 2013; Jensen et al., 2013), indicate that such residual events are mediated by recruitment of δ-containing extrasynaptic (e)GABAARs in response to nRT spike bursts. Importantly, acute application of etomidate greatly enhanced the peak amplitude (Table 2, Fig. 1B and D, n = 7, P < 0.001) and τw (Table 2, Fig. 1B and E, P < 0.001) of z1000 eIPSCs, leading to a significant increase in their charge transfer (Table 2, Fig. 1F, P < 0.001). However, the overall effect of etomidate on eIPSC charge transfer was reduced for z1000 relative to WT neurons (Fig. 1F, drug × genotype interaction, F2,19 = 7.21, P = 0.005, mixed ANOVA), suggesting that a significant proportion of the effects of etomidate on WT eIPSCs are mediated by synaptic GABAARs. In support, etomidate also enhanced eIPSCs recorded from δ000 neurons (Fig. 1C). In agreement with previous findings (Herd et al., 2013), eIPSCs recorded from δ000 VB neurons exhibited faster decay kinetics, due to the abolished eGABAAR component normally engaged by neuronal released GABA acting on WT neurons (Table 1). Nevertheless, despite the absence of the eGABAAR contribution to eIPSC properties, etomidate (3 μm) significantly increased eIPSC peak amplitude (Table 2, Fig. 1C and D, n = 8, P = 0.003), τw (Table 2, Fig. 1C and E, P < 0.001) and charge transfer (Table 2, Fig. 1F, P = 0.003). However, as observed for z1000 recordings, deletion of the δ subunit significantly reduced the overall effect of etomidate on eIPSC charge transfer relative to WT (Fig. 1F, drug × genotype interaction, F2,19 = 7.21, P = 0.005, mixed ANOVA).

Etomidate prolongs phasic suppression of VB tonic firing and delays post-inhibitory rebound burst firing

During conscious brain states, thalamocortical relay neurons exist in a relatively depolarized state, and respond to incoming excitatory inputs in the tonic ‘relay’ mode (Steriade et al., 1993). Given the shared roles of synaptic and extrasynaptic GABAARs in regulating the kinetics of inhibitory synaptic transmission (see above), we hypothesized that these receptor populations should combine to regulate the effect of etomidate on phasic inhibition of tonic VB spike output.

To explore these questions we performed whole-cell current-clamp recordings of tonic firing from WT, z1000 and δ000 slices. VB neurons were slowly depolarized from the resting membrane potential (Table 3) by DC injection until they discharged trains of tonic spikes, occurring at ~4–8 Hz (mean spike frequency: WT = 5.8 ± 0.3 Hz, n = 13; z1000 = 5.7 ± 0.3 Hz, n = 12; δ000 = 5.5 ± 0.1 Hz, n = 8; F2,32 = 0.35, P = 0.71, one-way ANOVA). We then evoked IPSPs by extracellular stimulation of the nRT (0.09–0.11 Hz stimulation frequency) at regular intervals during the tonic train, and quantified the duration of the eIPSP-induced suppression of tonic firing across strains (i.e., ISI), before and after etomidate (3 μm) (Figs 2 and 3). As illustrated in the representative example in Fig. 2A, stimulation of the nRT in WT brain slices evoked reproducible IPSPs, which transiently suppressed tonic spiking of VB neurons (baseline ISI = 173 ± 8 ms; ISI during IPSP = 1106 ± 118 ms; n = 13, P < 0.001; Fig. 2D). Remarkably, despite the absence of a detectable synaptic receptor population in VB, equivalent stimulation of the nRT in z1000 slices generated eIPSPs that transiently inhibited tonic spiking, albeit to a lesser extent than in WT tissue (baseline ISI = 176 ± 10 ms; ISI during IPSP = 672 ± 67 ms; n = 12, P < 0.001; stimulation × genotype interaction, F2,28 = 7.04, P = 0.003, mixed ANOVA; Fig. 2B and D). Similarly, in δ000 slices, eIPSPs induced a reproducible blockade of VB tonic firing, although the duration of spike suppression was significantly (and similar to z1000) reduced relative to WT recordings (baseline ISI = 180 ± 5 ms; ISI during IPSP = 650 ± 49 ms; n = 8, P < 0.001, stimulation × genotype interaction, F2,28 = 7.04, P = 0.003, mixed ANOVA; Fig. 2C and D), consistent with voltage-clamp experiments demonstrating faster eIPSC decay time in the absence of δ-GABAARs.

Consistent with the robust potentiation of eIPSCs observed in voltage-clamp experiments, etomidate (3 μm) significantly prolonged the duration of tonic firing suppression induced by eIPSPs in WT (ISI during IPSP: control = 1000 ± 148 ms; + etomidate = 1624 ± 161 ms; n = 8, P < 0.001; Fig. 3A and D), z1000 (ISI during IPSP: control = 737 ± 67 ms; + etomidate = 1383 ± 78 ms; n = 9, P < 0.001; Fig. 3B and D) and δ000 brain slices (ISI during IPSP: control = 670 ± 54 ms; + etomidate = 1427 ± 206 ms; n = 6, P < 0.001; Fig. 3C and D). Surprisingly, however, the effect of etomidate on eIPSP-induced suppression of tonic firing (i.e. the difference in ISI during an IPSP before and after etomidate application) was not significantly different between strains (Fig. 3D, F2,20 = 0.56, P = 0.58, mixed ANOVA), despite the ablation of either major subcellular population of GABAARs.

In a subset of these recordings, eIPSPs recorded at or slightly depolarized to the resting membrane potential were of sufficient amplitude to generate a rebound low-threshold Ca2+ potential and associated spike burst at the offset of the eIPSP (von Krosigk et al., 1993). We therefore investigated the effect of etomidate on the post-inhibitory rebound properties by measuring the latency to rebound. Consistent with the prolongation of evoked phasic inhibition, etomidate significantly delayed the timing of rebound bursts in WT recordings (Fig. 4A and C, n = 4, P = 0.009) at similar Vm (pre-eIPSP Vm = −68.3 ± 1.2 and −69.0 ± 1.2 mV for control and etomidate, respectively). Similarly, in δ000 recordings, the latency to post-inhibitory rebound was significantly increased in the presence of etomidate (Fig. 4B and C, n = 5, P < 0.001) at equivalent Vm (pre-eIPSP
TABLE 2. Comparison of the effect of etomidate on eIPSCs recorded from WT, α1<sup>100</sup> and δ<sup>00</sup> mice

<table>
<thead>
<tr>
<th></th>
<th>WT (n = 7)</th>
<th>α1&lt;sup&gt;100&lt;/sup&gt; (n = 7)</th>
<th>δ&lt;sup&gt;00&lt;/sup&gt; (n = 8)</th>
</tr>
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<tbody>
<tr>
<td><strong>Peak amplitude (pA)</strong></td>
<td>599 ± 101</td>
<td>1037 ± 152&lt;sup&gt;***&lt;/sup&gt;</td>
<td>198 ± 31</td>
</tr>
<tr>
<td><strong>τ&lt;sub&gt;w&lt;/sub&gt; (ms)</strong></td>
<td>53 ± 7</td>
<td>144 ± 9&lt;sup&gt;***&lt;/sup&gt;</td>
<td>73 ± 14</td>
</tr>
<tr>
<td><strong>Charge transfer (pC)</strong></td>
<td>40 ± 8</td>
<td>184 ± 32&lt;sup&gt;**&lt;/sup&gt;</td>
<td>17 ± 3</td>
</tr>
</tbody>
</table>

**P < 0.01, ***P < 0.001, paired t-test.**

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European Journal of Neuroscience, 40, 2487–2501
TABLE 3. Comparison of resting membrane potential (V<sub>m</sub>) and input resistance recorded from WT, α<sup>100</sup> and δ<sup>000</sup> mice

<table>
<thead>
<tr>
<th></th>
<th>WT (n = 38)</th>
<th>α&lt;sup&gt;100&lt;/sup&gt; (n = 19)</th>
<th>δ&lt;sup&gt;000&lt;/sup&gt; (n = 27)</th>
</tr>
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<tbody>
<tr>
<td>V&lt;sub&gt;m&lt;/sub&gt; (mV)</td>
<td>−72.5 ± 0.5</td>
<td>−72.0 ± 0.9</td>
<td>−73.5 ± 0.5</td>
</tr>
<tr>
<td>Input resistance (MΩ)</td>
<td>288 ± 13</td>
<td>274 ± 22</td>
<td>328 ± 20</td>
</tr>
</tbody>
</table>

V<sub>m</sub> = −67.8 ± 0.3 and −68.3 ± 0.2 mV for control and etomidate, respectively. eIPSPs recorded from α<sup>100</sup> slices did not elicit rebound bursts, although in all seven recordings, the duration of eIPSPs recorded from equivalent membrane potentials were prolonged (data not shown). These observations imply that both synaptic and extrasynaptic receptor modulation can influence post-inhibitory rebound timing.

Deletion of the α<sub>1</sub> or δ subunit influences the etomidate-induced enhancement of the VB tonic current

While quantifying the effects of etomidate on eIPSCs, we simultaneously monitored the VB holding current in each mouse strain to further understand the contribution of α1β2 and α4βδ GABA<sub>A</sub>Rs

Fig. 2. Synaptic and extrasynaptic GABA<sub>A</sub>Rs influence phasic suppression of VB tonic firing. (A–C) Representative current-clamp recordings illustrating the suppression of tonic action potential discharge induced by evoked IPSPs in WT (A), α<sup>100</sup> (B) and δ<sup>000</sup> (C) thalamic slices. The plots to the right of each trace depict the inter-spike interval (ISI) over the course of the illustrated recordings, and demonstrate the increase in ISI (i.e. spike suppression) during delivery of the eIPSP. Evoked IPSPs were generated by extracellular stimulation of the nRT. (D) Bar graph comparing the effect of eIPSPs on ISI during tonic spike trains recorded from WT (n = 13), α<sup>100</sup> (n = 12) and δ<sup>000</sup> (n = 8) VB neurons. ** p < 0.001 vs. baseline ISI, paired t-test. † P < 0.05 vs. WT, mixed ANOVA, Tukey’s post hoc test.
Fig. 3. Etomidate prolongs phasic suppression of VB tonic firing by potentiating synaptic and extrasynaptic GABA\(_{\text{A}}\)Rs. (A–C) Whole-cell current-clamp recordings illustrating the suppression of VB tonic firing under control conditions (Ai–Ci), and in the presence of 3 \(\mu\)M etomidate (Aii–Ci), in WT (A), \(\alpha^{10/0}\) (B) and \(\delta^{0/0}\) (C) thalamic slices. The plots to the right of each trace depict the inter-spike interval (ISI) during the illustrated tonic spike train, and demonstrate blockade of action potentials during the eIPSP. Note that the ISI during the eIPSP is increased in the presence of etomidate across all mouse strains. (D) Bar graph comparing the effect of eIPSPs on the ISI of tonic spike trains, before and after etomidate, in WT (\(n = 8\)), \(\alpha^{10/0}\) (\(n = 9\)) and \(\delta^{0/0}\) (\(n = 6\)) VB neurons. The bar representations are depicted in the symbol key. \(* * * P < 0.001\) vs. baseline ISI, paired \(t\)-test. \(*** P < 0.001\), ISI during eIPSP after etomidate application vs. control. The increased duration of the eIPSP-induced spike suppression observed in the presence of etomidate is not significantly influenced by mouse genotype (\(P = 0.58\), mixed ANOVA).
recordings (20.7 ± 2.9 pA, n = 9) concurrently with potentiation of residual eIPSCs (Fig. 5C and D), the magnitude of the outward current shift was significantly reduced relative to WT (Fig. 5G, drug × genotype interaction, \( F_{2,21} = 20.11, P < 0.001, \text{mixed ANOVA} \)). In recordings from VB neurons of \( \delta^{\text{lo}} \) mice, bath application of 3 \( \mu \)M etomidate induced only a minimal outward current (5.0 ± 1.0 pA, \( n = 7 \); \( P < 0.001 \) vs. WT; \( P = 0.012 \) vs. \( \delta^{\text{lo}} \), mixed ANOVA, Tukey’s post-hoc test; Fig. 5E and G), while robustly potentiating eIPSCs (Fig. 5F).

**Deletion of \( z1 \) or \( \delta \) subunits attenuates the effect of etomidate on VB neuron excitability**

Given that etomidate modulation of VB tonic inhibition is significantly reduced in \( z1^{\text{lo}} \) and \( \delta^{\text{lo}} \) neurons, such genetic manipulations are thus predicted to influence the effect of the anaesthetic on cellular excitability. To investigate this hypothesis, we examined the effect of etomidate on VB input-output (I-O) relationships generated in response to a family of current steps (−200 to +300 pA, 400 ms duration, 50-pA increments every 15 s). Current steps were delivered while neurons were held at a membrane potential of ~66 mV; at such potentials, \( I_f \) is largely inactivated (Coulter et al., 1989). First, we compared I-O relationships across WT, \( z1^{\text{lo}} \) and \( \delta^{\text{lo}} \) under control conditions to determine if basic electrical excitability was affected by the mutations (Fig. 6). Across all three strains, in accordance with classical TC neuron physiology, hyperpolarizing currents steps of sufficient amplitude generated a sagging, hyperpolarization-activated cation current \((I_{\text{H}})\)-dependent membrane potential response, followed by a rebound low-threshold Ca\(^{2+}\) potential, crowned by a high-frequency spike burst (spikes per burst, WT = 2–10, \( n = 23 \); \( z1^{\text{lo}} = 1–9, n = 12 \); \( \delta^{\text{lo}} = 2–10, n = 22 \)). Although the amplitude of the hyperpolarizing current step significantly influenced the mean number of spikes per rebound burst \((F_{3,162} = 104.80, P < 0.001, \text{mixed ANOVA})\), there was no significant effect of genotype on this relationship \((F_{3,162} = 1.00, P = 0.43, \text{mixed ANOVA}, \text{Fig. 6D})\). Supra-threshold depolarizing current steps of increasing amplitude generated tonic spikes of increasing frequency in VB neurons derived from each mouse strain (Fig. 6A–C and E). Although visual inspection of the resulting I-O curves (Fig. 6E) suggests a modest leftward trend of the relationship towards increased excitability for \( \delta^{\text{lo}} \) relative to WT recordings, this did not reach statistical significance \((F_{3,162} = 1.46, P = 0.20, \text{mixed ANOVA})\). Indeed, estimation of the current amplitude generating half maximal spike output \((\text{EA}_{50})\) from Boltzmann sigmoidal curve fits (see Methods) indicated a small, but non-significant effect of \( \delta \) subunit deletion on the I-O \( \text{EA}_{50} \) (WT = 173 ± 7 pA, \( n = 23 \); \( z1^{\text{lo}} = 168 ± 10 \text{ pA, } n = 12 \); \( \delta^{\text{lo}} = 158 ± 8 \text{ pA, } n = 22 \), \( F_{2,54} = 1.00, P = 0.37, \text{one way ANOVA} \)). Similarly, neither the resting membrane potential nor IR of VB neurons was significantly influenced by deletion of the \( \delta \) subunit, although again a trend towards an increased IR was observed relative to WT (IR, WT = 288 ± 13 mV, \( n = 38 \); \( \delta^{\text{lo}} = 328 ± 20 \text{ mV, } n = 27 \), \( P = 0.1, \text{unpaired t-test, Table 3} \)).

In accordance with the large potentiation of tonic inhibition, bath application of 3 \( \mu \)M etomidate hyperpolarized the resting \( V_m \) reduced membrane IR and exerted a clear reduction in the excitability of WT VB neurons (Table 4, Fig. 7A–D). Thus, etomidate significantly reduced the number of spikes per rebound burst upon release from hyperpolarizing stimulations, although the effect was particularly evident for smaller amplitude current steps (Fig. 7A and B). Similarly, etomidate reduced the tonic spike output, resulting in a clear rightward shift of the I-O relationship of VB neurons (Fig. 7A and C). Indeed, the \( \text{EA}_{50} \) determined from Boltzmann fits to I-O

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**Fig. 4.** Etomidate delays the timing of post-inhibitory rebound burst firing in VB neurons. (A, B) Exemplar traces from a subset of current-clamp recordings obtained from WT (A) and \( \delta^{\text{lo}} \) (B) mice in which rebound burst firing was observed at the offset of eIPSPs. The responses to three consecutive stimuli recorded before (black traces) and after (grey traces) application of 3 \( \mu \)M etomidate are superimposed, illustrating the delay to rebound burst firing in the presence of the anaesthetic. Action potentials are truncated for clarity. (C) Summary bar graph comparing the latency to rebound spiking (black bars) and after etomidate (grey bars) application for WT (\( n = 4 \)) and \( \delta^{\text{lo}} \) (\( n = 5 \)) VB neurons. **\( P < 0.01 \), ***\( P < 0.001 \), paired \( t \)-test.

To the effects of the anaesthetic on VB tonic inhibition. Bath application of 3 \( \mu \)M etomidate induced a clear outward shift in VB holding current (37.2 ± 5.0 pA, \( n = 8 \); Fig. 5A and G), with a time course similar to that observed for potentiation of eIPSCs (Fig. 5B, panels i and ii). In previous work, we found that deletion of \( z1 \)-containing receptors did not significantly influence baseline tonic current amplitudes, consistent with the hypothesis that VB tonic inhibition is mediated predominantly by \( z4 \beta 2\delta \) receptors (Chandra et al., 2006; Herd et al., 2009). Thus, it was surprising that while etomidate application induced a clear outward current in \( z1^{\text{lo}} \) VB recordings (20.7 ± 2.9 pA, \( n = 9 \)) concurrently with potentiation of residual eIPSCs (Fig. 5C and D), the magnitude of the outward current shift was significantly reduced relative to WT (Fig. 5G, drug × genotype interaction, \( F_{2,21} = 20.11, P < 0.001, \text{mixed ANOVA} \)). In recordings from VB neurons of \( \delta^{\text{lo}} \) mice, bath application of 3 \( \mu \)M etomidate induced only a minimal outward current (5.0 ± 1.0 pA, \( n = 7 \); \( P < 0.001 \) vs. WT; \( P = 0.012 \) vs. \( \delta^{\text{lo}} \), mixed ANOVA, Tukey’s post-hoc test; Fig. 5E and G), while robustly potentiating eIPSCs (Fig. 5F).
Fig. 5. Deletion of the α1 or δ subunit reduces the effect of etomidate on VB tonic inhibition. (A, C, E) Representative whole-cell recordings (left) and corresponding all points histograms (right), illustrating the effect of etomidate on the holding current of WT (A), α1°°° (C) and δ°°° (E) VB neurons. Evoked IPSCs were simultaneously monitored throughout the recordings, but have been truncated to emphasize changes in holding current (I_hold). Non-truncated eIPSCs denoted by i (control) and ii (etomidate) are shown in B, D and F for WT, α1°°° and δ°°°, respectively, to confirm the effect of etomidate on eIPSCs. (G) Bar graph comparing the outward current induced by the bath application of etomidate (expressed as a change in I_hold) for WT (black bar, n = 8), α1°°° (dark grey bar, n = 9) and δ°°° (light grey bar, n = 7) VB neurons. *p < 0.05, **p < 0.01, ***p < 0.001, one-way ANOVA, Tukey’s post hoc test.
curves was significantly increased in the presence of the anaesthetic (EA_{50} control = 170 ± 12 pA; + etomidate = 195 ± 12 pA, n = 12, P < 0.001). By contrast, although etomidate also hyperpolarized the resting V_m and decreased the membrane IR of α_{10}/δ_{0} VB neurons (Table 4), the consequent reduction of their excitability was less than for WT neurons (Fig. 7D–F). These findings are consistent with
the reduced enhancement of tonic inhibition by the anaesthetic following α1 subunit deletion. This was particularly evident upon inspection of the I-O relationship generated from depolarizing stimulations, where the rightward shift in the curve induced by etomidate (EA_{et} control = 166 ± 11 pA, + etomidate = 178 ± 13 pA, n = 11, P = 0.03) was significantly reduced relative to WT recordings (Table 5). Deletion of the δ subunit virtually abolished the effect of etomidate on VB burst and tonic firing (Fig. 7G–I), consistent with the limited etomidate-induced outward current observed in δ^{00} voltage-clamp experiments. In addition, for δ^{00} neurons, etomidate had minimal effect on the resting V_m, or on the membrane input resistance (Table 4). Thus, with the exception a small effect at the smallest current step, which was significantly reduced relative to that observed in WT recordings (P = 0.0191, paired t-test), etomidate did not significantly influence rebound intra-burst spike number in response to hyperpolarizing current steps (Fig. 7H). Similarly, for δ^{00} neurons, the I-O curve generated in response to positive current steps remained relatively unaffected by etomidate (EC_{et} control = 163 ± 14 pA, + 3 μm etomidate = 163 ± 15 pA, n = 9, P = 0.95, Fig. 7I; Table 5).

### Discussion

In this study, we investigated the relative contribution of synaptic and extrasynaptic GABA_αRs to the etomidate-induced inhibition of neuronal activity in a network putatively involved in general anaesthetic mechanisms. Specifically, deletion of the δ subunit reduced the effect of etomidate on VB tonic inhibition. Consequently, the ability of etomidate to inhibit burst and tonic spiking was virtually abolished in δ^{00} mice, suggesting that etomidate facilitation of tonic inhibition exerts a dominant effect on TC cell output. Unexpectedly, deletion of the α1 subunit also reduced the impact of etomidate on VB tonic inhibition and the relay neuron I-O relationship, thus implicating the α1-GABA_αR population during anaesthetic actions in thalamus. Importantly, etomidate also greatly prolonged the duration of nRT burst-mediated phasic inhibition, by potentiating both synaptic and extrasynaptic GABA_αRs. Thus, further to conventional phasic and tonic inhibition, burst-mediated spill-over may represent a novel mechanism whereby anaesthetics can modulate circuit function.

While anaesthetics probably exert their behavioural effects by altering neuronal activity across multiple brain regions (Franks, 2008), human neuro-imaging studies have identified a consistent thalamic deactivation during anaesthetic-induced sedation and/or loss of consciousness (Fiset et al., 1999; Alkire et al., 2000; White & Alkire, 2003; Hofbauer et al., 2004; Gili et al., 2013). Such studies are consistent with the canonical view of the thalamus as a sensory ‘gate’ that governs the transmission of multi-modal information to the cortex (Steriade, 2003). In vivo, anaesthetics may modulate thalamic activity directly by influencing the intrinsic excitability of resident thalamic neurons or indirectly by altering the activity of extrinsic excitatory inputs (e.g. descending cortico-thalamic, or ascending reticular activating system inputs). Indeed, single-unit recordings from cats showed a reduction of the spontaneous tonic firing rate of TC neurons by etomidate, alongside similar decreases in the neocortex and reticular formation (Andrada et al., 2012). Computer simulations utilizing thalamocortical network models have also provided important insights, revealing that elevated inhibition of cortical and thalamic neurons imposes a slowed, hyper-synchronous rhythm upon the thalamocortical loop that is reminiscent of electroencephalography dynamics during anaesthetized states (Talavera et al., 2009; Ching et al., 2010; Purdon et al., 2013). Collectively, such studies favour a co-ordinated action of anaesthetics across key interconnected networks, rather than a specific localized effect. Nevertheless, our data support the hypothesis that etomidate mediates at least a component of its actions by a direct reduction of TC neuron output, resulting from enhanced synaptic and extrasynaptic GABA_αR function. In agreement, propofol and the volatile agent isoflurane also inhibit relay neuron tonic spiking in a manner that is at least partially GABA_αR-dependent (Ying & Goldstein, 2005a,b; Jia et al., 2008).

That some general anaesthetics exert their spectrum of behavioural effects via a selective interaction with the GABA_αR is well established (Hales & Lambert, 1991; Franks & Lieb, 1994). In support, mutant mice harbouring point-mutated GABA_αRs engineered to be etomidate-insensitive (β2N265S; Belelli et al., 1997; Reynolds et al., 2003) or propofol- and etomidate-insensitive (β3N265S; Pistas et al., 1999; Jurd et al., 2003) display dramatically altered behavioural responses to these anaesthetics, revealing that specific anaesthetic behaviours can be assigned to individual GABA_αR populations. Specifically, the sedative and immobilizing effects of both etomidate and propofol are mediated by β2- and β3-containing receptors, respectively, whereas both receptor populations contribute to the hypnotic component (Jurd et al., 2003; Reynolds et al., 2003). However, these studies did not address which β2- or β3-expressing neuronal populations may be involved in the respective behaviours. Thus, it is noteworthy that the effects of etomidate on relay cell phasic and tonic currents are greatly involved in β2N265S slices (Belelli et al., 2005). In light of the present evidence for a reduced etomidate-induced inhibition of relay neuron spiking in α1^{00} and δ^{00} mice, such studies implicate synaptic (α1β2γ2) and extrasynaptic (α4β2δ2) GABA_αRs as important molecular targets for the thalamic actions of this anaesthetic.

The effect of etomidate on VB tonic inhibition was virtually abolished in δ^{00} mice, leading to a reduced inhibitory effect on burst and tonic spiking. Such observations are consistent with the tonic conductance generating an inhibitory charge which greatly outweighs that produced by brief, spontaneous phasic inhibition. In agreement, computational studies utilizing a thalamocortical network model suggest that enhanced tonic inhibition is critical for the etomidade-induced depression of thalamic firing (Talavera et al., 2009). Thus, the significant effect of the α1 subunit deletion (albeit lesser than for the δ^{00}) was unexpected. One potential interpretation

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**Table 4. Comparison of the effect of etomidate on the resting membrane potential (V_m) and input resistance of WT, α1^{00} and δ^{00} VB neurons**

<table>
<thead>
<tr>
<th></th>
<th>WT (n = 19)</th>
<th>α1^{00} (n = 12)</th>
<th>δ^{00} (n = 11)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Control</td>
<td>3 μm etomidate</td>
<td>Control</td>
</tr>
<tr>
<td>V_m (mV)</td>
<td>−71.7 ± 0.8</td>
<td>−73.2 ± 0.9***</td>
<td>−71.7 ± 0.8</td>
</tr>
<tr>
<td>Input resistance</td>
<td>258 ± 14</td>
<td>193 ± 11***</td>
<td>285 ± 26</td>
</tr>
</tbody>
</table>

*P < 0.05, **P < 0.01, ***P < 0.001, paired t-test.
suggests that a subpopulation of δ-GABA<sub>Rs</sub> may contain the α<sub>1</sub> subunit instead of α<sub>4</sub>. Indeed, α<sub>1</sub> can form a functional partnership with the δ subunit in some hippocampal interneurons (Glykys et al., 2007). However, while α<sub>1</sub> subunit deletion abolishes spontaneous phasic inhibition in VB (Peden et al., 2008), we did not detect a significant reduction of baseline tonic inhibition, or the current induced by the δ-selective agonist THIP (4,5,6,7-tetrahydroisoxazolo[5,4-c]pyridin-3-ol), which does not discriminate between α<sub>1</sub>δ and

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**Fig. 7.** The effect of etomidate on VB neuron excitability is attenuated by deletion of the α<sub>1</sub> subunit and abolished by deletion of the δ subunit. (A) Representative whole-cell current-clamp recording from a WT VB neuron, illustrating superimposed voltage responses to the indicated current steps (middle) under control conditions (top) and in the presence of 3 μM etomidate (bottom). (B) Graph comparing the intra-burst spike number observed on rebound from a range of hyperpolarizing current steps (−200 to −50 pA), before (black symbols) and after (grey symbols) etomidate (n = 12). (C) Graph comparing the number of spikes generated in response to depolarizing current steps of increasing amplitude (50–300 pA) under control conditions (black symbols), and in the presence of etomidate (grey symbols). (D–F) Effect of etomidate on the excitability of α<sub>1</sub><sup>0/0</sup> VB neurons (n = 11). Figure details are as described for A–C. (G–I) Effect of etomidate on the excitability of δ<sup>0/0</sup> VB neurons (n = 9). Figure details are as described for A–C. The input-output relationships determined for each condition are fitted with a Boltzmann curve. *P < 0.05, **P < 0.01, ***P < 0.001, paired t-test. Etom, etomidate.
TABLE 5. Comparison of the effect of etomidate on the input-output relationship of WT, z1\(^{10/0}\) and \(\delta^{10/0}\) VB neurons

<table>
<thead>
<tr>
<th>Condition</th>
<th>EA50 (pA)</th>
</tr>
</thead>
<tbody>
<tr>
<td>WT (n = 12)</td>
<td>25 ± 4</td>
</tr>
<tr>
<td>z1(^{10/0}) (n = 11)</td>
<td>12 ± 5*</td>
</tr>
<tr>
<td>(\delta^{10/0}) (n = 19)</td>
<td>0 ± 6**</td>
</tr>
</tbody>
</table>

*P < 0.05, **P < 0.01, vs. WT, one-way ANOVA, Tukey’s post-hoc test.

\(\alpha\)β\(\delta\) receptors (B. Ebert, personal communication; Herd et al., 2009), suggesting that \(\alpha\)β\(\delta\) receptors are not abundantly expressed in VB neurons. Alternatively, the loss of on-going spontaneous phasic inhibition, which is robustly prolonged by etomidate in WT neurons, may partially account for the reduced effect of the anaesthetic on spike discharge for z1\(^{10/0}\) relay neurons. Additionally, a substantial pool of z1\(\beta\)2 receptors may exist extrasynaptically (Kasugai et al., 2010). While this receptor population does not appear to contribute to baseline tonic inhibition, etomidate, which increases GABA\(\alpha\)R channel open probability in response to low GABA concentrations (Yang & Uchida, 1996), may enhance the activation of extrasynaptic z1\(\beta\)2 receptors by ambient GABA. Finally, we cannot exclude that direct activation of z1\(\beta\)2 receptors by etomidate (Hill-Venning et al., 1997) may influence VB neuron excitability.

Using paired nRT–VB recordings, we recently reported that peri/extrasynaptic \(\alpha\)β\(\delta\) receptors are recruited during nRT burst firing, leading to a substantial prolongation of burst-mediated phasic inhibition (Herd et al., 2013). In support of this, despite the absence of synaptic GABA\(\alpha\)ARs, we observed a residual eIPSC in z1\(^{10/0}\) VB neurons, which was greatly potentiated by the \(\delta\)-selective positive allosteric modulator DS2 (Herd et al., 2013). Here, we observed a similar potentiating of the residual eIPSC upon application of etomidate to z1\(^{10/0}\) slices. Consequently, the remarkable ability of the residual eIPSPs to suppress tonic spiking of z1\(^{10/0}\) neurons was greatly prolonged by etomidate. In z2\(^{10/0}\) and \(\delta^{10/0}\) mice, eIPSCs are shorter due to the abolished spill-over component, and are insensitive to DS2, consistent with the loss of \(\alpha\)β\(\delta\) receptors (Herd et al., 2013). In contrast to DS2, in \(\delta^{10/0}\) recordings, etomidate significantly prolonged eIPSCs and therefore the eIPSP-induced suppression of tonic spiking. Curiously, for the latter paradigm, the duration of spike suppression induced by the eIPSP in the presence of etomidate was not significantly different from WT in either z1\(^{10/0}\) or \(\delta^{10/0}\) neurons (Fig. 3D), despite a clear effect of both gene deletions under control conditions (Fig. 2D). In current-clamp recordings the duration of IPSPs is determined partially by the kinetic properties of GABA\(\alpha\)ARs, but also by the ensuing activation of voltage-dependent conductances (e.g. I\(\text{H}\) and I\(\text{T}\)), which may act to truncate the IPSP time course (Thomson et al., 2000). Thus, we speculate that the duration of the IPSP may be maximized in the presence of the anaesthetic and that this ‘ceiling effect’ may be achieved via the potentiation of either synaptic or extrasynaptic GABA\(\alpha\)ARs in the absence of the complementary receptor population. Nevertheless, the results suggest that etomidate facilitates burst-mediated inhibition via a combined action at synaptic and peri/extrasynaptic GABA\(\alpha\)ARs, leading to an amplified inhibitory effect on VB tonic firing.

As noted above, the strength of synaptic inhibition arising from nRT neurons interacts with intrinsic voltage-dependent conductances to control the timing of post-inhibitory rebound bursts, and thus the period and synchrony of thalamus-dependent oscillations, which prevail in the unconscious brain (Huguenard & McCormick, 2007). In addition to an enhanced tonic inhibition, which induces a membrane hyperpolarization that is likely to favour burst firing (Cope et al., 2005), etomidate also delayed the timing of post-inhibitory rebound burst firing by prolonging the IPSP. Given the importance of burst firing during thalamocortical rhythmosgenesis, the etomidate-induced increase of IPSP duration may therefore contribute to the slowed oscillatory activity of the anaesthetized brain.

Our in vitro data, together with a computational study (Talavera et al., 2009), suggest that the etomidate-induced inhibition of TC output is dependent on synaptic and extrasynaptic GABA\(\alpha\)ARs. Thus, it is surprising that deletion of z1, z4 or \(\delta\) subunits exerts little or no influence on the sedative or hypnotic actions of etomidate (Mihalek et al., 1999; Kralic et al., 2003; Iyer et al., 2013). However, the interpretation of behavioural data obtained from mice harbouring global GABA\(\alpha\)R subunit deletions is complicated by the non-uniform up-regulation of alternative GABA\(\alpha\)R isoforms following subunit loss (Sur et al., 2001; Kralic et al., 2002a,b; Peng et al., 2002). This caveat is exemplified by studies investigating the subunit-dependence of benzodiazepine-induced behaviours; mice expressing a benzodiazepine-insensitive z1 subunit (H101R) are resistant to the sedative effects of diazepam (Rudolph et al., 1999; McKernan et al., 2000), yet z1\(^{10/0}\) mice exhibit enhanced diazepam-induced sedation (Kralic et al., 2002b). Thus, caution is warranted when comparing results from a slice preparation demonstrating minimal functional GABA\(\alpha\)R compensation to behavioural studies underpinned by complex interactions between multiple brain networks subject to varying degrees of compensation. We note that an as yet unidentified neuronal adaptation (e.g. as described by Brickley et al., 2001) is likely to compensate for the loss of tonic or phasic inhibition in relay neurons, as demonstrated by the lack of any significant alteration in membrane IR or the I-O relationship in the mutant mice. Crucially, however, such adaptations do not appear to include upregulation of the remaining palette of GABA\(\alpha\)AR subunits (Kralic et al., 2006; Peden et al., 2008; Herd et al., 2009).

In summary, we have shown that etomidate reduces the electrical excitability of thalamocortical relay neurons via a specific interaction with synaptic and extrasynaptic GABA\(\alpha\)ARs. Given the established role of the thalamus in arbitrating conscious state transitions, our results support the putative role of these distinct thalamic GABA\(\alpha\)AR populations in general anaesthetic actions. Finally, we suggest that burst-mediated spill-over, a kinetically intermediate form of inhibition that bridges the kinetic extremes of fast phasic and slow tonic inhibition, represents a novel means by which GABA-modulatory anaesthetics may influence the relay of sensory information from thalamus to cortex.

Acknowledgements

This work was supported by an Epilepsy Research UK fellowship (M.B.H., F1001), the BBSRC (D.B. & J.J.L., Grant CS09923), Tenevus Scotland (M.B.H., D.B. & J.J.L.) and the Anonymous Trust (M.B.H., D.B. & J.J.L.). We thank Dr Bjarte D. Landbeck (Uppsala) for sharing unpublished data, and Dr Thomas Rosahl (Merk) and Professor Gregg Homanics (University of Pittsburgh) for provision of the z1\(^{10/0}\) and \(\delta^{10/0}\) mice, respectively. The authors declare no competing financial interests.

Abbreviations

ANOVA, analysis of variance; DMSO, dimethyl sulphoxide; DS2, \(\delta\)-selective compound 2; EA50, excitatory current required for half maximal spike frequency; ECS, extracellular solution; GABA\(\alpha\)R, extrasynaptic \(\gamma\)-aminobutyric acid, type A receptor; eIPSC, evoked IPSC; eIPSP, evoked IPSP; \(E_{\text{rev}}\), reversal potential; GABA, \(\gamma\)-aminobutyric acid; GABA\(\alpha\)R, \(\gamma\)-aminobutyric acid, type A receptor; GABA\(\beta\), \(\gamma\)-aminobutyric acid, type B receptor; I\(\text{H}\), current mediated by hypopolarization-activated, cyclic nucleotide-gated cation channels; I-O, input-output relationship; IPSC, inhibitory postsynaptic
current; IPSP, inhibitory postsynaptic potential; IR, membrane input resistance; ITA, inter-spike interval; IV, current mediated by low threshold T-type Ca2+-channel (IPSC, nRT, nRT). The subunit nomenclature contains: TC, thalamocortical; VB, ventrobasal complex; vRC, membrane potential; WT, wild-type; z1α300, z1 subunit knock-out mouse; z1α-GABAR, z1 subunit-containing GABAAR receptor; β326M, asparagine to methionine mutation at position 265 of the GABAAR receptor β3 subunit; β226MS, asparagine to serine mutation at position 265 of the GABAAR receptor β2 subunit; δ260, δ subunit knock-out mouse; δ-GABAR, δ subunit-containing GABAAR receptor; τm, weighted decay time constant.

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