Inactivation properties of sodium channel Na\textsubscript{v}1.8 maintain action potential amplitude in small DRG neurons in the context of depolarization

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Abstract

**Background:** Small neurons of the dorsal root ganglion (DRG) express five of the nine known voltage-gated sodium channels. Each channel has unique biophysical characteristics which determine how it contributes to the generation of action potentials (AP). To better understand how AP amplitude is maintained in nociceptive DRG neurons and their centrally projecting axons, which are subjected to depolarization within the dorsal horn, we investigated the dependence of AP amplitude on membrane potential, and how that dependence is altered by the presence or absence of sodium channel Na\textsubscript{v}1.8.

**Results:** In small neurons cultured from wild type (WT) adult mouse DRG, AP amplitude decreases as the membrane potential is depolarized from -90 mV to -30 mV. The decrease in amplitude is best fit by two Boltzmann equations, having $V_{1/2}$ values of -73 and -37 mV. These values are similar to the $V_{1/2}$ values for steady-state fast inactivation of tetrodotoxin-sensitive (TTX-s) sodium channels, and the tetrodotoxin-resistant (TTX-r) Na\textsubscript{v}1.8 sodium channel, respectively. Addition of TTX eliminates the more hyperpolarized $V_{1/2}$ component and leads to increasing AP amplitude for holding potentials of -90 to -60 mV. This increase is substantially reduced by the addition of potassium channel blockers. In neurons from Na\textsubscript{v}1.8(-/-) mice, the voltage-dependent decrease in AP amplitude is characterized by a single Boltzmann equation with a $V_{1/2}$ value of -55 mV, suggesting a shift in the steady-state fast inactivation properties of TTX-s sodium channels. Transfection of Na\textsubscript{v}1.8(-/-) DRG neurons with DNA encoding Na\textsubscript{v}1.8 results in a membrane potential-dependent decrease in AP amplitude that recapitulates WT properties.

**Conclusion:** We conclude that the presence of Na\textsubscript{v}1.8 allows AP amplitude to be maintained in DRG neurons and their centrally projecting axons even when depolarized within the dorsal horn.

Background

Peripheral nociceptors have a highly specialized function: transducing noxious stimuli into neural activity and transmitting that activity to the central nervous system. Encoded nociceptive information is transmitted via small diameter axons that originate from a population of small cell bodies (20–30 μm) contained within dorsal root ganglia (DRG), which send their central projections to the

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Research

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**Conclusion:** We conclude that the presence of Na\textsubscript{v}1.8 allows AP amplitude to be maintained in DRG neurons and their centrally projecting axons even when depolarized within the dorsal horn.

Background

Peripheral nociceptors have a highly specialized function: transducing noxious stimuli into neural activity and transmitting that activity to the central nervous system. Encoded nociceptive information is transmitted via small diameter axons that originate from a population of small cell bodies (20–30 μm) contained within dorsal root ganglia (DRG), which send their central projections to the
spinal cord dorsal horn. It is well-established that extracellular potassium levels ([K⁺]o) within the dorsal horn can rise significantly as a result of neuronal activity induced by many stimuli including peripheral injury and noxious stimuli [1]: these changes in [K⁺]o can, in turn, lead to depolarization which can inactivate sodium channels, producing conduction block of neurons and neuronal processes such as axons and their terminals [2-4]. Yet some forms of peripherally induced pain have a persistent quality, suggesting that the nociceptive afferent barrage can be maintained even in the face of this depolarization. This raises the possibility that the membranes of nociceptive DRG neurons and their centrally projecting axons are constructed so as to permit the conduction of action potentials even when depolarized.

Small DRG neurons, which include nociceptors, express five of the nine functional voltage-gated sodium channels that have been sequenced thus far, including three that are selectively expressed within DRG neurons. Three tetrodotoxin (TTX)-sensitive (TTX-s) channels, Na\textsubscript{1.1}, 1.6 and 1.7 are expressed in adult small DRG neurons [5], as are the two TTX-resistant (TTX-r) channels, Na\textsubscript{1.8} and 1.9 [6-8]. These channels differ with regard to their biophysical properties. TTX-s channels in DRG neurons are characterized by relatively rapid activation and inactivation kinetics [9-13]. By comparison, the TTX-r channel Na\textsubscript{1.8} activates and inactivates more slowly [6,7]. Na\textsubscript{1.9} activates too slowly to contribute significantly to the upstroke of the action potential (AP), and its inactivation is so slow near activation threshold that the current is often referred to as persistent [14]. With regard to the voltage-dependence of channel opening and closing, TTX-r Na\textsubscript{1.8} channels in small DRG neurons have activation and steady-state fast inactivation functions that are depolarized compared to those of the TTX-s channels [6,7,9-13]. In DRG neurons, Na\textsubscript{1.7} channels have a relatively slow recovery from fast inactivation that presumably reduces its contribution to high frequency firing [15]. On the other hand, Na\textsubscript{1.7} channels have a relatively slow rate of inactivation from the closed state, which allows them to produce inward current in response to slow depolarizations.

Before the identification of the TTX-r channels Na\textsubscript{1.8} and Na\textsubscript{1.9}, it was known that DRG neurons were different from other neurons because they could generate sodium-dependent action potentials (AP) in the presence of TTX [16-20]. Investigation of AP electrogenesis in small DRG neurons from Na\textsubscript{1.8} (-/-) animals by Renganathan et al [20] demonstrated that cells with a resting membrane potential (RMP) close to -50 mV produced small, graded responses to depolarization, whereas cells with a more hyperpolarized RMP (-65 mV) produced nearly full-sized APs. These findings suggested that TTX-s sodium channels were capable of generating AP’s in small DRG neurons, but that most of these channels are inactivated at the typical RMP for these cells while, since the inactivation of Na\textsubscript{1.8} channels is more depolarized, more Na\textsubscript{1.8} channels are available at RMP to contribute to the AP. Similar conclusions based on voltage and current clamp recordings have been reached by other groups [21,22]. Na\textsubscript{1.8} is known to be expressed within the spinal cord dorsal horn, where it is present within the central terminals of primary sensory neurons [23,24]. In the present study we investigated the contribution of Na\textsubscript{1.8} to AP electrogenesis in DRG neurons from a different perspective, that of the reduction of AP amplitude in current clamp mode. We hypothesized that i) AP amplitude should decrease as a function of membrane potential, because membrane potential determines the degree of inactivation for the various sodium channel subtypes, and that ii) the presence of Na\textsubscript{1.8} should shift the voltage-dependence of AP amplitude in a depolarized direction, permitting relatively large APs to be generated even in depolarized neurons.

### Results

We first tested the dependence of AP amplitude on the membrane potential of small (< 25 um) DRG neurons from adult wild type mice. This was achieved in current clamp mode, by using injected current to hold the cell at a series of membrane potentials starting at -90 mV, with successive depolarizing steps of approximately 5 mV. At each holding potential (V\textsubscript{h}), the cell was stimulated with a positive-going square pulse of current (100 ms) starting just below threshold and proceeding in 10 pA intervals to a level 40–50 pA beyond threshold. A plot of AP amplitude (see Methods section for details on measurement procedure) and V\textsubscript{h} (Figure 1) revealed that AP amplitude decreased gradually as a function of cell membrane potential, but with a clear shoulder in the relationship. The voltage-dependent decrease is clearly visible in representative AP responses (waveform insets a-e in Figure 1). The relationship between AP amplitude and V\textsubscript{h} was best fit with 2 Boltzmann equations, one having a mean V\textsubscript{1/2} of -73 ± 2 mV and accounting for 27 ± 6% of the decrease in AP amplitude, the other having a mean V\textsubscript{1/2} of -37 ± 2 mV and accounting for 81 ± 6% of the decrease in AP amplitude. The more hyperpolarized V\textsubscript{1/2} is consistent with the voltage-dependence of steady-state fast inactivation that is characteristic of the TTX-s sodium channels present in small DRG neurons (V\textsubscript{1/2}s of -64 to -75 mV), while the more depolarized V\textsubscript{1/2} is consistent with a contribution from TTX-r channels, which have fast inactivation V\textsubscript{1/2}s of -30 to -45 mV [6,7,9-13].

To investigate the hypothesized contribution of TTX-s currents to AP amplitude, we performed the same analysis of AP amplitude and V\textsubscript{h} in DRG from WT mice with the inclusion of TTX (300 nM). As seen in Figure 2, for V\textsubscript{h} values from approximately -50 mV to -25 mV, there was a
gradual decrease in AP amplitude that was well fit with a single Boltzmann equation. The mean $V_{1/2}$ characterizing this voltage-dependence was $-36 \pm 2$ mV, which was similar to the depolarized component in absence of TTX and consistent with the elimination of TTX-s currents responsible for the decrease in AP amplitude at more hyperpolarized membrane potentials. An unexpected finding was that AP amplitude actually increased in amplitude as $V_h$ was increased from -90 to about -60 mV. The average AP amplitude at a $V_h$ of -90 mV was $46.2 \pm 4.7$ mV, representing a decrease of $45 \pm 4\%$ compared to the maximum AP amplitude. In the absence TTX-s sodium currents and any other influence on AP amplitude, it would be expected that AP amplitude would remain constant at these hyperpolarized potentials because TTX-r sodium channels would be in closed and activatable states.

We hypothesized that, in addition to voltage-gated sodium channels, a strong influence on AP amplitude would be from outward potassium currents. To test this hypothesis, we performed the same AP amplitude/$V_h$ analysis in the presence of TTX (300 nM), TEA (25 mM) and 4-AP (3 mM) to block a significant portion of outward potassium current. Just before switching to current clamp recording mode, the extent of potassium channel block was determined by holding cells at -100 mV and stepping for 100 ms to membrane voltages between -40 and +25. An example of currents produced by this protocol for two cells, one in the absence (A) and one in the presence (B) of TEA and 4-AP are shown in Figure 3. In the absence of TEA and 4-AP, maximum outward current often exceeded 20 nA, thus saturating the amplifier. To quantify the extent of block, maximum outward current was measured at steps to 0 mV. For cells in the absence of TEA and 4-AP, the maximum outward current at 0 mV was $13.8 \pm 1.9$ nA ($N = 8$), whereas for cells in the presence of TEA and 4-AP the maximum outward current was $1.4 \pm 0.3$ nA ($N = 6$), representing a reduction of 90%. For this group of cells tested in the presence of TEA, 4-AP as well as TTX, AP amplitude decreased gradually for $V_h$ values between -60 and -20 (Figure 4). This voltage dependence was fit with a single Boltzmann equation producing a $V_{1/2}$ of $-40 \pm 1.3$ mV, which was similar to the single component fit for WT neurons in the presence of TTX. In addition, the increase in AP amplitude observed in the presence of TTX alone for $V_h$ from -90 to -60 mV was significantly reduced. The average AP amplitude at $V_h$ -90 mV was $83.6 \pm 6.4$ mV, representing a decrease of $9 \pm 3\%$ of the maximum AP amplitude. Thus, blocking 90% of the outward potassium current led to a reduction in the decrease in AP amplitude at the most hyperpolarized membrane potential (-90 mV) from 45 to 9%. The remaining decrease may be attributable to the unblocked outward current or additional factors that can influence AP amplitude.

To further investigate the contribution of TTX-r currents to the decrease in AP amplitude associated with the more depolarized $V_{1/2}$ we applied the AP amplitude/$V_h$ analysis to small DRG neurons obtained from Na$_v$1.8(-/-) mice transfected with either GFP alone, or GFP plus a construct containing Na$_v$1.8. As shown in Figure 5, AP amplitude in cells transfected with GFP alone decreased gradually as $V_h$ increased from -90 to -30 mV, and this decrease was best fit with a single Boltzmann equation. The mean $V_{1/2}$ for 10 cells was $-55.0 \pm 1.5$ mV, a value that is depolarized by
about 20 mV compared to the TTX-s component in WT neurons. When Na\textsubscript{v1.8} was transfected back into small DRG neurons from Na\textsubscript{v1.8}(−/−) animals, analysis of the decrease in AP amplitude as a function of V\textsubscript{h} two populations revealed two populations: for one group, decrease in AP amplitude was best fit by a single Boltzmann equation (Figure 6A) having a mean V\textsubscript{1/2} of -52 ± 1.5 mV (N = 7). This value was not statistically different from the V\textsubscript{1/2} value for the cells from Na\textsubscript{v1.8}(−/−) animals transfected with GFP alone. Presumably, this group of cells did not express the Nav1.8 construct. In a second group of cells (N = 8), the decrease in AP amplitude was best fit by two Boltzmann equations with mean V\textsubscript{1/2}'s of -71 ± 2.1 mV and -35.6 ± 3.0 mV. These values are not significantly different from the corresponding values for the small DRG neurons from WT animals. In addition, the portion of the decrease in AP amplitude accounted for by each V\textsubscript{1/2} was similar to the WT data (24 ± 3% for the more hyperpolarized component, 77 ± 3% for the more depolarized component). These data indicate that the shift in V\textsubscript{1/2} observed in the Nav1.8(−/−) neurons, presumably dependent on the fast inactivation of TTX-s sodium currents, is due to the absence of Na\textsubscript{v1.8} channels.

Discussion

This work was motivated by the question of how AP conduction can be maintained in the central projections of nociceptive DRG neurons which course through the dorsal horn where they are subjected to depolarization [1,2] and by the observations that Na\textsubscript{v1.8} channels are present within the central terminals of primary sensory neurons within the dorsal horn [23,24] and that the amplitude of APs recorded from small DRG neurons is dependent on the holding potential of the cell, and that this voltage
dependence is best fit with two Boltzmann equations. Because sodium channels provide the primary source of current for the AP upstroke, the latter finding suggested a method for investigating the contribution of the various sodium channels expressed by small DRG neurons to AP generation. The $V_{1/2}$ values for the two Boltzmann fits (-73 mV and 37 mV) are similar to the $V_{1/2}$ values for fast steady-state inactivation associated with TTX-s channels (-65 to -75 mV) and $\text{Na}_v1.8$ TTX-r channels (-30 to -40 mV), respectively, that are expressed in small DRG neurons [6,7,9-13]. The substantial contribution of the component with the more depolarized $V_{1/2}$ to AP amplitude (81%), is consistent with previous estimates of the relative contribution of $\text{Na}_v1.8$ current to AP amplitude. Using the Goldman-Hodgkin-Katz equation to estimate sodium ion permeability during the AP in DRG from $\text{Na}_v1.8(+/-)$ and $\text{Na}_v1.8(-/-)$ animals, Renganathan concluded that $\text{Na}_v1.8$ channels contribute 80–90% of the current that flows at the peak of the AP [20]. Blair and Bean isolated TTX-s, TTX-r and high voltage-activated calcium currents in small DRG neurons stimulated with a simulated AP voltage protocol [21]. They estimated that TTX-r currents contributed 60% of the AP upstroke, as compared to 40% for the TTX-s component.
To confirm the contribution of TTX-s channels, we determined the voltage-dependence of AP amplitude in the presence of 300 nM TTX. For \( V_h \) values of approximately -60 to -20 mV, AP amplitude decreased gradually, and this voltage-dependence was well fit with a single Boltzmann equation having a \( V_{1/2} \) of -36 mV. The absence of the more hyperpolarized component demonstrates a contribution of TTX-s sodium currents. In place of a decrease in AP amplitude, we observed that AP amplitude increased for \( V_h \) values of -90 to -60 mV. It is unlikely that an increasing availability of TTX-r channels could account for increasing AP amplitude in this voltage range because it does not fall with the range for steady-state fast inactivation of these channels. Alternatively, we hypothesized that outward current generated by potassium channel opening might be responsible for the change in AP amplitude in the voltage range. To test this hypothesis, we recorded AP amplitude under conditions known to block the majority of voltage-gated potassium channels (TEA and 4-AP). Estimates of outward current amplitude obtained before switching to current clamp mode (see Methods) indicated that 80–90% of the outward current produced by depolarizing voltage steps was blocked. In current clamp mode, analysis of AP amplitude and \( V_h \) revealed that the 45% increase in AP amplitude from -90 to -60 mV in the presence of TTX alone decreased to only 9% in the presence of TEA and 4-AP, representing a reduction of 80%. Although we cannot state with certainty the identity of the potassium channels reducing AP amplitude from -90 to -60 mV, possible candidates are the inwardly rectifying current \( I_{IR} \) [25] and \( I_h \) [26], a slowly activated inward current, both of which are initiated by membrane hyperpolarization. It is interesting to consider the possibility that one of the roles of TTX-s sodium currents in small DRG neurons is to boost AP amplitude at these hyperpolarized membrane potentials.

In the absence of well-established blockers of the \( N_{a1.8} \) sodium current, we used DRG neurons from \( N_{a1.8}(-/-) \) animals to further investigate the identity of the currents contributing to the depolarized \( V_{1/2} \) characterizing AP amplitude reduction. Previous studies have demonstrated that small DRG neurons from these animals completely lack the slow TTX-r current produced by \( N_{a1.8} \) channels.

Figure 6
Transfection with \( N_{a1.8} \) produces two populations of cells with respect to voltage-dependence of AP amplitude. Data are from two small DRG neurons from \( N_{a1.8}(-/-) \) mice transfected with \( N_{a1.8} \). The cell in (A) is representative of a population of cells (N = 7) for which the decrease in AP amplitude was best fit with a single Boltzmann equation having a \( V_{1/2} \) of -53.9 mV (solid line). The cell in (B) is representative of a population of cells (N = 8) for which the decrease in AP amplitude was best fit by two Boltzmann equations, with \( V_{1/2} \) values of -72.6 mV (solid line) and -33.3 mV (dashed line).
[27]. AP amplitude in these cells decreased with a voltage dependence that was well fit by a single Boltzmann equation with a $V_{1/2}$ of $-55$ mV. This value represents a depolarizing shift of 20 mV compared to the $V_{1/2}$ for the TTX-s component in WT neurons. One possible explanation is that the inactivation properties of the remaining TTX-r sodium channels, Na$_v$1.9, are combining with those of TTX-s sodium channels to produce a single depolarized $V_{1/2}$. The midpoint of steady-state fast inactivation for Na$_v$1.9 in DRG cells ranges from -44 to -54 mV [14,28]. However, the slow onset of Na$_v$1.9 channel openings indicates that these channels make only a minor contribution to AP amplitude [22,28]. Akopian et al [27] observed an up-regulation of Na$_v$1.7 in Na$_v$1.8(-/-) DRG neurons; however, this would not account for the shift in $V_{1/2}$ because the $V_{1/2}$ for steady-state fast inactivation of Na$_v$1.7, -71 mV to -78 mV [15,29], is similar to the more hyperpolarized $V_{1/2}$ associated with the TTX-s component of AP decrease in WT neurons. An alternative explanation for the shift in the $V_{1/2}$ of the TTX-s component is that the inactivation properties of TTX-s sodium channels are modified in DRG neurons of Na$_v$1.8(-/-) animals. Such a modification has been observed in a previous study [20]. In a comparison of TTX-s currents in small DRG neurons from Na$_v$1.8(+/-) and Na$_v$1.8(-/-) mice, the authors observed a 20 mV depolarizing shift in the voltage dependence of fast inactivation. The mechanism for this shift in TTX-s voltage dependence of inactivation was not determined. Possible mechanisms include G-protein activation, which has been shown to depolarize inactivation $V_{1/2}$ of Na$_v$1.8 currents by 3–4 mV in DRG neurons [30], the presence of arachidonic acid, which hyperpolarizes the inactivation $V_{1/2}$ of both TTX-s and TTX-r currents in DRG neurons [31] and tyrosine kinase phosphorylation, which has been shown to both depolarize the $V_{1/2}$ of fast inactivation in cardiac Na$_v$1.5 channels in HEK293 cells, [32] and hyperpolarize the $V_{1/2}$ of fast inactivation in sodium currents of differentiated PC-12 cells [33]. Because expression of $\beta$ subunits can influence sodium channel inactivation properties [34], differential expression of $\beta$ subunits between Na$_v$1.8(+/-) and Na$_v$1.8(-/-) animals might also account for the observed shift. Irrespective of the modulatory process involved, our additional results with transfection of Na$_v$1.8 into cells from Na$_v$1.8(-/-) animals demonstrates that the process is not permanent and is dependent on the absence of Na$_v$1.8 channels. After transfections, AP amplitude dependence on $V_h$ was again fit best by two Boltzmann equations in half of the cells, and the $V_{1/2}$ values (-71 and -36 mV) were similar to values for the wild type DRG neurons. These results indicate that the reintroduced Na$_v$1.8 channels led to a shift in the inactivation voltage-dependence of the TTX-s component back to the WT value.

Conclusion

In summary, we have shown that the AP amplitude of small DRG neurons decreases as a function of membrane depolarization, and that this relationship is best fit with two Boltzmann components. The $V_{1/2}$ characterizing the more hyperpolarized component is similar to the $V_{1/2}$ associated with TTX-s sodium channel steady-state fast inactivation and is eliminated in the presence of TTX. The $V_{1/2}$ characterizing the more depolarized component is similar to the $V_{1/2}$ of steady-state fast inactivation for Na$_v$1.8. In small DRG neurons from Na$_v$1.8(-/-) animals, the voltage-dependence of the AP amplitude decrease is characterized by a single Boltzmann component with a $V_{1/2}$ of -55 mV, a value that is depolarized by about 20 mV compared to the TTX-s component in neurons from WT animals. We believe that this shift is due to a shift in the inactivation properties of TTX-s sodium channels in small DRG neurons from Na$_v$1.8(-/-) animals, but that this shift is not permanent. When these same cells are transfected with Na$_v$1.8, the voltage-dependence of AP amplitude recapitulates the voltage-dependence observed in neurons from WT animals. Taken together, these results suggest that the expression of Na$_v$1.8 within nociceptive DRG neurons permits the central projections of these cells to conduct APs even when depolarized within the dorsal horn.

Methods

Animal care

All animal care and surgical procedures followed a protocol approved by the Animal Care and Use Committee of Yale University. A colony of Na$_v$1.8(-/-) mice was raised from a breeding pair of Na$_v$1.8(+/-) mice [27] generously provided by John Wood (University College, London). Animals were housed in a pathogen-free area at the Veterans Affairs Medical Center in West Haven, CT and fed ad libitum.

Culturing and transfection of DRG neurons

DRGs were harvested from deeply anesthetized (ketamine/xylazine; 80/10 mg/kg; i.p.) adult (4–6 weeks) C57BL6 mice or adult Na$_v$1.8(-/-) mice (in C57/BL6 genetic background). Neurons were isolated as previously described [35]. Briefly, tissue was enzymatically digested in collagenase A (1 mg/ml, Roche) and sterile complete saline solution (CSS) for 20 min at 37°C, and then for 20 min in CSS solution with collagenase D (1 mg/ml, Roche). The pRK-Na$_v$1.8 plasmid used for transfections of Na$_v$1.8 into DRG from Na$_v$1.8(-/-) mice was a gift from John Wood (University College, London). The green fluorescent protein plasmid pEGFP was purchased from Clontech (Palo Alto, CA). Sodium channel and GFP constructs (channel:GFP ratio of 5:1) were electroporated into DRG neurons using Nucleofector Solution (Amaxa, Gaithersburg, MD) as previously described [36,37]. Transfected
DRG neurons were incubated at 37°C in Ca²⁺- and Mg²⁺-free culture medium (DMEM plus 10% fetal calf serum) for 5 min. to increase cell viability. The cell suspension was then diluted in culture medium, plated on 12 mm circular coverslips coated with laminin and poly-ornithine, and incubated at 37°C and 5% CO₂.

**Current and voltage clamp electrophysiology**

Small (20–25 μm) DRG neurons were used for current clamp recording after 24–40 hours in culture (with or without transfection). Electrodes had a resistance of 1–2 MΩ when filled with the pipette solution, which contained (in mM): 140 KCl, 0.5 EGTA, 5 HEPES and 3 Mg-ATP (pH 7.3 with KOH, adjusted to 320 mOsm with dextrose). The extracellular solution contained (in mM): 140 NaCl, 3 KCl, 2 MgCl₂, 2 CaCl₂, 10 HEPES (pH 7.3 with NaOH, adjusted to 320 mOsm with dextrose). The whole-cell recording configuration was obtained in voltage clamp mode with an Axopatch 200B amplifier (Molecular Devices, Sunnyvale, CA). Data was stored on a personal computer via a Digidata 1322a A/D converter (Molecular Devices) at an acquisition rate of 50 kHz with a lowpass Bessel filter setting of 5 kHz. In voltage clamp recording mode, voltage errors were minimized with 85% series resistance compensation, and linear leak currents and capacitance artifacts were subtracted out using the P/N method provided by Clampex (Molecular Devices) acquisition software. Clampfit (Molecular Devices) and Origin (Microcal Software, Northampton, MA) were used for data analysis. To obtain an estimate of inward and outward current amplitudes, two activation protocols were applied before proceeding to the current clamp recording mode. Both protocols involved steps to a range of membrane potentials from -45 to 40 mV, in 5 mV increments. For the first protocol, cells were held at -50 mV to eliminate the majority of TTX-s sodium channels. For the second protocol, cells were held at -100 mV to activate both TTX-s and TTX-r channels. A rough estimate of TTX-s current was obtained by subtraction of the currents obtained from the two protocols. After switching to current clamp recording mode, passive and active cell properties were determined (RMP, input resistance, spike amplitude and threshold at RMP). Cells with RMPs more negative than -50 mV were used for the analysis. AP amplitude was defined as the difference between the peak and the onset of the AP. Onset of the AP was further defined as the point at which dV/dt for the AP was greater than 5 mV/ms (this value was chosen to be approximately 2 times the maximum baseline noise of the dV/dt trace). AP amplitude was plotted as a function of V_h and fit with either 1 or 2 Boltzmann equations of the form V_{AP}/V_{APmax} = 1/(1 + exp((V_h - V_{1/2}))/k), for which V_{AP} is the AP amplitude, V_{APmax} is the maximum AP amplitude, V_h is the holding potential, V_{1/2} is the V_h at which the AP amplitude is 50% of maximum, and k is the slope of the fit. Goodness of fit for 1 and 2 Boltzmann equations was determined by eye. Data is expressed as mean ± sem, and for all data comparisons, Student's t-test was used to assess statistical differences between means from different groups.

**Competing interests**

The author(s) declare that they have no competing interests.

**Authors' contributions**

SGW and TPH planned the experiments. TPH conducted all the experiments. Both authors contributed to the writing of the paper.

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