Brief Communications

TASK-Like Conductances Are Present within Hippocampal CA1 Stratum Oriens Interneuron Subpopulations


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TASK-1 (KCNK3) and TASK-3 (KCNK9) are members of the two-pore domain potassium channel family and form either homomeric or heteromeric open-rectifier (leak) channels. Recent evidence suggests that these channels contribute to the resting potential and input resistance in several neuron types, including hippocampal CA1 pyramidal cells. However, the evidence for TWIK-related acid-sensitive potassium (TASK)-like conductances in inhibitory interneurons is less clear, and mRNA expression has suggested that TASK channels are expressed in only a subpopulation of interneurons. Here we use immunocytochemistry to demonstrate prominent TASK-3 protein expression in both parvalbumin-positive- and a subpopulation of glutamic acid decarboxylase (GAD)67-positive interneurons. In addition, a TASK-like current (modulated by both pH and bupivacaine) was detected in 30-50% of CA1 stratum oriens interneurons of various morphological classes. In most neurons, basic shifts in pH had a larger effect on the TASK-like current than acidic, suggesting that the current is mediated by TASK-1/TASK-3 heterodimers. These data suggest that TASK-like conductances are more prevalent in inhibitory interneurons than previously supposed.

Key words: potassium channels; interneurons; TASK; KCNK; hippocampus; inhibition

Introduction

Within the nervous system, local and global changes in pH occur under a variety of physiological and pathophysiological conditions, including synaptic transmission, high-frequency action potential firing, ischemia, and stroke (Kaila and Chester, 1998; Katsura and Siesjo, 1998). These alterations in pH influence the function of nervous tissue, leading to increases or decreases in neuronal excitability (Balestrino and Somjen, 1988). Although numerous channels are sensitive to changes in pH, Twik-related acid-sensitive potassium (TASK) channels are emerging as target pH-sensitive ion channels that may be involved in pH-dependent activity changes (Bayliss et al., 2003).

TASK channels are voltage independent, potassium selective channels belonging to the two-pore domain potassium channel family (Goldstein et al., 2001; Bayliss et al., 2003). TASK-1 (KCNK3) and TASK-3 (KCNK9) are the predominant TASK subunit types expressed in the brain (Talley et al., 2001). High levels of TASK-3 mRNA and lower levels of TASK-1 mRNA were observed in presumed stratum oriens (SO) inhibitory interneurons (Talley et al., 2001). However, in a separate study, single cell reverse transcription (RT)-PCR suggested that only ~15% of CA1 SO interneurons with horizontally oriented dendrites contain TASK mRNA (Taverna et al., 2005). Furthermore, SO interneurons displayed “leakage” currents with weak acid sensitivity, consistent with expression of a TASK-like current (Taverna et al., 2005).

Because CA1 SO interneurons are a heterogeneous group (Somogyi and Klausberger, 2005), the possibility exists that TASK channels are expressed in other subpopulations, accounting for the disparity between in situ hybridization and single cell RT-PCR data. It is well established that interneuron subpopulations often have distinct resting membrane potentials, input resistances, and membrane time constants, consistent with each having different complements of potassium leak channels.

Here we use immunohistochemistry to show that TASK-3 mRNA localized to the CA1 subfield (Talley et al., 2001). High levels of TASK-3 mRNA and lower levels of TASK-1 mRNA were observed in presumed stratum oriens (SO) inhibitory interneurons (Talley et al., 2001). However, in a separate study, single cell reverse transcription (RT)-PCR suggested that only ~15% of CA1 SO interneurons with horizontally oriented dendrites contain TASK mRNA (Taverna et al., 2005). Furthermore, SO interneurons displayed “leakage” currents with weak acid sensitivity, consistent with expression of a TASK-like current (Taverna et al., 2005).

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We use immunohistochemistry to show that TASK-3 is expressed in most parvalbumin-positive interneurons as well as a subpopulation of somatostatin-containing interneurons. Furthermore, we demonstrate a TASK-like (pH and bupivacaine sensitive) potassium current in a large number of horizontal CA1 SO interneurons. Based on the greater sensitivity of these neurons to alkalization than acidification, we suggest that channels are comprised of TASK-1/TASK-3 heterodimers.

Materials and Methods

All experiments were conducted in accordance with animal protocols approved by the National Institutes of Health.

Immunohistochemistry. Adult (35–60 d) or postnatal day 16 (P16)
mice C57BL/6 mice were deeply anesthetized and transcardially perfused with PBS followed by 4% paraformaldehyde in 0.1 M phosphate buffer (PFA). Brains were postfixed (2 h) in PFA and placed in 25% sucrose, 10% glycerol in 0.1 M phosphate buffer overnight and sectioned (50 μm) with a freezing microtome.

Sections were permeabilized and blocked with 0.5% Triton X-100, 1% bovine serum albumin, and 10% goat serum in PBS (in 1 h at 22°C). Primary antibody was added to carrier solution (with 0.1% Triton X-100, 1% bovine serum albumin, and 1% goat serum in PBS) with an additional 5% goat serum at the following concentrations: 1:1000 rabbit anti-TASK-3, and generated and characterized in house (Berg et al., 2004) or obtained from Santa Cruz Biotechnology (Santa Cruz, CA); 1:1000 mouse anti-parvalbumin (Sigma, St. Louis, MO); and 1:2000 mouse anti-GAD67 (Chemicon International, Temecula, CA). Sections were incubated for 48–96 h at 4°C, washed with carrier solution, and incubated in secondary antibody [1:500 AlexaFluor 488 or 633 (Invitrogen, Carlsbad, CA); or 1:100 HRP-linked (Jackson ImmunoResearch, West Grove, PA) for TASK-3] in carrier solution (1 h at 22°C). For adult TASK-3, a tyramide signal amplification system was used at 1:50 in ammonium pyrrolidinedithiocarbamate (APDC) and sodium borohydride (S3787, Sigma; 100 μM Dl-APV (Tocris Bioscience); 5 μM SR95531 (Tocris Bioscience); or 100 μM picrotoxin (Tocris Bioscience), and 250 nm CGP55845 (Tocris Bioscience) was included to block AMPA-, NMDA-, GABA<sub>A</sub>, and GABA<sub>B</sub>-receptor-mediated synaptic transmission, respectively.

**Electrophysiology.** Hippocampal slices from P14–P18 C57BL/6 mice were prepared as described previously (Pelkey et al., 2005). Slices were placed in a recording chamber and continuously perfused (2–4 ml/min) with artificial CSF containing the following (in mM): 130 NaCl, 3.5 KCl, 1.25 NaH<sub>2</sub>PO<sub>4</sub>, 25 NaHCO<sub>3</sub>, 2 CaCl<sub>2</sub>, 2 MgCl<sub>2</sub>, and 10 glucose saturated with 95% O<sub>2</sub>/5% CO<sub>2</sub>. pH 7.4. CA1 SO interneurons were visually identified using a 40× objective and infrared differential interference contrast video microscopy (Axioskop; Zeiss, Oberkochen, Germany). Whole-cell patch-clamp recordings were performed at room temperature (22°C) using a Multiclamp 700A amplifier (Molecular Devices, Union City, CA). Recording electrodes (2.5–4 MΩ) pulled from boro-silicate glass (World Precision Instruments, Sarasota, FL) were filled with a solution containing the following (in mM): 130 NaCl, 3.5 KCl, 1.25 NaH<sub>2</sub>PO<sub>4</sub>, 25 NaHCO<sub>3</sub>, 2 CaCl<sub>2</sub>, 2 MgCl<sub>2</sub>, and 10 glucose saturated with 95% O<sub>2</sub>/5% CO<sub>2</sub>. pH 7.4. CA1 SO interneurons were visually identified using a 40× objective and infrared differential interference contrast video microscopy (Axioskop; Zeiss, Oberkochen, Germany).

**Analysis of electrophysiological recordings.** Data were analyzed using IgorPro 5.0 (WaveMetrics, Lake Oswego, OR). In voltage clamp, cell input resistance was determined by delivering a 50 ms, −5 mV step and calculated from the reciprocal of the average current amplitude of the last 10 ms divided by the voltage step. I–V curves were fit with the following Goldman–Hodgkin–Katz (GHK) current equation (Hille, 2001):

\[
I = P \frac{E}{RT} [S]_o - [S] \exp(-zFE/RT)
\]

where \(P\) is the permeability, \([S]_o\) and \([S]\) are the intracellular and extra-cellular concentrations of potassium, \(z\) is the valence, \(F\) is Faraday’s constant, \(R\) is the gas constant, \(E\) is the voltage, and \(T\) is the absolute temperature. The pH-sensitive current was obtained by subtracting the control (pH 7.4) current from the current measured in either acidic (pH 6.4) or basic (pH 8.4) solution. Data were corrected for the measured junction potentials of −6.5 mV (potassium gluconate internal solution) or −5.5 mV (potassium methyl sulfate internal solution). Changing the pH did not alter the junction potential. Data are presented as mean ± SEM.

**Results**

**TASK-3 is primarily expressed in parvalbumin-positive cells**

Previous studies using *in situ* hybridization techniques suggested that TASK-3 mRNA is localized to interneurons in CA1 SO (Talley et al., 2001). Using immunohistochemistry, we labeled coronal sections of mouse hippocampus with a TASK-3 antibody. TASK-3 expression was observed throughout the adult hippocampus (Fig. 1A) in a pattern consistent with that shown by *in situ* hybridization (Talley et al., 2001). TASK-3-immunoreactive interneurons were scattered throughout CA1 SO, primarily in cells with large vertically oriented primary dendrites (Fig. 1B) but also in cells with horizontally oriented primary dendrites (Fig. 1C). A similar pattern of TASK-3 immunoreactivity was also observed in juvenile (P16) mice (Fig. 1C). Double labeling with GAD67, a marker for GABAergic interneurons, revealed that 83 ± 3% of TASK-3-immunoreactive CA1 SO neurons were also GAD67 positive (Fig. 1D, G), confirming that TASK-3-positive neurons in the stratum oriens are indeed GABAergic interneurons. However, only 49 ± 6% of GAD67-positive neurons contained TASK-3 (Fig. 1D, H), suggesting that not all GABAergic interneurons contain TASK-3.

GABAergic interneurons can be divided into several subpopulations based on the markers that they express. Double labeling with antibodies to parvalbumin and TASK-3 revealed that 95 ± 3% of TASK-3-positive cells were parvalbumin positive (Fig. 1E, G) and that 97 ± 4% of parvalbumin-immunoreactive cells were also TASK-3 immunoreactive (Fig. 1H). Similarly, in juvenile (P16) mice, 93 ± 1% of parvalbumin-immunoreactive cells also expressed TASK-3. In mice expressing EGFP under the GAD67 promoter (GAD67–EGFP) a subset of GAD67-positive interneurons express EGFP (Oliva et al., 2000). These interneurons are somatostatin positive and project their axons to lacunomolecular. In tissue obtained from these mice, 66 ± 5% of GAD67–EGFP-positive interneurons exhibited TASK-3 immunoreactivity (Fig. 1F, H) and represented 22 ± 2% of TASK-3-positive neurons (Fig. 1G). The combined total of TASK-3-positive cells expressing parvalbumin or EGFP is >100%, suggesting overlap between these two populations, consistent with previous reports (Somogyi and Klausberger, 2005). Attempts to determine the TASK-3 content of other neurochemically defined populations of interneurons was equivocal and suggests that TASK-3 is largely expressed in parvalbumin-containing, and a subpopulation of somatostatin-containing interneurons. Importantly, these data indicate that TASK-3 protein is expressed in a greater number of CA1 interneurons than suggested previously (Tavares et al. 2005).

**CA1 SO interneurons contain a TASK-like current**

Although no specific inhibitors of TASK channels exist, the conductance of recombinant TASK channels is decreased by extracellular acidification and increased by alkalization (Bayliss et al., 2003). To determine the prevalence of TASK-like conductances in CA1 SO inhibitory interneurons, we made whole-cell patch-clamp recordings from visually identified and anatomically confirmed interneurons. We tested 27 cells for sensitivity to both basic and acidic changes in pH, of which nine displayed changes in holding current, input resistance, and reversal potential that were consistent with a TASK-like current. In these nine cells, both alkalization caused a positive shift in the amount of current re-
input resistance of the cell (pH 7.4, 447 ± 69 MΩ; pH 6.4, 478 ± 68 MΩ; n = 9) (Fig. 2Ai, D). Notably, in one of those nine cells, bath acidification led to a large change (−90 pA) in the holding current as well as a 26% (pH 7.4, 78 MΩ; pH 6.4, 98 MΩ) increase in the input resistance (Fig. 2B).

Although little change in holding current was observed at holding potentials close to rest, a larger change was apparent at more negative potentials in these same nine cells. At a holding potential of −115 mV, a 24 ± 6 pA outward current was detected in response to 1 pH unit of acidification, whereas a −65 ± 7 pA inward current was detected in response to 1 pH unit of alkalization (current values are relative to current measured at pH 7.4) (Fig. 2E). Because bath alkalization resulted in a larger TASK-like current at resting potentials, subsequent analysis was performed only on the currents evoked by high pH.

To determine whether the current activated by alkalization was both time and voltage independent, hallmark properties of TASK-like conductances, we next measured the current–voltage (I–V) relationship of the pH-sensitive response in CA1 interneurons using either voltage steps or ramps delivered from −125 mV. To obtain the pH-sensitive current, the control (pH 7.4) current was subtracted from the current in basic (pH 8.4) solution (Fig. 2Aii, Bii, F). Voltage and time independence was tested by fitting the isolated pH-sensitive current to the modified GHK equation. Figure 2 shows that the I–V relationship of the base-sensitive current is well described by the GHK equation with a measured reversal potential close to the calculated EK (measured reversal, −86.5 ± 2.4 mV; n = 9; calculated EK, −95.5 mV). However, to verify that this isolated current was indeed potassium selective, in a separate series of experiments, the base-induced current was measured in an increased extracellular potassium concentration (8.5 mM). If the activated current is K selective, EK should shift by a calculated +22.5 mV. Indeed, in elevated potassium solution, 10 of 19 cells responded to alkalization with a current whose I–V relationship was again well fit by the GHK equation and reversed at −63.4 ± 0.7 mV (n = 10), representing a shift of 23.1 mV (Fig. 2F). These data indicate that the alkalization-activated current is a K-selective openly rectifying current.

To verify that these changes in holding current during voltage-clamp recordings manifested as changes in membrane potential, we next performed whole-cell current-clamp recordings in a separate set of experiments. Under control conditions, the average resting potential of CA1 SO interneurons was −54.2 ± 1.1 mV (n = 16). After bath alkalization, 10 of 16 neurons hyperpolarized, with an average change in resting potential of −12.8 ± 2.3 mV (p < 0.001) (Fig. 2G, H). The resting potential of the subpopulation of neurons that hyperpolarized in response to alkalization was −55.9 ± 1.4 mV (n = 10), which was significantly more negative than that of neurons that did not respond (−51.5 ± 0.9 mV; n = 6; p < 0.05). Furthermore, the average resting potential of nonresponders was unchanged after 5 min in alkaline solution (−50.3 ± 0.7 mV; p > 0.05).

Another hallmark property of TASK conductances is their block by local anesthetics such as bupivacaine (Bayliss et al., 2003). In another set of experiments, we first applied basic solution to increase the TASK-like current, and then applied the local anesthetic bupivacaine (20 μM; Sigma). In 9 of 22 CA1 SO inter-
neurons, alkalization resulted in a more positive holding current (88 ± 21 pA), decrease in input resistance (pH 7.4, 323 ± 42 MΩ; pH 8.4, 129 ± 36 MΩ), and a voltage-independent current that reversed near $E_K$ (−88 ± 1.1 mV), consistent with those cells containing a TASK-like current. In these nine interneurons, bupivacaine almost completely reversed the alkalization-activated current (mean decrease in holding current, −82 ± 22 pA), concomitant with an increase in the input resistance (pH 8.4, 129 ± 36 MΩ; pH 8.4 and bupivacaine, 256 ± 31 MΩ) (Fig. 3).

To determine whether this TASK-like current was found in morphologically distinct interneuron classes, cells were filled with biocytin during whole-cell recordings and anatomical reconstruction performed post hoc. We recovered the morphology of 26 cells that contained a TASK-like current. Of these 26 cells, 25 had horizontally oriented dendrites and cell bodies in CA1 SO but varied in their axonal morphologies. The large percentage of horizontal cells is likely the result of a selection bias while patching cells. In 16 cells, the axon crossed stratum (st.) pyramidale and projected to str. lacunosum molecular, where they branched extensively, consistent with them being oriens-lacunosum molecular (O-LM) cells (Fig. 2Aiii). Three cells had axons confined to str. pyramidale or proximal str. radiatum (basket/chandelier cells), and one was a back-projecting cell with axon ramiﬁying both in CA1 and CA3 str. radiatum (Fig. 3C). The remaining five cells had insuﬃcient axon for positive identiﬁcation. Finally, one cell had a cell body in CA1 SO but had vertically oriented dendrites, consistent with a basket cell morphology; however, the axon for this cell was not recovered (Fig. 2Biii). Of interest, this was the one cell with the largest response to acidification (Fig. 2B). Together these anatomical features are consistent with the immunohistochemical detection of TASK-3 in both somatostatin- and parvalbumin-containing interneurons.

Many CA1 SO interneurons do not exhibit a TASK-like current

Although we observed many cells with TASK-like currents, just over half of the interneurons tested under voltage-clamp conditions (39 of 67) did not display a TASK-like current, i.e., they did not respond to pH changes, did not display the expected change in input resistance, and the pH-sensitive current was not described by the GHK relationship. Eighteen neurons fit all three of these criteria. Figure 4A–C shows one example of a neuron that clearly did not contain TASK-like current. Both the holding current and input resistance were resistant to both alkalization and bupivacaine (Fig. 4A). Furthermore, the subtracted I–V showed no current activated in either condition (Fig. 4B). The morphologies of 10 interneurons that clearly did not display TASK-like current were recovered, and all of them had horizontal dendrites and cell bodies in CA1 SO. Five of these were identiﬁed as O-LM cells, and two were identiﬁed as basket/chandelier cells (Fig. 3C). The remaining neurons consisted of one bistratiﬁed cell with axon ramifying in SO and str. radiatum and two cells with insuﬃcient axon for positive identiﬁcation.

Twenty-one neurons partially ﬁtted the criteria for containing a TASK-like current, ﬁtting at least one of the criteria but not all
three. Figure 4D–F illustrates an example of a neuron in which the input resistance increased during acidification and decreased input resistance with alkalization, which were associated with changes in holding current (Fig. 4D). However, the isolated pH-sensitive I–V relationship was not adequately described by the GHK equation (Fig. 4E). All 13 recovered neurons had horizontally oriented dendrites. Of these, nine were identified as O-LM cells, and four had insufficient axon for additional characterization. Because of their ambiguity, these cells were omitted from the overall TASK-positive dataset.

For the 16 neurons tested in current clamp, six neurons did not hyperpolarize after addition of basic solution; rather, they had an average change in resting potential of 1.2 ± 0.7 mV, which was not significant. Of the three recovered neurons, two were O-LM neurons, and one was a horizontal neuron with insufficient axon for additional characterization.

Discussion

GABAergic interneurons vary widely in both their morphological and functional properties. The major source of functional heterogeneity arises from the distinct ion channel complements expressed in particular cell types (McBain and Fishah, 2001). In the present study, we observed TASK-3 immunoreactivity in ~50% of GAD67-positive interneurons. Similarly, TASK-like conductances were observed in just under half of all interneurons tested. TASK-3 protein was expressed primarily in parvalbumin-containing and a subset of GAD–EGFP (somatostatin)-containing SO neurons, which are mainly basket/chandelier and O-LM neurons, respectively. Consistent with this observation, morphologically identified CA1 SO neurons exhibiting TASK-like current were predominantly O-LM and basket/chandelier neurons.

TASK channel expression in SO interneurons has been previously investigated by Taverna et al. (2005), who concluded that TASK-like currents are not consistently detectable in horizontal CA1 SO interneurons, and only ~15% of CA1 SO interneurons express TASK-3 mRNA. However, in the present study, ~50% of morphologically identified horizontal interneurons displayed a TASK-like current. Furthermore, TASK-3 immunoreactivity was present in 66% of GAD67–GFP CA1 SO interneurons, of which the O-LM represents a subpopulation. This apparent discrepancy between our results and those of Taverna et al. (2005) likely results from the different methods used to detect the TASK-like conductance. First, Taverna et al. (2005), used single-cell PCR to determine TASK expression, which may have underestimated the number of cells expressing TASK mRNA. Consistent with this hypothesis, in the Taverna et al. (2005) study, electrophysiological data suggested that all pyramidal cells expressed TASK-like current, whereas mRNA was only detected in ~50% of pyramidal cells. Second, in our recordings the acid-sensitive current was small enough such that subsequent detection of the base-sensitive current, it is likely that we also would have identified these cells as pH insensitive. Because Taverna et al. (2005) relied on only acid sensitivity, the small (~2 mV) depolarization that they observed in acidic solutions did not conclusively identify them as pH sensitive.

TASK-like current in CA1 SO interneurons was only weakly sensitive to acid pH, consistent with a previous report that acidification results in only a small (~2 mV) depolarization of the resting potential in horizontal interneurons (Taverna et al., 2005). In contrast, the TASK-like current was much more sensitive to alkalization, which resulted in a more than twofold greater change in current than acidification (Fig. 2). This pH-sensitivity profile provides a tentative indication of the molecular identity of the underlying TASK channels. Although all combinations of TASK-1 and TASK-3 dimers have some sensitivity to acidification, TASK-1 homodimers and TASK-1/TASK-3 heterodimers have large increases in conductance during bath alkalization (Czirjak and Enyedi, 2002; Berg et al., Kang et al., 2004), whereas TASK-3 homodimers exhibit only a small increase in conductance during bath alkalization (Kim et al., 2000; Berg et al., 2004). In our recordings the current increase during alkalization is larger than the decrease during acidification, suggesting that TASK channels in SO interneurons may comprise either TASK-1/TASK-3 heterodimers or TASK-1 homodimers. Consistent with this hypothesis, low levels of TASK-1 mRNA have been observed in CA1 by both single-cell PCR (Taverna et al., 2005)
CA1 hippocampus, periods of high action-potential frequency transiently elevates extracellular pH (Kaila and Chesler, 1998). This transient pH elevation will increase the interneuron TASK current, decreasing excitability and potentially minimizing the role played by these interneurons during periods of excessive activity. The two subpopulations that we have identified as containing TASK, namely basket and O-LM cells, are involved in feedforward and feedback inhibition, respectively. Thus, TASK-dependent alterations in excitability may serve to regulate both feedforward and feedback inhibition.

References


