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TASK-3 Two-Pore Domain Potassium Channels Enable Sustained High-Frequency Firing in Cerebellar Granule Neurons

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The ability of neurons, such as cerebellar granule neurons (CGNs), to fire action potentials (APs) at high frequencies during sustained depolarization is usually explained in relation to the functional properties of voltage-gated ion channels. Two-pore domain potassium (K2P) channels are considered to simply hyperpolarize the resting membrane potential (RMP) by increasing the potassium permeability of the membrane. However, we find that CGNs lacking the TASK-3 type K2P channel exhibit marked accommodation of action potential firing. The accommodation phenotype was not associated with any change in the functional properties of the underlying voltage-gated sodium channels, nor could it be explained by the more depolarized RMP that resulted from TASK-3 channel deletion. A functional rescue, involving the introduction of a nonlinear leak conductance with a dynamic current clamp, was able to restore wild-type firing properties to adult TASK-3 knock-out CGNs. Thus, in addition to the accepted role of TASK-3 channels in limiting neuronal excitability, by increasing the resting potassium conductance TASK-3 channels also increase excitation by supporting high-frequency firing once AP threshold is reached.

Key words: accommodation; action potential; cerebellum; excitability; granule cell; potassium channels

Introduction

Two-pore domain potassium (K2P) channels hyperpolarize the resting membrane potential (RMP) by increasing potassium permeability (Millar et al., 2000; Talley et al., 2000), reducing cell excitability by moving the potential away from action potential (AP) threshold (Bean, 2007). The increased input conductance associated with K2P channel opening will further reduce excitability by attenuating excitatory drive (Goldstein et al., 2001; Patel and Honore, 2001). Many neuromodulators promote closure of K2P channels (Talley et al., 2000; Chemin et al., 2003; Heurteaux et al., 2006; Meuth et al., 2006; Mathie, 2007), and the conventional view would be that K2P closure would increase neuronal excitability.

Here, we explore the possibility that K2P channel opening does more than simply limit excitability. Membrane hyperpolarization could reduce steady-state inactivation of voltage-gated sodium channels (VGSCs), making sustained AP firing more likely once threshold is reached. An increased input conductance will also reduce depolarization block of VGSCs and aid recovery from inactivation because of a faster membrane time constant. In addition, under physiological conditions, the current–voltage relationship of a K2P-mediated conductance deviates significantly from a linear ohmic response (Goldman, 1943; Cole, 1949; Hodgkin and Katz, 1949). This open rectification will result in voltage dependence of the input conductance additional to that attributed to conventional voltage-gated potassium channels (Brickley et al., 2001; Cathala et al., 2003). These additional biophysical properties could promote, rather than inhibit, AP firing.

The potassium leak conductance present in adult cerebellar granule neurons (CGNs) correlates with the strong expression of five K2P genes: TWIK-1, TREK-2c, THIK-2, TASK-1, and TASK-3 (Millar et al., 2000; Brickley et al., 2001; Karschin et al., 2001; Rajan et al., 2001; Talley et al., 2001; Gu et al., 2002; Han et al., 2002; Aller et al., 2005). Recombinant TASK-1 and TASK-3 channels combine as homodimers and heterodimers, and it is likely that TASK-1/TASK-3 heterodimers are a prominent combination in adult CGNs (Aller et al., 2005). However, because of dif-
ferences in single-channel properties (Han et al., 2002; Kang et al., 2004), it is likely that, at physiological pH, TASK-3-containing channels (homodimers or heterodimers) will increase potassium permeability to a greater extent than TASK-1 homodimeric channels. We previously showed that genetic deletion of TASK-1 does not alter the leak conductance, the RMP, or AP firing in adult CGNs (Aller et al., 2005). In the current study, we examined the consequence of genetically removing TASK-3 expression on the excitability of adult CGNs. In contrast to the situation for TASK-1 knock-out (KO) neurons (Aller et al., 2005; Meuth et al., 2006), a significant depolarization of the RMP was observed in CGNs lacking TASK-3. In addition, a clear increase in the degree of AP accommodation was apparent both at threshold and suprathreshold levels of depolarization. To directly test the hypothesis that loss of TASK-3 channels was responsible for this phenotype, we reintroduced a nonlinear leak conductance into adult TASK-3 KO CGNs using a dynamic current-clamp procedure. This restored the ability of TASK-3 KO CGNs to fire at high frequencies during sustained depolarization.

**Materials and Methods**

*Generation of TASK-3 knock-out mice.* As deduced by aligning the TASK-3 coding sequences (Chapman et al., 2000; Kim et al., 2000; Rajan et al., 2000; Vega-Saenz De Miera et al., 2001) against the mouse genomic sequence, the mouse TASK-3 gene (alternative gene names, Kv9.1, KCNK9, Kt3.2) lies on chromosome 15E1 (gene reference number, ENSMUSG00000036760; see http://www.ensembl.org/Mus_musculus/ geneview?gene=ENSMUG00000036760) (supplemental Fig. S2A, available at www.jneurosci.org as supplemental material).

We obtained the mouse TASK-3 gene by screening a mouse 129 BAC library (BAC mouse ES Release A, BAC-4921, Library Screening Services; Genome Systems, St. Louis, MO) with a 1 kb intronic probe (derived from the 5′ end of intron 1 in the TASK-3 gene). This intron 1 probe was generated by PCR on C57BL/6 mouse genomic DNA using the following primers: 5′-CCA TCA TCG TCA TCA CAA CTA TCG-3′ (sense) and 5′-CTC ACA GAT CCA AAG AGA TGG ACC-3′ (antisense). The targeting vector backbone was constructed by placing an 8 kb EcoRI fragment of the TASK-3 gene, containing exon 1, into the pLitmus38 vector (New England Biolabs, Beverly, MA) (supplemental Fig. S1A, available at www.jneurosci.org as supplemental material). The Sall site in the pLitmus38 polynuker was first converted to Nool, permitting later linearization of the targeting vector (supplemental Fig. S2A, available at www.jneurosci.org as supplemental material). An 5′-EcoRV-lloxP site (5′-GATATCCCAATACTGCTTAGTACCATATTAGGATTAT-3′) was then inserted into the unique BstI site, 560 bp upstream of the initiator ATG codon of exon 1 in the TASK-3 gene. The EcoRV site allows detection of homologous recombination. The final targeting vector was obtained by insertion of a 5′-Xhol-frt-neomycin resistance-frt-loxp-Sal-I cassette (Cope et al., 2004) into an engineered Sall site, 275 bp downstream of the 3′ end of exon 1 of the TASK-3 gene (note: the Sall site for receiving the neomycin resistance cassette was introduced by first ligating Sall linkers into an unique AflI site in intron 1) (supplemental Fig. S1B, available at www.jneurosci.org as supplemental material). A plasmid was selected with bothloxP sites in the same orientation. The targeting vector was linearized with Nool and electroporated into mouse embryonic stem (ES) cells (strain 129/Sv, B6, gift from G. Schuetz Laboratory, Deutsches Krebsforschungszentrum, Heidelberg, Germany). The cells were grown on mouse fibroblast cells in medium containing G418. Genomic DNA from ES cell colonies was screened by Southern blotting and hybridization; colony genomic DNA was digested with Nool, and hybridized with a specific 5′ external probe, giving a 5.8 kb band for the wild-type allele and a 4.2 kb band for the targeted allele (supplemental Fig. S1D,E, available at www.jneurosci.org as supplemental material); the 5′ external probe was generated by PCR on C57BL/6 mouse tail genomic DNA using the following primers: 5′-CCA TGG CTA GGG ACA GAG AAT A3′ (sense) and 5′-TCT CAT TTC CCC TCT CCA GCA GTA-3′ (antisense). The presence of the 5′ loxP site was confirmed by digestion with EcoRV and hybridizing with the same 5′ external probe, which gives a 15 kb wild-type band and a 5.6 kb targeted allele band (supplemental Fig. S1D,E, available at www.jneurosci.org as supplemental material). We undertook three electroporations and screenings of ES cell colonies; in the first two screens (of 850 colonies in total screened), no correctly targeted colonies were identified; in the third attempt, of 700 colonies screened, one was positive with the 5′ external probe (Nool) digest and also contained the 5′ loxP site (EcoRV digestion) (supplemental Fig. S1D, available at www.jneurosci.org as supplemental material). Cells from this targeted stem cell colony were injected (by F. Zimmermann, Heidelberg University, Heidelberg, Germany) into C57BL/6 blastocysts; male chimeras were bred with C57BL/6 wild-type mice; germline transmission of the targeted TASK-3 allele was confirmed by Southern blots of tail DNA digested with Nool (which marks the neo gene) and EcoRV (which marks the 5′ loxP site) respectively, hybridized with the 5′ external probe (supplemental Fig. S1D, available at www.jneurosci.org as supplemental material).

The subsequent generations of TASK-3 knock-out mice have been maintained as heterozygotes on a C57BL/6 background, and can be genotyped by PCR on genomic DNA with primers flanking the 5′ loxP site (supplemental Fig. S1G, available at www.jneurosci.org as supplemental material), giving a 400 bp band for the knock-out and 350 bp band for the wild type. For PCR genotyping the primers were as follows: 5′-TCTGTCGGGCAGGATGACAGTCG-3′ and 5′-TTTCCGTTGGGGCGACTCGCCG-3′. More information on the TASK-3 knock-out construction is given in the supplemental material (available at www.jneurosci.org).

Quantification of K<sub>γ</sub> expression in wild-type and TASK-3 knock-out brains by real-time PCR. Mouse brain K<sub>γ</sub> and GABA<sub>γ</sub> receptor α6 subunit gene expression were assayed by real-time PCR as described by Aller et al. (2005).

**In situ hybridization.** In situ hybridization with 35S-labeled oligonucleotide probes was as described by Wisden and Morris (1994) and Aller et al. (2005). Nonspecific labeling of the sections was assessed by hybridizing labeled oligonucleotide in the presence of a 100-fold excess of unlabeled oligonucleotide. Images were generated by exposure to Biomax MR (Kodak, Rochester, NY) x-ray film, and then scanned and processed in Adobe Photoshop. The TASK-1 and TASK-3 oligonucleotides were identical to those used by us previously (Aller et al., 2005).

**Recombinant expression.** The tsA-201 cells were transiently transfected using the calcium phosphate method. Briefly, 0.5 or 1 µg of cDNA expression vector encoding mouse TASK-3 (Aller et al., 2005) was added to each 15 mm well, and 1 µg of a plasmid encoding the cDNA of green fluorescent protein was included to identify cells expressing channels. After a 24 h incubation period, the cells were rinsed with saline and fresh growth medium was added to the wells. The cells were incubated at 37°C with 5% CO<sub>2</sub> for 24–60 h before whole-cell voltage-clamp electrophysiological recording measurements were made.

**Slice preparation.** Adult wild-type (81 ± 6 d of age) and TASK-3 KO mice (85 ± 10 d of age) were used for preparation of parasagittal cerebellar slices (250 µm thick). Slices were transferred to a recording chamber and constantly perfused (2 ml/min) with standard recording solution that contained the following (in mM): 125 NaCl, 2.5 KCl, 1 CaCl<sub>2</sub>, 2 MgCl<sub>2</sub>, 26 NaHCO<sub>3</sub>, 1.25 NaH<sub>2</sub>PO<sub>4</sub>, 25 glucose, pH 7.4, when bubbled with 95% O<sub>2</sub> and 5% CO<sub>2</sub>. Whole-cell recordings from CGNs were made using either an Axopatch 200A or 700B amplifier (Molecular Devices, Foster City, CA) run via a National Instruments digitization board (PCI-6052E; National Instruments, Austin, TX) on a dedicated PC. All experiments were performed at room temperature (22–23°C) with 1 mM kynurenic acid and 30 µM picrotoxin added to the external solution to block glutamatergic and GABAergic synaptic transmission. The pipette solution contained the following (in mM): 120 KCl, 2.5 KCl, 1 CaCl<sub>2</sub>, 2 MgCl<sub>2</sub>, 26 NaHCO<sub>3</sub>, 1.25 NaH<sub>2</sub>PO<sub>4</sub>, 25 glucose, pH 7.4, adjusted to pH 7.3 with KOH. Data acquisition and analysis was performed using WINWCP (version 3.3.3) kindly provided by John Dempster (University of Strathclyde, Glasgow, UK).

**Recording procedures and analysis.** Voltage recordings were passed through a 2 kHz, −3 dB, eighth-pole Bessel filter before digitization at a rate of 20 kHz. The RMP of CGNs was measured during the first minute of whole-cell recording. AP waveform analysis was performed on the first
5–10 APs elicited at current injection threshold for each cell. Average waveforms were constructed from these initial APs by aligning on the AP threshold. AP threshold was estimated using phase plane plots (see Fig. 5B) of the first derivative as a function of voltage, thus enabling a simple estimate of the initial voltage inflection (Bean, 2007). The phase plane plots also enabled a fit-independent method for estimating the maximum rate of rise and fall for the AP upstroke and repolarizing phase. AP width was estimated at the AP half-height, defined as the distance between the AP threshold and the AP overshoot. The accommodation index was calculated during a suprathreshold depolarizing current pulse of 20 pA above threshold. AP frequency was calculated during the first and last 20 ms epoch of a current pulse lasting either 200 or 300 ms. APs were considered to have failed if their peak amplitude fell <80% of the AP peak amplitude recorded at threshold. Current recordings were digitized at 50 kHz when measuring fast transients and 10 kHz for slower conduc-
tance changes such as \( I_{	ext{Na}} \). In all cases, a 2 kHz, −3 dB, eight-pole Bessel filter was applied before digitization. Particular attention was taken to eliminate possible capacitive and resistive artifacts when action potential waveforms were being used to control the command voltage during voltage-clamp experiments. Action potential waveforms were first recorded from a wild-type adult CGN in response to a series of 400 ms suprathreshold depolarizing current injections. These particular recording were made at a total gain of 50 mV/mV and were saved onto a DAT recorder at a bandwidth of 50 kHz after being passed through a 5 kHz, −3 dB, eight-pole Bessel filter. Voltage records could then be played back into the external voltage command of the patch-clamp amplifier. Subsequent voltage-clamp recordings were only used for these experiments if the seal resistance was >10 GΩ and the initial series resistance was <25 MΩ. Eighty percent series resistance compensation was then applied using a lag time of 3 μs. A series of control experiments were performed on a model cell to ascertain the validity of this approach.

When examining sodium channel activation, a protocol consisting of a 50 ms conditioning step at −100 mV was followed by a series of 25 ms steps from −105 to +80 mV in 5 mV increments. For inactivation, the initial 100 ms step was delivered at command potentials ranging from −100 to −15 mV in 5 mV increments before the voltage was stepped to a test command at +20 mV for 25 ms. Data from these experiments were converted into conductance plots and fitted with a Boltzmann function (\( y = \frac{y_{\text{max}}}{1 + \exp([V - V_{0.5}]/k]) \)) to calculate \( V_{0.5} \) and \( k \) values. When examining the recovery from inactivation, a 50 ms conditioning step to −100 mV was followed by a 25 ms step to +20 mV. The membrane was then returned to −100 mV for a maximum of 600 ms in 0.5 ms increments to allow recovery from inactivation, before the membrane was stepped back to +20 mV for 25 ms. The recovery from inactivation plots were fitted with exponential functions (\( I = [0.25 \exp(-t/\tau) + 0.75] \)). The protocols used to examine voltage-gated potassium channel properties consisted of a family of 200 ms duration voltage steps from a control potential of −70 mV to test voltages ranging from −120 to +100 mV. These experiments were performed in the presence of 500 nM TTX, and 1 mM tetraethylammonium (TEA) was applied when the properties of tran-
sient A type potassium channels were examined. Once again, data from these experiments were converted into conductance plots and fitted with a Boltzmann function to calculate \( V_{0.5} \) and \( k \) values. In all experiments, a 1 s delay was given between each protocol during which the CGN was held at −70 mV. All data fitting was performed in Origin 6.0 (Microcal, Northampton, MA). CGN input conductance was measured from a 1000 ms duration 10 mV hyperpolarizing voltage step from which we also calculated the input capacitance. The membrane time constant (\( \tau_m \)) was then calculated from the calculated input conductance and capacitance.

**Dynamic current-clamp procedure.** The dynamic current clamp was implemented in a Labview 7.2 (National Instruments) software environment following adaptation of the original G-clamp software (Kullmann et al., 2004). A dynamically variable current was injected into the soma of neurons using a patch-clamp amplifier (interfaced to a slave computer via a National Instruments PCI-6052E DAQ board). To make use of the computers full bandwidth, the slave computer was running embedded software controlled by a second master PC via an ethernet connection. The constant field or Goldman–Hodgkin–Katz (GHK) model provides us with a quantitative description of a nonlinear \( I-V \) relationship of the type produced by \( K_{\text{sp}} \) channel populations. From the GHK equation, we know that the current carried by potassium ions is equal to the perme-
ability of that ion multiplied by a nonlinear function of voltage as follows:

\[
I = \frac{PF \cdot V_m}{RT} \cdot \left( \frac{c_m - c_{out} \exp \left( \frac{-V_{0.5}}{RT} \right)}{1 - \exp \left( \frac{-V_{0.5}}{RT} \right)} \right).
\]

Abbreviations are as follows: \( P \), permeability (centimetres per second); \( z \), valence of charge; \( F \), Faraday’s constant (coulombs per mole); \( T \), temper-
ature (kelvin); \( R \), gas law constant [Joules per (mole kelvin)]; \( |c_{\text{in}}|/|c_{\text{out}}| \), concentration of ions inside/outside cell. However, we need to describe this relationship in terms of conductance not permeability. Given that the slope conductance \( (g_m) \) at \( V \rightarrow \infty \) is given by the following:

\[
g_m = \frac{PF \cdot c_m}{RT},
\]

substituting \( g_m \) into the GHK equation,

\[
I = \frac{PF \cdot V_m}{RT} \cdot \left( \frac{c_m - c_{out} \exp \left( \frac{-V_{0.5}}{RT} \right)}{1 - \exp \left( \frac{-V_{0.5}}{RT} \right)} \right).
\]

The resting conductance of this relationship could then be modified during the experiment to observe the impact of this conductance on the firing properties of CGNs. Reliable performance of the system requires minimal variation, or jitter, from cycle to cycle so that the requested arithmetical operations are achieved during each real-time loop. Opera-
tionally, the maximum jitter reported by the system when implementing this simple GHK-type leak was 2 ± 0.05 μs, well within the sample interval of 50 μs used during these experiments.

All statistical tests were performed using STATISTICA 5.1 (StatSoft, Tulsa, OK) and considered significant at \( p < 0.05 \). A Shapiro–Wilk test was used to determine whether measures were normally distributed, and differences between groups were examined using the Student’s t test or the Mann–Whitney U test when distributions were not normal. Any possible correlation between variables was examined using a Spearman rank correlation test.

**Results**

As seen by in situ hybridization on wild-type adult mouse brains (Fig. 1A), high levels of TASK-3 mRNA were observed in the CGN layer of the cerebellum, the colliculi, hippocampal CA1 and dentate gyrus, anterior dorsal thalamus, the medial habenule, cholinergic cells in the caudate–putamen (Berg and Bayliss, 2007) and many cells in neocortex and olfactory bulb (Aller et al., 2005; Linden et al., 2006). The expression of the TASK-3 gene in mouse brain is identical with that in rat (Karschin et al., 2001; Talley et al., 2001; Vega-Saenz De Miera et al., 2001). On the other hand, although TASK-1 appears to be expressed at slightly higher levels than TASK-3 in CGNs (Fig. 1A, C), in the rest of the brain TASK-1 expression is more limited (Fig. 1C). Because of a lack of reliable antibodies, it is not yet known whether these mRNA levels correlate with relative TASK-1 and -3 protein levels.

We generated mice with a disabled TASK-3 gene by combined homologous and insertional recombination, so that exon 1 and intron 1 of the gene were crippled (supplemental Fig. S1, available at www.jneurosci.org as supplemental material). The first exon normally encodes the first one-third of the TASK-3 protein up to the first potassium selectivity filter residue, glycine Y55. Both in situ hybridization with a TASK-3 exon1-specific probe (Fig. 1B) and real-time PCR (Fig. 1E) demonstrated that TASK-3 transcripts were absent in the TASK-3 knock-out brain. A commercial polyclonal antibody (anti-K2P9.1; Alomone, Jerusalem, Israel) exhib-
Loss of TASK-3 channels reduces the potassium permeability of adult CGNs

In the acute slice preparation, the steady-state outward current (Fig. 1 F), recorded at −20 mV, was significantly reduced in adult TASK-3 KO CGNs (41.4 ± 2.3 pA/pF, n = 55 in the wild type; vs 22.9 ± 2.9 pA/pF, n = 16 in TASK-3 KO). The outward rectification of the potassium leak current is characteristic of a Kv_{leak} channel population recorded with asymmetrical potassium ion concentrations, and the magnitude of this conductance across the entire voltage range examined is clearly reduced in the TASK-3 KO CGNs (Fig. 1 G). The ensuing reduction in the potassium permeability of the membrane resulted in a significant −10 mV depolarization of the RMP from −80.3 ± 1.1 mV (n = 55) in wild type to −70.9 ± 1.8 mV (n = 27) in TASK-3 KO CGNs. This is consistent with a movement away from the potassium equilibrium potential (−104 mV) when TASK-3 channels are removed from CGNs. Indeed, it can be seen in Figure 1 G that the intersection between the wild-type and TASK-3 KO average I–V relationships is close to this potassium equilibrium potential. A reduced leak conductance was not observed in TASK-1 knockout mice, and, therefore, we observed no change in the RMP (Aller et al., 2005). This highlights the relative importance of TASK-3-containing Kv_{leak} channels in maintaining the hyperpolarized RMP of adult CGNs. A reduced potassium leak conductance in TASK-3 KO CGNs (compared with wild type) was also apparent in the presence of TTX, TEA, and 4-AP (see Fig. 7 A). The non-inactivating property of the Kv_{leak} leak conductance allowed us to quantify the reduction in this conductance in the absence of potassium channel blockers. This was fortunate given that pharmacological isolation of the potassium leak conductance is complicated by the fact that recombinant TASK channels do exhibit some TEA sensitivity (IC_{50} ~ 10 μM) (S. G. Brickley and A. Mathie, unpublished observations).
AP accommodation is more pronounced in TASK-3 KO CGNs

As shown in Figure 2A, the current required to reach AP threshold was clearly reduced in TASK-3 KO CGNs, reflecting the more depolarized RMP and decreased input conductance of these cells. In the examples shown, only 12 pA of current was required to reach AP threshold in the TASK-3 KO compared with 24 pA in the wild-type CGN. The AP height recorded was also clearly reduced in the TASK-3 KO example (see below). The increased excitability of TASK-3 KO CGNs was further apparent in the subthreshold membrane behavior illustrated in Figure 2B, both for the individual examples and the CGN populations as a whole (Fig. 2B). Once AP threshold was reached, high-frequency AP firing was still observed in TASK-3 KO adult CGNs. The most striking phenotype observed in TASK-3 KO CGNs was the marked AP accommodation during suprathreshold current injections. In all TASK-3 KO CGNs examined, the AP peak amplitude was heavily attenuated during sustained depolarization, and in 17 of 27 CGNs this resulted in complete AP failure by the end of a suprathreshold depolarizing current pulse. Correlated with the reduction in AP height, it is also clear that in TASK-3 KO CGNs the afterhyperpolarization (AHP) is also reduced in magnitude during suprathreshold depolarization (Figs. 2A, 3C, 4A). Therefore, during sustained depolarization, the smaller AHP will hinder recovery of VGSCs from inactivation, perhaps trapping channels in a slow inactivated state. The difference in the input–output curves constructed from the wild-type and TASK-3 KO data sets is explained by the altered level of accommodation (Fig. 2C). In the TASK-3 KO data set, accommodation at suprathreshold depolarization results in a decline in AP instantaneous frequency. No decline in output frequency was observed in the wild-type CGN population at the current intensities examined.

The level of accommodation was quantified both at threshold depolarizations and during suprathreshold current injections. When examining AP accommodation at threshold, AP properties were compared at current injections that elicited the first burst of three or more spikes. In this way, the degree of AP accommodation could be quantified at similar membrane voltages (−46.1 ± 1.0 mV, n = 44 in wild-type; vs −44.9 ± 0.9 mV, n = 27 in TASK-3 KO CGNs). As shown in Figure 3A, the degree of AP accommodation at threshold depolarizations was greater in TASK-3 KO CGNs (Fig. 3B). In wild-type CGNs, no change in AP height was observed during the first three spikes with initial average amplitudes of 80.8 ± 1.8 mV and final amplitudes of 79.2 ± 1.8 mV. In contrast, the average AP height measured in the TASK-3 KO CGNs reduced significantly from 55.4 ± 2.8 to 49.9 ± 3.5 mV. At suprathreshold depolarizations, the AP frequency was compared during the first and last 20 ms epoch of the current pulse (30 pA above threshold). As expected from previous studies (D’Angelo et al., 1998, 2001; Brickley et al., 2001; Chaderton et al., 2004), the resulting accommodation index (ratio of the instantaneous frequency recorded at the end and start of a suprathreshold current pulse) in wild-type CGNs was high (0.97 ± 0.05; n = 18), and consistent with our previous findings (Aller et al., 2005), this measure was no different in TASK-1 KO CGNs (0.94 ± 0.07; n = 9) (Fig. 3D). However, in TASK-3 KO CGNs, this measure had significantly reduced to 0.20 ± 0.06 (n = 26).

Depolarized RMP is not responsible for increased accommodation in TASK-3 KO CGNs

VGSC availability for activation will be influenced by the initial RMP, and this could influence the degree of accommodation observed in TASK-3 KO CGNs. As seen from Figure 3D, there was considerable overlap in the recorded RMPs of wild-type and TASK-3 KO CGNs. However, TASK-3 KO CGNs with relatively hyperpolarized RMPs still exhibit marked accommodation; conversely, accommodation was not observed in the more depolarized wild-type CGNs. To directly examine how AP generation was influenced by the RMP of CGNs, we injected steady-state current to impose an RMP on wild-type and TASK-3 KO CGNs. It is unclear to what extent this procedure would uniformly influence the entire CGN membrane, but, as shown in Figure 4A and C, in TASK-3 KO CGNs, hyperpolarizing the RMP did not prevent accommodation (n = 7). Conversely, in wild-type CGNs, accommodation was only observed at imposed RMPs more depolarized than −60 mV (Fig. 4B, D). In the wild-type example shown, a CGN was chosen that exhibited a similar RMP to the TASK-3 KO CGN (around −70 mV). The same current injection that resulted in AP accommodation in the TASK-3 KO CGN elicited continuous AP firing in this wild-type example. Steady-state depolarization of this wild-type CGN to around −40 mV is expected to reduce the availability of VGSCs for activation, and this would be consistent with the absence of accommodation in this example. In TASK-3 KO CGNs, the availability of VGSCs for activation is not reduced significantly by similar levels of depolarization (Figs. 3C, 4A). This result is consistent with the notion that the accommodation phenotype observed in TASK-3 KO CGNs is due to the presence of TASK-3 channels.
mV resulted in tonic AP firing, and accommodation was superimposed on subsequent current injection pulses. Therefore, it is possible to induce AP accommodation in wild-type CGNs by steady-state depolarization, but this only occurred at voltages near AP threshold. Collectively, these data suggest that steady-state inactivation of VGSCs was not responsible for the accommodation observed in the TASK-3 KO CGNs.

Changes in action potential shape in TASK-3 KO adult CGNs
As shown in Figure 5A, the AP threshold in TASK-3 KO adult CGNs was similar to that we previously reported for adult CGNs (Cathala et al., 2003) and no different to that calculated for the wild type. In the current study, we used phase plane plots (Fig. 5B) to estimate the AP threshold in wild-type (−39.4 ± 1.1 mV; n = 49) and TASK-3 KO (−40.6 ± 0.8 mV; n = 29) adult CGNs. As shown in Figure 5C, the AP height [calculated according to Bean (2007)] had significantly reduced from 106.4 ± 1.6 mV (n = 49) in the wild type to 84.3 ± 3.3 mV (n = 29) in TASK-3 KO CGNs reflecting a reduction in the AP overshoot (36.5 ± 1.5 mV in wild type vs 14.2 ± 2.7 mV in the TASK-3 KO) with little change in the AHP (−69.9 ± 0.9 mV in wild type vs −70.0 ± 1.0 mV in the TASK-3 KO). Gaussian fits to histograms constructed from AP half-width estimates (Fig. 5D) indicated a significantly broader AP in TASK-3 KO adult CGNs, with an average half-width of 624 ± 61 μs in the TASK-3 KO compared with 435 ± 21 μs in wild-type CGNs. This broader AP reflected a significantly slower rate of AP rise during the upstroke: 250.4 ± 10.0 mV·ms⁻¹ in wild type compared with only 159.9 ± 18.1 mV·ms⁻¹ in the TASK-3 KO. The repolarizing phase of the AP was also slower: −220.4 ± 9.5 mV·ms⁻¹ in wild-type compared with −166.3 ± 21.2 mV·ms⁻¹ in TASK-3 KO adult CGNs. From these data, it is clear that the rising phase of the AP is more affected in the TASK-3 KO than the falling phase (Fig. 5E). This is to be expected, given that voltage-gated potassium channels do not contribute to the rising phase of the AP (Bean, 2007). In contrast, Kᵥ channel opening is expected to influence the input conductance during the AP upstroke in a manner not predicted of standard voltage-gated potassium channels. During the repolarizing phase of the AP, voltage-gated potassium channels will make a significant contribution to the membrane time constant, therefore minimizing the impact of the reduced Kᵥ-mediated leak conductance in TASK-3 KO adult CGNs.

We also examined AP broadening during repetitive firing in the two strains. APs generated from adult CGNs during threshold current injection (Fig. 5F) were analyzed in a total of 43 wild-type and 26 TASK-3 KO CGNs. As shown in Figure 5G, in wild-type CGNs, there was little change in AP width at any instantaneous frequency, but in TASK-3 KO recordings the AP width was significantly broadened at all frequencies. It was also clear that the average instantaneous frequency had significantly increased from 46 ± 4.5 Hz (n = 422) in wild-type to 84.5 ± 3.8 Hz (n = 277) in TASK-3 KO CGNs. The mechanisms responsible for this apparent increase in instantaneous AP frequency in the TASK-3 KO are unclear at present and require additional examination. However, a reduction in the Kᵥ-mediated leak conductance can explain the spike broadening apparent at all these instantaneous frequencies, because a slower rising AP will occur regardless of instantaneous frequency in the TASK-3 KO CGNs.

Changes in action potential shape cannot be explained by differences in voltage-gated channels
It could also be that the functional properties of the underlying voltage-gated channels have been altered in the TASK-3 KO mice. Conductance changes resulting from VGSC activation were directly examined in adult CGNs using standard voltage-clamp protocols (Fig. 6). These experiments indicated no significant difference in the properties of VGSCs when comparing wild-type and TASK-3 KO adult CGNs. For example, the peak sodium conductance (Nᵥ) in wild-type CGNs was 15.9 ± 3.3 nS (n = 6) compared with 14.5 ± 2.2 nS (n = 6) in TASK-3 KO CGNs. As shown in Figure 6, A and B, the 10–90% rise time of the peak sodium conductance measured after a step from −100 to 0 mV was also not altered in the TASK-3 KO adult CGNs (180 ± 20 μs,
only reduces the AP frequency at imposed RMPs more depolarized than -50 mV. C, Quantification of the relationship between RMP and the normalized AP frequency in TASK-3 KO adult CGNs. At all imposed RMPs the output of the neurons declines at high stimulus intensities because of the accommodation phenotype. D, Quantification of the relationship between RMP and the normalized AP frequency in wild-type adult CGNs. AP accommodation only reduces the AP frequency at imposed RMPs more depolarized than -50 mV. Error bars indicate SEM.

Figure 4. AP accommodation in TASK-3 KO CGNs is not attributable to the depolarized RMP of TASK-3 KO CGNs. A, Example of voltage responses from an adult CGN recorded from a TASK-3 KO mouse that exhibits considerable AP accommodation at different imposed RMPs. B, Example of a wild-type CGN that only exhibits AP accommodation at depolarized potentials more positive than -50 mV. C, Quantification of the relationship between RMP and the normalized AP frequency in TASK-3 KO adult CGNs. At all imposed RMPs the output of the neurons declines at high stimulus intensities because of the accommodation phenotype. D, Quantification of the relationship between RMP and the normalized AP frequency in wild-type adult CGNs. AP accommodation only reduces the AP frequency at imposed RMPs more depolarized than -50 mV. Error bars indicate SEM.

n = 6 in wild type; vs 192 ± 11 μs, n = 6 in TASK-3 KO). Furthermore, the midpoint for activation (-45.9 ± 1.8 vs -46.4 ± 1.8 mV) and inactivation (-44.8 ± 1.4 vs -45.6 ± 2.7 mV) for Na+/K+ type conductance were no different as was the time course for recovery from inactivation (1.7 ± 0.5 vs 1.5 ± 0.4 ms). These data would indicate that slowing of the AP upstroke is not attributable to changes in VGSCs but reflects the slower membrane time constant that would result from a reduction in the CGN input conductance on removal of a K_{aP}-mediated leak conductance.

In the presence of 500 nM TTX, voltage-step protocols were also used to elicit the delayed rectifier type conductance (K_v) and the transient A-type potassium conductance (K_a) in adult CGNs. As shown in Figure 6E, the K_v-type conductance was isolated using 1 mM TEA and the K_{aP} conductance was estimated in the presence of TTX, TEA, and 4-AP as illustrated in Figure 7A. The K_v-like conductance was estimated from the steady-state outward current elicited during voltage-step protocols once the contribution from the TEA-insensitive steady-state component (K_{aP} conductance) was subtracted. For the K_v-like conductance (Fig. 6D), the steady-state outward current measured during voltage-step protocols exhibited a similar midpoint for activation in wild type (10.5 ± 4.3 mV; n = 18) and TASK-3 KO (11.0 ± 3.0 mV; n = 18). The maximum conductance of this steady-state component was not altered in TASK-3 KO CGNs (12.1 ± 1.7 nS in wild type vs 11.2 ± 0.5 nS in TASK-3 KO). The peak of the transient K_v-type conductance was then examined in the presence of 1 mM TEA (Fig. 6E), and there was no significant difference in the maximum conductance (7.3 ± 1.7 nS, n = 3 in wild-type; vs 7.4 ± 2.0 nS, n = 5 in TASK-3 KO CGNs), or the midpoint for activation (6.7 ± 9.8 mV compared with 5.6 ± 3.5 mV). Next, we compared the Na_v, K_v, K_{aP}, and K_{aP} conductance present at 0 mV (Fig. 6F, G) using the average activation curves shown in Figure 6, C–E, and the open rectification fits of the TEA-insensitive conductance shown in Figure 7A. From this analysis, we would predict that during the repolarizing phase of the AP the K_v- and K_{aP}-type conductance will contribute the largest proportion of the total potassium conductance. However, during the AP upstroke, the K_{aP} component is the only potassium conductance active, and, as discussed previously, this could explain why we see a larger change in the maximum rate of rise in the TASK-3 KO data set.

**TASK-3 channels contribute to excitability changes during action potential generation**

To directly examine the contribution a TASK-3 conductance makes to the input conductance during AP generation, we performed AP clamp experiments during which the command voltage was controlled by previously recorded AP waveforms (Fig. 7A). In addition to the glutamate and GABAA receptor antagonists, these experiments were performed in the presence of 10 mM TEA, 5 mM 4-AP, and 500 nM TTX. Figure 7A shows that, during high-frequency firing, the input conductance contains a voltage-dependent component involving rapid activation/deactivation. The hysteresis observed in the wild-type CGN current–voltage behavior was not evident in TASK-3 KO CGNs. Therefore, the input conductance at -20 mV was significantly reduced from 5.0 ± 0.5 nS (n = 34) in the wild-type to 3.3 ± 0.3 nS in the TASK-3 KO CGNs (n = 16). The very fast activation/deactivation kinetics are reminiscent of K_v3 channels (Rudy and McBain, 2001), but K_v3 channels have very positive activation thresholds with V_{0.5} for activation of +10 to +20 mV (for review, see Rudy and McBain, 2001) and, in any case, would be completely blocked during these experiments (see above). Therefore, we studied recombinant mouse TASK-3 channels to establish whether their behavior could indeed explain these voltage properties seen in native CGN membranes. Mouse TASK-3 channels exhibited similar voltage-dependent activation/deactivation to wild-type CGNs when the command voltage was controlled by AP waveforms (Fig. 7B). The hysteresis that characterized wild-type CGN current–voltage behavior was also observed in recombinant TASK-3 channels. Therefore, the voltage dependence of TASK-3 channels contributes to the increased input conductance of adult wild-type CGNs observed in our experiments. Previous studies have shown fast activation kinetics for TASK-3 channels (time constant, ~2–10 ms) (Rajan et al., 2000; Meadows and Randall, 2001), with an apparently instantaneous deactivation of...
TASK-3 that could explain the hysteresis loop signature. The impact of TASK-3 channel expression on the voltage dependence of the input conductance is further illustrated by comparing the CGN membrane time constant ($\tau_m$) at specific membrane voltages. In wild-type CGNs, the $\tau_m$ was 7.6 ± 1.2 ms ($n = 14$) at a holding potential of −70 mV, but when the membrane was depolarized to −20 mV the input conductance increased, resulting in a $\tau_m$ of 0.5 ± 0.1 ms. This effect was significantly attenuated in TASK-3 KO CGNs ($n = 16$) and the $\tau_m$ was slower both at −70 mV ($\tau_m = 9.4 ± 0.01$ ms) and −20 mV (1.0 ± 0.1 ms).

**Action potential accommodation in adult CGN TASK-3 KOs can be rescued by a nonlinear potassium leak conductance**

TASK-3 gene expression in wild-type CGNs increases throughout development (Zanzouri et al., 2006), but TASK-3 KO CGNs develop with no TASK-3 protein. To rule out the possibility that the electrophysiological mutant phenotype we have described in adult TASK-3 KO CGNs had a developmental origin involving some form of compensation, and to further highlight the role of TASK-3 channels in influencing AP firing, we restored to adult TASK-3 KO CGNs the nonlinear behavior of the input conductance using a dynamic current-clamp approach (Kullmann et al., 2004; Akemann and Knöpfel, 2006). The firing properties of TASK-3 KO CGNs were then examined in the presence of a nonlinear leak conductance. The nonlinear conductance was implemented using the Goldman–Hodgkin–Katz equation with a limiting conductance of 5 nS to approximate the input conductance of wild-type CGNs during AP firing (Fig. 7A). A significant hyperpolarization of 18.1 ± 4.4 mV ($n = 7$) was observed in response to the 5 nS nonlinear potassium-like leak, and therefore, the current required to reach AP threshold was significantly increased from 23.4 ± 2.5 to 28.3 ± 3.1 pA. As shown in Figure 8A, at threshold depolarizations the degree of AP accommodation was modest (height of the third AP was reduced by 5.4 ± 2.8%), and in all CGNs examined this accommodation phenotype was no longer apparent in the presence of a 5 nS nonlinear leak (Fig. 8B). Addition of this nonlinear leak conductance in TASK-3 KO CGNs also resulted in a significant increase in the AP height and a reduction in the AP width (Fig. 8C). Five of the seven CGNs used in these experiments exhibited complete AP failure by the end of a 200 ms suprathreshold depolarization, and all CGNs exhibited accommodation (accommodation index, 0.19 ± 0.12; $n = 7$). However, in the presence of a 5 nS nonlinear leak conductance, the ability of TASK-3 KO CGNs to fire APs at high frequencies during sustained depolarization was restored (Fig. 8D). Therefore, at suprathreshold depolarizations, the average accommodation index in TASK-3 KO CGNs was restored to a wild-type value of 1.02 ± 0.14 ($n = 7$) after insertion of a 5 nS nonlinear leak conductance. The speed of the dynamic clamp is highlighted in Figure 8E in which we plot the altered current injection ($I_{\text{com}}$) in response to the recorded change in membrane voltage ($V_m$). The speed of the dynamic current clamp is sufficient to respond during the fast APs that occur in adult CGNs. However, this speed is only attainable when implementing a simple GHK-type leak. When we attempted to introduce additional
Another series of voltage-step protocols (gray traces) were applied to examine the voltage dependence of sodium channel activation (NaT).

Current traces of the type shown in Figure 7. A Boltzmann function was fitted to the conductance data for all cells examined to estimate both the peak conductance and rate of activation. A Boltzmann function was fitted to the peak conductance data for all cells examined to estimate both the peak conductance and rate of activation. A series of voltage-step protocols (gray traces) were applied to examine the voltage dependence of sodium channel activation (NaT).

A Boltzmann function was fitted to the peak conductance data for all cells examined to estimate both the $V_{0.5}$ and rate of inactivation. $V_{0.5}$ and rate of inactivation. $V_{0.5}$ and rate of inactivation.

Functional properties of voltage-gated channels in adult CGNs. A, Current traces showing the rising phase of a voltage-gated sodium conductance in response to a maximal activation elicited after a voltage step from $-100$ mV (gray lines) recorded from a wild-type (black trace) and a TASK-3 KO (gray trace) adult CGN. B, Plot of 10–90% rise times calculated from current traces of the type shown in A from wild-type (black circles; $n = 6$) and TASK-3 KO (gray circles; $n = 6$) adult CGNs. C, A series of voltage-step protocols (gray traces) were applied to examine the voltage dependence of sodium channel activation (NaT). A Boltzmann function was fitted to the peak conductance data for all cells examined to estimate both the $V_{0.5}$ and rate of activation. D, Another series of voltage-step protocols (gray traces) examined the voltage dependence of activation for the sustained outward potassium currents (Kv) recorded in the presence of 500 nM TTX, and once again a Boltzmann function was fitted to the conductance data to estimate the $V_{0.5}$ and rate of inactivation. E, Another series of voltage-step protocols (gray traces) examined the voltage dependence of activation for the transient potassium currents (KA) recorded in the presence of 500 nM TTX and 1 mM TEA, and once again a Boltzmann function was fitted to the conductance data to estimate the $V_{0.5}$ and rate of inactivation. F, Bar graph illustrating the relative contribution of the $K_D$, $K_T$, $K_{Na}$, and $K_{SP}$ conductance in wild-type CGNs. The conductance of the voltage-gated currents was estimated at 0 mV using the Boltzmann functions in A–C. The $K_D$ component represented the TEA-sensitive component of the steady-state outward current, whereas the $K_{SP}$ component was estimated from the TEA-insensitive outward potassium currents shown in Figure 7. G. Same conventions as D but for TASK-3 KO CGNs. Note that the only component that has significantly changed is the $K_{SP}$-like TEA-insensitive conductance reflecting the loss of TASK-3 channels.

Discussion

Kv channels can make a significant contribution to the resting potassium permeability of neuronal membranes (Millar et al., 2000; Talley et al., 2000; Heurteaux et al., 2006). This becomes particularly apparent when G-protein activation results in their closure and the RMP is depolarized (Goldstein et al., 2001; Patel and Honore, 2001; Mathie, 2007). In this study, we demonstrated that, in adult CGNs, TASK-3-containing channels do hyperpolarize the RMP leading to reduced excitability at rest, but in addition this TASK-3-mediated conductance also becomes permissive for AP firing once the resting potassium leak has been overcome. We hypothesize that the increased potassium conductance provided by TASK-3 channels reduces the degree of depolarization block that VGSCs experience during excitatory drive. In addition, it is also possible that the voltage-dependent increase in input conductance through TASK-3 channel opening can facilitate the removal of VGSCs from inactivation by reducing the membrane time constant. This represents an additional consequence of TASK-3 channel expression that has not been examined in previous studies.

TASK-1 removal did not alter the resting potassium permeability of adult CGNs and certainly did not alter any aspect of AP firing (Aller et al., 2005) (Fig. 3D). Nevertheless, there was an alteration in the modulation of the leak conductance in TASK-1 KO CGNs by pH, zinc, and ruthenium red consistent with the loss of TASK-1 (Aller et al., 2005). In this previous study, we postulated that TASK-1/TASK-3 heterodimers were replaced by TASK-3 homodimers in the TASK-1 KO CGNs. We suggested that the increased open probability of the TASK-3 homodimers compared with the wild-type heterodimer population (cf. Han et al., 2002) explained the normalization of the resting potassium permeability (Aller et al., 2005). The depolarized RMP of TASK-3 KO CGNs is consistent with this argument because any remaining TASK-1 homodimers would exhibit a low open probability relative to wild-type TASK-1/TASK-3 heterodimers. However, we cannot rule out that, in TASK-3 KO CGNs, TASK-1 protein is degraded, although TASK-1 mRNA is unaltered (cf. Jones et al. (1997) for an analogous situation with GABA$_A$ receptor subunits, in which, in the GABA$_A$ receptor $a6$ subunit knock-out, the $\delta$ subunit protein is degraded because it is an obligatory partner for $a6$, but $\delta$ mRNA is still present at wild-type levels). It is also conceivable that, in adult CGNs at least, the TASK-1 protein serves as a modulatory or accessory subunit for TASK-3. Although there are few mouse brain regions in which TASK-1 expression predominates over TASK-3 as assessed by RNA levels (Linden et al., 2006), the majority of nuclei appear to have higher TASK-3 expression than TASK-1 (Fig. 1) [Linden et al. (2006), their Fig. 1; Aller et al.
At present, we know nothing about the assembly and subcellular targeting of TASK-1/TASK-3 channels, and our understanding of the cellular interactions that occur between these subunits will require the availability of good antibodies.

A surprising electrophysiological phenotype observed in this study is that AP accommodation is more pronounced in adult CGNs recorded from TASK-3 KO mice (Fig. 2). This was most apparent at suprathreshold levels of depolarization when complete AP failure was commonly observed by the end of a current pulse (Fig. 3). During suprathreshold current injection in TASK-3 KO recordings, the reduced input conductance of CGNs results in a greater level of depolarization leading to increased VGSC inactivation. This highlights the physiological importance of TASK-3-mediated conductance.
of a TASK-3-mediated conductance in limiting voltage-dependent inactivation of VGSCs. The accommodation phenotype is not greatly influenced by steady-state inactivation of VGSCs (Fig. 4) and does not appear to involve biophysical alterations in the VGSCs themselves (Fig. 6). We postulate that the ability of a TASK-mediated conductance to limit AP accommodation at threshold voltages could relate to the nonlinear behavior of the membrane. The increased input conductance that occurs during the AP will reduce the membrane time constant, leading to faster recovery of VGSCs from inactivation. Changes in the membrane time constant also explain the briefer AP width in wild-type CGNs (Fig. 5). This is most notable during the AP upstroke when conventional voltage-gated potassium channels have little influence, but the increased membrane conductance provided by TASK channels contributes significantly to the membrane time constant.

We previously suggested that open rectification of a TASK-mediated conductance contributes to the voltage dependence of the input conductance described in adult CGNs (Brickley et al., 2001; Cathala et al., 2003; Aller et al., 2005). In support of this argument, the dynamic current-clamp experiments demonstrated that a GHK-type nonlinear conductance on its own was sufficient to restore wild-type firing to CGNs recorded from TASK-3 KO mice both at threshold (Fig. 8A) and suprathreshold (Fig. 8D) levels of depolarization. Insertion of a nonlinear leak conductance also altered AP width and height in the TASK-3 KO CGN recordings (Fig. 8C), consistent with our postulated role of TASK-3 channels in the wild type. However, as shown during AP injection experiments (Fig. 7), we also saw some hysteresis of the TASK-3-mediated conductance between the rising and falling phase of the AP waveform, consistent with a modest voltage dependence of channel opening for TASK-3 (Rajan et al., 2000; Meadows and Randall, 2001). Addition of these voltage-dependent parameters was not possible using our dynamic current-clamp configuration (because of insufficient processing power), but we would predict that these parameters would further accentuate the ability of a TASK-mediated conductance to support sustained AP firing.

The mossy fiber excitatory input to CGNs can maintain transmitter release at frequencies of up to 300 Hz (Saviane and Silver, 2006), and adult CGNs respond to continued depolarization with nonaccommodating AP firing of >200 Hz, both in vitro (Brickley et al., 2001) and in vivo (Chadderton et al., 2004; Jorntell and Ekerot, 2006). Based on our findings, CGNs lacking TASK-3 will be compromised in their ability to support high-frequency firing during prolonged mossy fiber excitation in vivo. However, because TASK-3 KO mice have no ataxic phenotype, it is likely that there has been some resetting of the system at the neuronal network level. This comes as no surprise: the cerebellum is remarkably robust to many disruptions that were anticipated to compromise motor function. For example, genetically deleting extrasynaptic GABA_A receptors that give rise to tonic inhibition in CGNs produces no motor phenotype (Jones et al., 1997; Brickley et al., 2001). Similarly, only mild motor impairments are found when the Na channel gene Scn8a (Nav1.6) is deleted selectively from CGNs (Levin et al., 2006); and even selective loss of synaptic GABA_A receptors from Purkinje cells, so that feedforward inhibition is removed, does not result in ataxia (Wulff et al., 2007). The ability of the cerebellum to compensate for chronic removal of a specific conductance is further demonstrated by the study by Watanabe et al. (1998) in which genetic ablation of Golgi cell inhibitory input to CGNs initially resulted in a motor defect that then reduced in severity over time. In contrast, we would predict that a selective chemical antagonist (not yet existing) of TASK-3 channels would immediately compromise motor performance (cf. Wulff and Wisden, 2005; Wulff et al., 2007). Nevertheless, at the cellular level, the genetic approach has given an informative insight into TASK-3 channel function, highlighting some unappreciated contributions that this conductance makes to the biophysical properties of CGNs.

This ability of a potassium conductance to allow high-frequency firing is usually associated with K_Ca channels (Brew and Forsythe, 1995) (for review, see Rudy and McBain, 2001; Bean, 2007). Our dynamic current-clamp experiments demonstrate that a TASK-like conductance can facilitate AP firing similarly to that observed when a more complicated K_Ca-like conductance was reintroduced into fast spiking interneurons (Lien and Jonas, 2003). This additional feature of a TASK-3 conductance highlights how regulation of TASK channels, such as through the activation of G-protein-coupled receptors, could allow neuronal excitability to be fine-tuned because these channels both select and sustain appropriate synaptic inputs. So, the hyperpolarized RMP and increased input conductance at rest will filter excitatory drive (by making EPSPs smaller and briefer) (cf. Cathala et al., 2003), increasing the requirement for closer timing of synaptic inputs before AP threshold is reached. However, once AP threshold is reached, the TASK-3 conductance will contribute to sustained AP firing by reducing the degree of sodium channel inactivation.

Because the TASK-3 gene expresses in other neurons that require sustained high-frequency AP firing (e.g., parvalbumin-positive hippocampal interneurons) (Torborg et al., 2006), it could be that the ability of a TASK-mediated conductance to sustain high-frequency firing is more general. Indeed, a potassium leak conductance with a pharmacological profile reminiscent of TASK-3 channels occurs in the myelinated axons of motor neurons in Xenopus laevis (Koh et al., 1992). It is possible that a TASK-3 conductance in combination with VGSCs is sufficient to support high-frequency firing, and it would be interesting to directly examine the contribution these potassium leak channels make to AP generation in mammalian myelinated axons.

References


Chapman CG, Meadows HJ, Godden RJ, Campbell DA, Duckworth M, Kelless RE, Murdock PR, Randall AD, Rennie GI, Gloger IS (2000) Clo-


