Aquaporin-1 Tunes Pain Perception by Interaction with Na$_v$1.8 Na$^+$ Channels in Dorsal Root Ganglion Neurons*

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Aquaporin-1 (AQP1) water channels are expressed in the plasma membrane of dorsal root ganglion (DRG) neurons. We found reduced osmotic water permeability in freshly isolated DRG neurons from AQP1$^{-/-}$ versus AQP1$^{+/+}$ mice. Behavioral studies showed greatly reduced thermal inflammatory pain perception in AQP1$^{-/-}$ mice evoked by bradykinin, prostaglandin E$_2$, and capsaicin as well as reduced cold pain perception. Patch clamp of freshly isolated DRG neurons showed reduced action potential firing in response to current injections. Single action potentials after pulse current injections showed reduced maximum inward current, suggesting impaired Na$_v$1.8 Na$^+$ potentials after pulse current injections showed reduced maximum inward current, suggesting impaired Na$_v$1.8 Na$^+$ function. Whole-cell Na$_v$1.8 Na$^+$ currents in Na$_v$1.8-expressing ND7-23 cells showed slowed frequency-dependent inactivation after AQP1 transfection. Immunoprecipitation studies showed AQP1-Na$_v$1.8 Na$^+$ interaction, which was verified in live cells by single-particle tracking of quantum dot-labeled AQP1. Our results implicate the involvement of AQP1 in DRG neurons for the perception of inflammatory thermal pain and cold pain, whose molecular basis is accounted for, in part, by reduced Na$_v$1.8-dependent membrane Na$^+$ current. AQP1 is, thus, a novel target for pain management.

Aquaporins (AQPs) are water-transporting proteins expressed in epithelial, endothelial, and other cell types. In the central nervous system AQP4 is expressed in glial cells, where it plays a role in cerebral edema (1, 2), glial cell migration (3, 4), and neuroexcitation (5, 6). The mechanisms of AQP4 modulation of seizure dynamics (5), cortical spreading depression (6), vision (7), hearing (8), and olfaction (9) remain unclear. AQP4-dependent Kir4.1 K$^+$ channel function has been suggested from delayed K$^+$ reuptake from brain extracellular space after neuroexcitation (6, 10); however, patch clamp analysis showed AQP4-independent Kir4.1 K$^+$ channel function (11). Extracellular space expansion, which has been found AQP4-deficient brain (12, 13), may contribute to the altered neuroexcitation phenotype. Neurons in the central nervous system do not express AQPs.

Water channel AQP1 is abundant in hematopoietic cells and kidney. In normal brain AQP1 expression is restricted to the choroid plexus, where it facilitates the secretion of cerebrospinal fluid (14). In the spinal cord and peripheral nervous system, AQP1 is expressed in sensory neurons in dorsal root ganglia (DRG) that are associated with pain nociception (15–17). These neurons, whose cell bodies reside in the DRG, carry sensory signals from the periphery through small diameter, non-myelinated fibers that synapse in the superficial lamina of the dorsal horn in the spinal cord (18). Two prior studies on pain phenotype in AQP1$^{-/-}$ mice have reported conflicting behavioral findings. Oshio et al. (16) reported mild impairment in pain nociception in AQP1$^{-/-}$ mice after thermal (tail flick) and chemical (capsaicin) stimuli, with no differences in response to mechanical stimuli. Shields et al. (15) confirmed AQP1 expression in DRG neurons and partially colocalization with TRPV1 and substance P; however, they reported no significant differences in behavioral pain tests. The role of AQP1 in neuronal function in the DRG has, thus, remained unclear, as does its role in neurons in trigeminal and nodose ganglia, where it is also expressed.

To clarify the role of AQP1 in pain physiology, we did more extensive behavioral testing as well as immunolocalization, water permeability, and patch clamp studies on freshly isolated dissociated DRG neurons from wild type (AQP1$^{+/+}$) and AQP1 null (AQP1$^{-/-}$) mice. We found greatly reduced behavioral response to inflammatory thermal and cold pain in litter-matched AQP1$^{-/-}$ mice and distinct electrophysiological defects related to impaired Na$_v$1.8 Na$^+$ channel functioning in AQP1-deficient DRG neurons. Patch clamp, immunoprecipitation, and single particle tracking studies in transfected cell models suggested a novel AQP1-Na$_v$1.8 interaction as responsible, in part, for the impairment in pain-sensing in AQP1 deficiency.

**EXPERIMENTAL PROCEDURES**

Mice—AQP1$^{-/-}$ mice in a CD1 (out-bred) strain were generated by targeted gene disruption as described (19). Mice in a C57/b16 (inbred) genetic background were generated by >10 back-crosses. 8-to-10-week-old, age-matched mice were used. Investigators were blinded to genotype information in all experiments. Protocols were approved by the University of California San Francisco Committee on Animal Research.

Isolation of DRG Neurons—Adult mice were decapitated after anesthesia. L4–6 DRGs were removed and treated with collagenase (type I A, 1.5 mg/ml, Sigma) and trypsin (1 mg/ml, Sigma) in Dulbecco’s modified Eagle’s medium at 37 °C for 30 min. After incubation, DRGs were washed five times and gently
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Aquaporin-1 (AQP1) is a water permeable channel critically involved in pain perception. The role of AQP1 in the peripheral nervous system is still an open question. In this study, we employed AQP1-overexpressing mouse neuroblastoma cells (ND7-23) and show that AQP1-overexpression reduces the Na+ current and cell excitability. This is associated with changes in AQP1 distribution and cell morphology. Immunohistochemistry and immunocytochemistry confirmed that AQP1 was localized to DRG neurons and was upregulated in DRG neurons following inflammation. Thus, AQP1 may be a potential therapeutic target for treating pain.

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AQP1 (AQP1.T120.mycΔ27 or AQP1.T120.mycΔ41). Two days after transfection, cell extracts were prepared using radioimmuno precipitation assay buffer lysis buffer containing protease inhibitor mixture (Pierce) according to the manufacturer’s instructions. Protein concentration was determined using the BCA protein reagent kit (Bio-Rad). The cell lysate was diluted to ~1 mg/ml of total protein with lysis buffer and mixed with rabbit anti-AQP1 polyclonal antibody (1 g/100 g of total protein, Santa Cruz), rabbit anti-Na⁺/1.8 polyclonal antibody (1 g/100 g of total protein, Millipore), or c-Myc monoclonal antibody (1 g/100 g of total protein, Covance, Emeryville, CA) for 1 h at 4 °C. Equal amounts of samples were mixed with either rabbit IgG or mouse IgG or vehicle as negative controls. Protein A/G beads were then added, and solutions were rocked for 1 h at 4 °C. The protein A/G beads were collected and washed three times with lysis buffer. Proteins were eluted by boiling in elution buffer (Pierce). Samples were resolved by SDS-PAGE and analyzed by immunoblot using mouse anti-AQP1 monoclonal antibody (Santa Cruz, 1:400), rabbit anti-Na⁺-1.8 antibody (Millipore), and rabbit anti-c-Myc antibody (Santa Cruz, 1:100). Clean-Blot IP detection reagent (Pierce) was used as the secondary antibody, and blots were detected using the SuperSignal (Pierce) detection system.

Behavioral Testing—All behavioral testing was done with genotype information blinded. Age-matched male and female mice were used. Several pain behavior studies were done using established procedures as follows. (a) For hotplate testing, mice were placed on a hotplate (model #35100, Ugo Basile, Comerio, Italy), and a ramp stimulus (45–52 °C, 2 °C/min) was applied. The temperature threshold was recorded at which paw-licking or jumping was observed. In some experiments the temperature was fixed at 50 °C, and the latency was determined for paw-licking or jumping (cut-off time 1 min). (b) For capsaicin testing, mice were kept in an empty cage. Total licking time over 5 min was recorded after intraplantar injection of capsaicin (10 µl, 1 or 3 µg). (c) Prostaglandin E₂-induced paw hypersensitivity was tested by measuring hotplate latency (50 °C) at specified times after intraplantar injection of PGE₂ (10 µl, 300 ng). (d) For bradykinin-induced acute pain testing, total licking time over 10 min was recorded after intraplantar injection of bradykinin (10 µl, 300 ng). In separate experiments hotplate latency (50 °C) was measured at 10 min after bradykinin injection. Paw edema was measured as paw thickness at 10 min. (e) For formalin testing, total licking time in 5-min intervals over 40 min was measured after intraplantar injection of 10 µl of 5% formalin in saline. (f) For cold pain testing, a cold-plate (model #35100, Ugo Basile) maintained at 4 °C was used to assess nocuous cold sensitivity of the plantar surface of the hind paws (27). Mice were placed on the cold plate for 5 min, and behavior was videoed and scored in a blinded manner. Severity of cold pain was scored in 5-s intervals using the following scale: 0; standing still; 1, walking; 2, each paw lifting occurrence; 3, each jumping occurrence.

Osmotic Water Permeability—Cell plasma membrane water permeability was measured by a calcein-quenching method (28). DRG cells were isolated after enzymatic digestion as described above, immobilized on a poly-L-lysine-coated cover glass, and loaded with calcein by 30-min incubation with 5 µM calcein-AM (Invitrogen). The cover glass was mounted in a custom perfusion chamber having a solution exchange time of <200 ms at 50 µl/min perfusion rate. The time course of cytoplasmic calcein fluorescence was measured in response to cell swelling produced by a 2-fold dilution of the extracellular bathing solution with water. Single cell calcein fluorescence was measured continuously using a Nikon inverted epifluorescence microscope equipped with 100× oil objective, halogen light source, calcein filter set (480-nm excitation, 490-nm dichroic mirror, 535-nm emission filter), photomultiplier detector, and 14-bit analog-to-digital converter.

Quantitative Real-time Reverse Transcription-PCR—L4–6 DRGs were collected after euthanasia, total RNA was isolated by a PureLink™ Micro-to-midi kit (Invitrogen), and cDNA was reverse-transcribed from mRNA with the Super-Script First Strand Synthesis System for reverse transcription-PCR (Invitrogen). Fluorescence-based quantitative real-time reverse transcription-PCR was carried out using the LightCycler 480 and with LightCycler FastStart DNA MasterPLUS SYBR Green I kit (Roche Diagnostics). Primers were as follows: 5′-TGTAT-GCCTCTGTCTGACACC-3′ (sense) and 5′-CAGGTCCAGA-CGAGGATG-3′ (antisense) for β-actin, 5′-CTCTCCTAGTG-GAATTC-3′ (sense) and 5′-ACGATACCCATGCG-3′ (antisense) for AQP1, 5′-ACGACATAGAATGGAC-AATCACC-3′ (antisense) for Na⁺,1.8, 5′-TGAGGCAACACTTCCACAAATG-3′ (sense) and 5′-AGCCAGAAACAGGTAATAGATG-3′ (antisense) for Na⁺,1.9, 5′-TCC-ATTATCATAATCCAGCCTCAC-3′ (sense) and 5′-GATCGGTCCGTTCTCTTTCG-3′ (antisense) for Na⁺,1.7, 5′-GTGCATCTCTTGTAAGGCCTTAG-3′ (sense) and 5′-ATCTCATAAGCGTGAATCTTCGAA-3′ (antisense) normalized to β1. Data were analyzed by LightCycler software 4.0 (Roche Diagnostics) and reported as calibrated ratios normalized to β-actin. Data were averaged from three mice of each phenotype.

Single Particle Tracking—Single particle tracking measurements were done on confluent cells grown on 18-mm cover glasses that were transfected with plasmid encoding AQP1.T120.myc at 18–24 h before measurements. Na⁺,1.8 or control pcDNA3 vector was co-transfected with AQP1 at a 5:1 ratio. In some experiments Na⁺,1.8 or control pcDNA3 vector was first co-transfected with plasmid encoding green fluorescent protein (10:1) for 36 h and then transfected with the AQP1.T120.myc, allowing measurements to be made selectively on green fluorescent protein-positive cells. Cells were labeled with c-Myc antibody (Covance) and followed by goat F(ab’2) anti-mouse Qdot 655 (Invitrogen) as described previously (29). Single particle tracking was done on a TE2000S inverted microscope (Nikon, Melville, NY) equipped with a 100× total internal reflection fluorescence oil immersion objective and EM-CCD (Hamamatsu, Bridgewater, NJ) with resolution 80 nm/pixel using continuous 11-ms acquisitions for 6 s (91 Hz). Single particle tracking image sequences were analyzed, and trajectories were constructed using IDL software (Research Systems, Boulder, CO) as described (29). Blinking of individual Qdots was analyzed by LightCycler 4.0 software (Roche Diagnostics) and reported as calibrated ratios normalized to β-actin. Data were averaged from three mice of each phenotype.
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Aquaporin-1 and IB4, a surface binding lectin specific for non-peptidergic neurons (21), 54 ± 4% of IB4-positive neurons were AQPI-positive. By double immunostaining with AQPI and CGRP, a neuropeptide specific for peptidergic small DRG neurons (18), 71 ± 1% of CGRP-positive neurons were AQPI-positive. The majority of small-diameter DRG neurons, thus, express AQPI.

To test whether AQPI in DRG neurons is functional as a plasma membrane water channel, water permeability was determined in freshly isolated DRG neurons from AQPI+/+ and AQPI−/− mice. Plasma membrane osmotic water permeability was measured by a calcein fluorescence quenching method (28), which quantifies the kinetics of cell volume change in response to osmotic gradients. Two types of responses were seen in small DRG neurons from AQPI+/+ mice, including neurons showing rapid changes in calcein fluorescence that were slowed by the AQPI inhibitor HgCl2 (Fig. 1C, left). Responses from many AQPI+/+ DRG neurons are summarized in Fig. 1C (right). All DRG neurons from AQPI−/− mice showed slow responses that were insensitive to HgCl2. These measurements indicate that AQPI is functional in DRG neurons.

Behavioral Analysis Shows Impaired Nocuous Thermal Pain Nociception in AQPI−/− Mice—The functional expression of AQPI in small diameter DRG neurons suggests its involvement in pain nociception. Initial studies of hotplate threshold (temperature ramp), hotplate latency (constant 50 °C temperature), paw licking time after high dose intraplantar injection of capsaicin (3 μg/paw), and paw licking time after intraplantar injection of formalin (5% in saline, 10 μl/paw), as done in the two prior studies (15, 16), showed no significant differences between AQPI+/+ and AQPI−/− mice (Fig. 2A).

Behavioral studies involving acute inflammatory pain were done in an attempt to discover differences in pain nociception in AQPI−/− mice that were not observed using the protocols in Fig. 2A. Bradykinin responses were tested as a measure of acute inflammatory pain (31). Intraplantar bradykinin injection produced an immediate paw licking response, which was greatly reduced in AQPI−/− mice (Fig. 2B, left). Also, after bradykinin injection, the paw withdrawal latency on a 50 °C hotplate was significantly reduced in AQPI+/+ mice but not changed significantly in AQPI−/− mice (Fig. 2B, middle). As a control, we found that paw edema, which is mediated after bradykinin injection by mast cell B1 receptors (31), was comparable in AQPI+/+ and AQPI−/− mice (Fig. 2B, right). Studies were also done using a different inflammatory mediator, PGE2, which is

RESULTS

AQPI Expression and Water Permeability in DRG Neurons—By immunofluorescence, AQPI immunoreactivity was seen in DRG neurons, mainly in the plasma membranes of small, ~25-μm diameter neurons (Fig. 1A, left). Immunoblot analysis showed non-glycosylated AQPI as a sharp band at 28 kDa and glycosylated AQPI as a diffuse band migrating at 36–42 kDa (Fig. 1A, right). Specific immunoreactivity was absent in DRGs from AQPI−/− mice. Double immunostaining of DRG sections with AQPI and peripherin, a small-diameter nociceptor marker (30), showed that 92 ± 2% of AQPI-positive neurons were peripherin-positive, indicating that nearly all AQPI-expressing cells are nociceptors (Fig. 1B). There are two subpopulations of nociceptive neurons, “peptidergic” and “non-peptidergic” types (18). By double immunostaining with AQPI and IB4, a surface binding lectin specific for non-peptidergic neurons (21), 54 ± 4% of IB4-positive neurons were AQPI-positive. By double immunostaining with AQPI and CGRP, a neuropeptide specific for peptidergic small DRG neurons (18), 71 ± 1% of CGRP-positive neurons were AQPI-positive. The majority of small-diameter DRG neurons, thus, express AQPI.

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involved in the development of acute inflammatory pain (32). Although intraplantar injection of PGE₂ did not itself produce a paw licking response, it produced a transient thermal hypersensitivity in AQP1⁺/⁻ mice which was absent in AQP1⁻/⁻ mice (Fig. 2C).

After discovery of these significant phenotype differences in bradykinin and PGE₂ testing, we re-examined capsaicin responses but using a lower dose (1 μg/paw) of capsaicin, which produces less nerve damage but a similar licking response (33, 34); 3 μg of capsaicin has been reported to destroy C fibers (35). Intraplantar injection of 1 μg of capsaicin produced a greatly reduced licking time in AQP1⁻/⁻ mice in both out-bred (CD1) and inbred (C57/bl6) mouse strains (Fig. 2D).

Altered Electrophysiological Responses of AQP1-deficient DRG Neurons—To investigate possible mechanisms to account for the greatly reduced pain nociception in AQP1⁻/⁻ mice, whole-cell patch clamp was done to compare electrophysiological responses from isolated, small DRG neurons (diameter <25 μm). Of 45 and 50 DRG neurons studied from AQP1⁺/+ and AQP1⁻/⁻ mice, respectively, there were no differences in cell size, resting membrane potential, or input resistance (Table 1). Because of the distinct electrophysiological properties of IB₄-positive and IB₄-negative small DRG neurons (21, 36), in subsequent patch clamp studies we labeled live cells with IB₄-Alexa 594 and only studied IB₄-negative small DRG neurons.

A first set of studies was done under current clamp conditions with 1-s current injections with a linear ramp current from 0–1 nA (Fig. 3A). The ramp current injection elicited a series of spikes over the 1-s injection with progressively reduced amplitude. Spikes with peak voltage >0 mV were defined as APs (23). The threshold was defined as the potential when the curve slope abruptly increased, as determined by derivative analysis, and the amplitude was defined as the potential change from the threshold to the peak. Of 26 small DRG neurons studied from AQP1⁺/+ mice, 22 responded, with the number of APs in response to the current ramp ranging from 6 to 28, with an average 16 ± 1 (S.E.). Of 28 small DRG neurons studied from AQP1⁻/⁻ mice, 24 responded, with a significantly reduced average number of APs of 11 ± 1 (S.E.). Fig. 3B summarizes the analyses done on data as in Fig. 3A. The threshold, initial latency (to the first AP peak), and the gap time (between the first two APs) showed no differences. The amplitude of the first AP showed no difference, but with continue firing the amplitudes of the fourth and fifth APs of AQP1⁻/⁻ neurons were significantly decreased. The total firing duration was reduced in AQP1⁻/⁻ DRG neurons. Counting of APs with an amplitude greater than 50% that of the first AP indicated remarkably

### TABLE 1
Electrophysiological properties of AQP1⁺/+ and AQP1⁻/⁻ small DRG neurons

| Membrane potential | Input resistance | Cell capacitance | Injection current threshold | AP 50% width | AHP |
|-------------------|------------------|-----------------|----------------------------|--------------|-----|
|                   | mV               | Gigaohms | Picofarads | nA        | ms  | mV  |
| AQP1⁺/+           | −62.7 ± 1.0      | (n = 45)  | 1.3 ± 0.1  | (n = 45)  | 18.7 ± 0.7     | (n = 45) | 0.6 ± 0.1        | (n = 26)  | 3.4 ± 0.4      | (n = 26)  | 12.3 ± 1.2    | (n = 26)  |
| AQP1⁻/⁻           | −62.9 ± 1.4      | (n = 50)  | 1.0 ± 0.1  | (n = 50)  | 17.6 ± 0.8     | (n = 50) | 0.6 ± 0.1        | (n = 28)  | 3.3 ± 0.4      | (n = 28)  | 11.3 ± 1.0    | (n = 28)  |
fewer APs and decreased the firing duration of AQP1<sup>−/−</sup> DRG neurons.

A second set of studies was done with a greater duration (10 s) of current injection using 100 pA constant current. Of 26 small DRG neurons studied from AQP1<sup>+/+</sup> mice, 22 responded with more than 1 AP, with 21 of 28 responders from AQP1<sup>−/−</sup> mice. Most DRG neurons from AQP1<sup>+/+</sup> mice fired throughout the depolarization, with minimal decay in the amplitude and frequency of APs; a substantially greater decay in APs was seen in DRG neurons from AQP1<sup>−/−</sup> mice (Fig. 3C). Analysis of all responding small DRG neurons showed significantly fewer APs in DRG neurons from AQP1<sup>−/−</sup> mice (47 ± 7 versus 28 ± 5 in 10 s) (Fig. 3D). Although the number of APs was comparable over the first second in the AQP1<sup>−/−</sup> neurons, the number of APs was much lower over the last second. Also, AP amplitudes in AQP1<sup>−/−</sup> DRG neurons decayed faster, taking 2.9 ± 0.8 s to decrease by 50% compared with 5.7 ± 0.5 s in AQP1<sup>+/+</sup> DRG neurons. At 1 s, the amplitude of APs in AQP1<sup>−/−</sup> DRG neurons was significantly decreased. The reduced AP firing in AQP1<sup>−/−</sup> DRGs, seen in two different current injection protocols, may account for the impaired pain nociception in AQP1<sup>−/−</sup> mice.

**Evidence for Nav1.8 as Responsible for the AQP1-sensitive Na<sup>+</sup> Current**—To identify the membrane channels responsible for the different responses in DRG neurons from AQP1<sup>+/+</sup> and AQP1<sup>−/−</sup> mice, we measured single APs after a short (1-ms) step-depolarizing current, leaving most of each AP free of the effect of injected current (22). Nociceptive DRG neurons generally show a long duration AP with a shoulder on the falling phase of the AP (21), as seen in the representative single APs in Fig. 4A. Comparing single APs from 26 and 28 small DRG neurons from AQP1<sup>+/+</sup> and AQP1<sup>−/−</sup> mice, respectively, no significant differences were found in AP threshold, AP amplitude, width at 50% amplitude, or maximum after-hyperpolarization (AHP) amplitude (Fig. 4B and Table 1). However, analysis of the time derivative of the voltage response (dV/dt) showed reduced
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**FIGURE 4. Impaired inward Na\(^+\) current in AQP1\(^{-/-}\) DRG neurons.** A, top, examples are shown of single APs from small DRG neurons elicited by injection of a 1-ms depolarizing current. Threshold, amplitude, 50% width, resting potential, and after-hyperpolarization (AHP) indices are indicated. Bottom, shown is ion current computed from the voltage waveform as \(-C dV/dt\), where C is the cell capacitance. B, analysis of data is as in A (S.E., 26 AQP1\(^{-/-}\) neurons and 28 AQP1\(^{+/+}\) neurons). Differences are not significant by t test, except as indicated by the asterisks (p < 0.05). C, left, examples are shown of whole-cell TTX-R Na\(^+\) currents. Right, shown is the current-voltage relationship of TTX-R Na\(^+\) currents (normalized by cell capacitance) (S.E., 22 AQP1\(^{-/-}\) neurons and 17 AQP1\(^{+/+}\) neurons, paired t test; *, p < 0.05). D, left, shown is a Na\(_{1.8}\) immunoblot of DRG in AQP1\(^{+/+}\) and AQP1\(^{-/-}\) mice (CD1 and C57/bl6 genetic backgrounds), with \(\beta\)-actin immunoblot for the same samples. Right, shown is relative mRNA expression of AQP1, Na\(_{1.8}\), Na\(_{1.7}\), Na\(_{1.9}\), and sodium channel \(\beta_1\) subunit quantified by real-time PCR (S.E., 3 mice per genotype). *, p < 0.05.

maximum \(dV/dt\) in AQP1\(^{-/-}\) neurons, giving a significant reduced maximum ionic current (\(I_{\text{ionic}} = -C dV/dt\)) in the inward direction. The maximum inward current represents a Na\(^+\) current with little contribution from other ion channels.

Both TTX-S and TTX-R Na\(^+\) channels are expressed on small DRG neurons. Several TTX-S Na\(^+\) channels are expressed on all DRG neurons, including Na\(_{1.1}\), Na\(_{1.2}\), and Na\(_{1.7}\) (37). TTX-R Na\(^+\) channels Na\(_{1.8}\) and Na\(_{1.9}\) are expressed mainly in small DRG neurons (38). In small DRG neurons >80% of the peak inward current during an AP upstroke is carried by Na\(_{1.8}\) (39, 40), whereas the TTX-S Na\(^+\) current contributes mainly to the AP initial threshold. We found that 300 nM TTX did not significantly change the maximal inward currents in AQP1\(^{+/+}\) or AQP1\(^{-/-}\) small DRG neurons (data not shown). To isolate Na\(_{1.8}\) Na\(^+\) currents, whole-cell currents were measured in the presence of TTX and reduced extracellular Na\(^+\) concentration of 35 mM. We studied 22 and 17 small DRG neurons from AQP1\(^{+/+}\) and AQP1\(^{-/-}\) mice, respectively. Although the voltage dependence of Na\(_{1.8}\) Na\(^+\) activation was similar in DRG neurons from AQP1\(^{+/+}\) and AQP1\(^{-/-}\) mice (Fig. 4C), the maximum current amplitude was significantly reduced in AQP1\(^{-/-}\) DRG neurons.

The reduced Na\(_{1.8}\) Na\(^+\) current in AQP1\(^{-/-}\) could be a consequence of reduced Na\(_{1.8}\) Na\(^+\) channel expression and/or a functional effect of AQP1 expression on Na\(_{1.8}\) Na\(^+\) current. Both possibilities were investigated. By immunoblot analysis, the expression of Na\(_{1.8}\) protein in DRGs from AQP1\(^{-/-}\) mice was reduced compared with that from AQP1\(^{+/+}\) mice, as seen in both out-bred (CD1) and inbred (C57/bl6) mouse strains (Fig. 4D). By quantitative reverse transcription-PCR, Na\(_{1.8}\) transcript expression was also significantly lower in AQP1\(^{-/-}\) mice, whereas that of Na\(_{1.7}\) and the sodium channel \(\beta_1\) subunit, which is widely expressed in all DRG neurons, were not changed (Fig. 4D).

**AQPI Modulates the Kinetic Properties of Na\(_{1.8}\) Na\(^+\) Currents in ND7-23 Cells**—The data above suggest impaired Na\(_{1.8}\) current in AQP1\(^{-/-}\) DRG neurons. To further study the involvement of AQP1 in Na\(_{1.8}\) function, we measured Na\(_{1.8}\) Na\(^+\) current in ND7-23 cells, an immortalized rat DRG/mouse neuroblastoma cell line, which is the only cell line that forms functional Na\(_{1.8}\) channels after transfection without the need...
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ND7-23 cells (Fig. 5A, middle). Analysis of the voltage-dependent activation and steady-state inactivation also showed no differences between control and AQP1-expressing cells. However, analysis of activation and inactivation kinetics of Na\textsubscript{1.8} current at 0 mV did reveal differences (Fig. 5B). The activation time (the time to peak current) was significantly shorter in the presence of AQP1, which was seen both in freshly isolated DRG neurons and transfected ND7-23 cells. The inactivation time did not differ significantly.

Patch clamp of DRG neurons in Fig. 3 showed decreased AP amplitude upon sustained stimulation, with a greater decrease seen in the AQP1\textsuperscript{--/--} neurons. Decreased AP amplitude with repetitive firing is produced by slow inactivation of TTX-R Na\textsuperscript{+} current (26). We measured Na\textsubscript{1.8} Na\textsuperscript{+} channel slow inactivation in Na\textsubscript{1.8}-expressing ND7-23 cells after repeated depolarizations to 0 mV from a −80 mV holding potential, with frequencies of 5, 10, and 20 Hz. Na\textsubscript{1.8} Na\textsuperscript{+} current amplitudes decreased progressively as a consequence of slow inactivation (Fig. 5C, top right). The reduced current at the 40th step compared with the first step was taken as a measure of slow inactivation. At 20 Hz, ∼60% reduction in Na\textsubscript{1.8} current was seen in control cells compared with ∼25% in AQP1-expressing cells, with the percentage reduction depending on frequency (Fig. 5C, top right). As a control, we studied intrinsic TTX-S Na\textsuperscript{+} current in ND7-23 cells, which has similar kinetics and activation/inactivation voltage dependence as the TTX-S current in DRG neurons (41). The TTX-S Na\textsuperscript{+} current was more sensitive to the slow inactivation protocol, with only ∼20% current remaining at the 40th pulse.

to transfact accessory subunits (41). These cells do not express AQP1 as seen by immunostaining (Fig. 5A, left) and immunoblot analysis (data not shown). For these studies Na\textsubscript{1.8} was transiently transfected (along with green fluorescent protein to identify transfected cells) into control and AQP1-expressing ND7-23 cells. The current-voltage relation of the Na\textsubscript{1.8} Na\textsuperscript{+} current was similar in the control and AQP1-expressing (Fig. 5C, bottom). However, in contrast to the TTX-R Na\textsuperscript{+} current, no significant differences in TTX-S current were found in control versus AQP1-expressing cells.

Evidence for Physical Interaction between Na\textsubscript{1.8} and AQP1—The above data indicate AQP1-modulation of Na\textsubscript{1.8} function. Both AQP1 and Na\textsubscript{1.8} are expressed in small DRG neurons. To test for physical association between Na\textsubscript{1.8} and AQP1, immunofluorescence (red) in non-transfected and AQP1 stably transfected ND7-23 cells is shown. Nuclei were counterstained blue with 4′,6-diamidino-2-phenylindole. Bar, 50 μm. Middle, shown is the current-voltage relationship of Na\textsubscript{1.8} currents in control and AQP1-stably expressing ND7-23 cells. Current was normalized to the current at 0 mV (analysis of variance, p = 0.52). Right, shown is voltage-dependent activation (G/G\textsubscript{max}) of Na\textsubscript{1.8} with fitted parameters: AQP1-expressing ND7-23 cells, V\textsubscript{1/2} = −4 ± 1 mV, k = 5 ± 1 mV, n = 8; control ND7-23 cells, V\textsubscript{1/2} = −3 ± 1 mV, k = 5 ± 1 mV, n = 8. Shown is steady-state inactivation (I/I\textsubscript{max}) with fitted parameters: AQP1-expressing ND7-23 cells, V\textsubscript{1/2} = −38 ± 2 mV, k = −10 ± 1 mV, n = 8; control ND7-23 cells, V\textsubscript{1/2} = −41 ± 2 mV, k = −10 ± 1 mV, n = 8 (t test, p = 0.35 and 0.45). B, left, examples of Na\textsubscript{1.8} current evoked by 100-ms pulses from −80 to 0 mV from control and AQP1-expressing ND7-23 cells; middle, activation time of Na\textsubscript{1.8} at 0 mV was significantly lower in AQP1-expressing ND7-23 cells and AQP1\textsuperscript{+/−} DRG neurons (S.E., t test; * p = 0.001 for the ND7-23 cell, 0.004 for DRG cells); right, inactivation time at 0 mV shows no differences (p = 0.26 for ND7-23 cells and 0.33 for DRG cells). C, left, examples of Na\textsubscript{1.8} current evoked by 30-ms pulses from −80 to 0 mV at 20 Hz and example of TTX-S Na\textsuperscript{+} current in AQP1-expressing cell. Na\textsuperscript{+} currents elicited by the 1st and 40th pulses are shown at the right. Right, TOP, peak Na\textsubscript{1.8} currents from control and AQP1-expressing ND7-23 cells (normalized to peak current from the 1st pulse) are shown as a function of pulse number and pulse simulation frequencies of 5, 10, and 20 Hz (S.E., 6 control and 6 AQP1 cells). Right, bottom, shown is normalized peak TTX-S Na\textsuperscript{+} currents at the 40th pulse at 5, 10, and 20 Hz frequencies in control and AQP1-expressing cells (S.E., t test, 4 control and 4 AQP1 cells).
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precipitations were done in ND7-23 cells at 48 h after co-transfection. As shown in Fig. 6A, top, Na$_{1.8}$ was detected in an AQP1 immunoprecipitate, and AQP1 was also detected in a Na$_{1.8}$ immunoprecipitate. Negative IgG and agarose controls indicated assay specificity. To test for involvement of the AQP1 C terminus in the AQP1-Na$_{1.8}$ interaction, immunoprecipitations were done using a series of AQP1 C terminus truncation mutants in which a c-Myc epitope was engineered into an extracellular site on AQP1. Fig. 6A, bottom, shows Na$_{1.8}$ in the c-Myc immunoprecipitates, even with the fully C terminus-truncated AQP1.T120.myc.$\Delta$41, indicating that the AQP1 C terminus is not required for Na$_{1.8}$-AQP1 association. Similar results were obtained in a separate set of co-immunoprecipitation studies done using co-transfected HEK293T cells (data not shown).

To further investigate the AQP1-Na$_{1.8}$ protein interaction demonstrated biochemically, we determined whether Na$_{1.8}$ expression would slow plasma membrane diffusion of co-expressed AQP1 in live cells. AQP1 diffusion was found previously to be rapid and unrestricted in multiple non-neural cell types (29). Single particle tracking was done to quantify the diffusion of Qdot-labeled AQP1 in ND7-23 cells. Examples of Qdot trajectories are shown in Fig. 6B, top, and analysis of many trajectories is summarized in Fig. 6B, bottom. Na$_{1.8}$ expression produced significant slowing of AQP1 diffusion in a subpopulation of AQP1 molecules, as shown in mean-squared displacement plots by a small reduction in average AQP1 diffusion coefficient and in cumulative probability distribution plots of diffusion coefficients and range at 1 s, where a distinct subpopulation of slowly moving AQP1 molecules was seen corresponding to ~30% of trajectories. These results provide evidence for AQP1-Na$_{1.8}$ interaction in live cells.

Reduced Cold Pain Perception in AQP1$^{+/-}$ Mice—The above data implicate Nav1.8 as an important determinant of AQP1-dependent DRG neuron function and pain perception. Based on a report showing that Na$_{1.8}$ is essential for the cold pain perception (27), we tested whether AQP1 deletion in mice affected cold pain perception. As summarized in Fig. 7, AQP1$^{-/-}$ mice showed remarkably reduced sensitivity to cold pain. During 5 min of observation on a 4 °C cold plate, the AQP1$^{+/+}$ litter-matched mice showed frequent jumping and lifting the whole time, whereas the AQP1$^{-/-}$ mice mainly sensed the cold pain within the first 1 min and showed little jumping/ lifting behavior after 3 min.

**DISCUSSION**

Our results establish a significant pain phenotype in AQP1$^{+/-}$ mice that was seen in assessments of acute thermal inflammatory, chemical, and cold pain. Evidence by patch clamp, immunoprecipitation, and single particle tracking analysis suggested that AQP1-Na$_{1.8}$ interaction is in part responsible for the
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Two prior studies on pain phenotype in AQP1−/− mice have reported conflicting behavioral findings. Oshio et al. (16) reported mild impairment in nociceptive response in AQP1−/− mice after thermal (tail-flick) and chemical (capsaicin) stimuli, with no differences in response to mechanical stimuli. Shields et al. (15) reported no significant differences in a series of acute and persistent behavioral pain tests. In this paper we carefully re-tested nociceptive behavior related to acute thermal and cold pain and inflammation-induced thermal pain. In agreement with the two prior studies (15, 16), we found no differences in hotplate threshold or latency or the response to formalin. We also found no differences in response to high dose capsaicin, in agreement with Shields et al (15). However, it was recently reported by the same group (35) that high dose capsaicin produces mainly mechanical rather than thermal hyperalgesia, probably by C-fiber destruction. C-fibers are involved in nociceptive thermosensation (both hot and cold), whereas mechanosensation is mediated by both non-myelinated C-fibers and myelinated A-type fibers (33). High dose capsaicin, thus, sensitizes a different subtype of DRG neurons. We found here that low dose capsaicin, which evokes a near-pure thermal hyperalgesia without C-fiber destruction (27, 33), produced much reduced pain perception in the AQP1−/− mice. Remarkable phenotype differences were also found upon challenge with bradykinin and PGE2, which are major components of the inflammatory soup and sensitize small DRG neurons through G protein-coupled receptors targeting TTX-R and TRP channels (43). Regarding PGE2 testing, Shields et al. (15) reported no differences for a hypoosmotic challenge in the setting of PGE2-induced inflammation. This response is mediated primarily by the TRPV4 (44), a channel expressed on both small and large DRG neurons (23), and cultured neuroblastoma cells (49) and is important to firing adaptation. Several molecular partners and membrane proteins have been reported to modulate Na+ channel slow inactivation, including β1−4 subunits, ankyrin G (50), and calmodulin (25). Our data suggest selective AQP1 modulation of Na+1.8 but not Na+1.7.

Na+ channels Na+1.1 to Na+1.9 are composed of a large α subunit containing the pore- and voltage-sensing machinery and several auxiliary β subunits (50). Only a few other proteins have been reported to interact directly with Na+1.8, including annexin II light chain (51), calmodulin (25), and clathrin-associated protein-1A (42). The immunoprecipitation and single molecule tracking indicated a physical interaction between AQP1 and Na+1.8. Impairment in cold pain sensing in the AQP1−/− mice provides further evidence for involvement of AQP1 in the Na+1.8 cold pain signaling pathway (27, 33).

In conclusion, we have discovered a significant pain phenotype in AQP1−/− mice that at the cell level appears to involve impairment in the firing of action potentials in small DRG neurons in AQP1 deficiency and at the molecular level appears to involve accelerated Na+1.8 Na+ channel inactivation. One clinical consequence of our results is the potential utility of AQP1 inhibitors to reduce pain nociception, which may provide a novel strategy to achieve analgesia at the presynaptic spinal level.

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