SUMMARY

Noxious pH triggers pungent taste and nociceptive behavior. While the mechanisms underlying acidic pH sensation have been extensively characterized, little is known about how animals sense alkaline pH in the environment. TMC genes encode a family of evolutionarily conserved membrane proteins whose functions are largely unknown. Here, we characterize C. elegans TMC-1, which was suggested to form a Na⁺-sensitive channel mediating salt chemosensation. Interestingly, we find that TMC-1 is required for worms to avoid noxious alkaline environment. Alkaline pH evokes an inward current in nociceptive neurons, which is primarily mediated by TMC-1 and to a lesser extent by the TRP channel OSM-9. However, unlike OSM-9, which is sensitive to both acidic and alkaline pH, TMC-1 is only required for alkaline-activated current, revealing a specificity for alkaline sensation. Ectopic expression of TMC-1 confers alkaline sensitivity to alkaline-insensitive cells. Our results identify an unexpected role for TMCs in alkaline sensation and nociception.

INTRODUCTION

The sense of pH is critical for the quality and survival of life (Holzer, 2009; Liman et al., 2014). Noxious pH produces pungent taste and also induces pain (Holzer, 2009; Liman et al., 2014). Animals have evolved a number of proton-activated channels (e.g., ASICs and TRP channels) and GPCRs to detect acidic pH (Holzer, 2009; Wemmie et al., 2006). By contrast, much less is known about how animals sense alkaline pH in the environment. For example, sensory neurons, such as trigeminal neurons, are excited by a wide range of external alkaline pH (Bryant, 2005); however, the underlying receptors/channels have not been identified. A couple of TRP channels (i.e., TRPV1 and TRPA1) are known to be activated by intracellular alkalinization but not by external alkaline pH. These channels, however, can also be activated by acidic pH (de la Roche et al., 2013; Tominaga et al., 1998) and thus do not exhibit specificity toward alkali. Some other channels, such as the constitutive-active sperm channel CatSper1 and the leaky K⁺ channels TASK/TALK, are potentiated rather than activated by alkaline pH (Kirichok et al., 2006). Apparently, unknown alkali-activated channels, which sense external alkali insults, must be present but remain to be identified.

Transmembrane channel-like proteins (TMCs) form a novel family of channel-like proteins conserved from worms to humans (Keresztes et al., 2003; Kurima et al., 2003). The mammalian genome encodes eight TMC genes, among which TMC1 and 2 are essential for hearing (Kawashima et al., 2011; Pan et al., 2013). However, their exact role in auditory transduction is unclear (Beurg et al., 2014; Kawashima et al., 2015; Pan et al., 2013), and it has not been possible to functionally express them in heterologous systems (Kawashima et al., 2015). The other six TMC genes remain uncharacterized. As such, little is known about the function and regulation of these channel-like proteins. The C. elegans genome encodes nearly all major classes of receptors and ion channels, offering an excellent genetic model for the study of their function and regulation in vivo (Bargmann, 1998). Two TMC family genes, tmc-1 and tmc-2, are present in C. elegans, of which tmc-1 has been suggested to encode a Na⁺-sensitive channel required for salt chemosensation and dietary signaling (Chatzigeorgiou et al., 2013; Zhang et al., 2015). We have characterized tmc-1 gene in the current study and found that it is required for worms to sense noxious alkaline environment mediated by the ASH nociceptive neurons. Alkaline pH evokes an inward current in ASH, which primarily depends on TMC-1 and to a lesser extent on the TRPV channel OSM-9. While OSM-9/TRPV is sensitive to both acidic and basic pH, TMC-1 demonstrates a specificity toward alkali, as it is required for alkali- but not acid-triggered behavioral response as well as electrical current in ASH. Importantly, ectopic expression of TMC-1 is sufficient to confer alkaline sensitivity to alkali-insensitive cells by promoting alkali-activated calcium transients and non-selective cation conductance. Our results identify an unexpected role of TMC-1 in alkaline sensation, demonstrate it as an essential component of a new type of alkali-activated cation channel, and also uncover a new function for TMC family genes in nociception.
ALKALINE pH-INDUCED AVOIDANCE BEHAVIOR REQUIRES TMC-1 AND OSM-9/TRPV

What mediates alkali-activated currents in ASH neurons? To address this question, we screened a select collection of ion channel mutants for defects in alkaline pH-induced avoidance behavior, including TRP, ENaC/DEG, TMC, and CNG family of channels (Figure 2). Wild-type worms began to avoid alkaline solution at pH 8.0 (Figure 3A). The response rate became saturated at pH 11 (Figure S2). As such, we focused on pH 12 for further analysis. As a positive control, osm-9 mutant worms, which are known to be defective in this behavior (Sassa et al., 2013), showed a strong defect (Figures 2, 3A, and 3B). Notably, we found that tmc-1 mutant worms were also severely defective in this avoidance behavior (Figures 2, 3A, and 3B). By contrast, loss of tmc-2 had no notable effect (Figures 2, 3A, and 3B). The guanylate cyclase Gcy-14, which mediates alkali-induced attractive responses (Murayama et al., 2013), did not have a notable role in this avoidance behavior (Figure 2). Transgenic expression of wild-type tmc-1 and osm-9 cDNA in ASH neurons was sufficient to rescue the behavioral phenotype (Figure 3B), suggesting that they act in ASH to mediate alkaline pH-induced avoidance behavior. In support of this view, laser ablation of ASH neurons led to a strong defect in this avoidance behavior (Figure 3C), which is consistent with previous results (Sassa and Maruyama, 2013). Notably, ASH-ablated worms retained a residual response to alkaline pH, indicating the presence of additional alkali-sensitive neurons, though these neurons probably only play a rather minor role (Figure 3C). This may also explain why the tmc-1;osm-9 double mutant did not exhibit a more severe behavioral phenotype than single mutants (Figure 3A). These results identify both TMC-1 and OSM-9 as important players in mediating alkaline sensation. Loss of OCR-2, another TMC-1 and OSM-9/TRPV channel.

### RESULTS

**Requirement of G Protein Signaling for TRPV/OSM-9 Channel Activation in ASH Neurons**

As TMC-1 is expressed in ASH (Chatzigeorgiou et al., 2013), a pair of polymodal nociceptive neurons, we first decided to gather some basic understandings of these neurons by characterizing their responses to noxious cues through electrophysiology. Previous work by calcium imaging showed that ASH neurons respond to a multitude of noxious stimuli such as high osmolality and various pungent chemicals (Hilliard et al., 2005). We first tested high osmolality and found that perfusion of glycerol-containing high osmotic solution to the nose tip, where the sensory endings of ASH neurons reside, evoked a robust inward current (Figure S1A). Consistent with previous calcium imaging results (Hilliard et al., 2005), we found that this current required OSM-9, a TRPV channel (Figures S1A and S1D). This current also depended on ODR-3 (Figures S1A and S1D), which is a G protein acting in sensory neurons including ASH (Roayaie et al., 1999). This supports the notion that osmosensation in ASH is mediated by G protein signaling, leading to the activation of the transduction channel OSM-9.

To provide direct evidence, we dialyzed GTPγS into ASH, a chemical that constitutively activates G protein signaling by locking G proteins in an active state. We found that it induced an inward current, showing that activation of G protein signaling was sufficient to stimulate ASH (Figures S1B and S1D). GTPγS-induced currents cannot be further enhanced by perfusion of high osmotic solution (glycerol), suggesting that the two may act in the same pathway (Figures S1B and S1D). Similarly, GTPγS-induced currents were absent in osm-9 and odr-3 mutant backgrounds (Figures S1C and S1D), indicating that these currents were mediated by ODR-3 signaling-dependent activation of OSM-9. We also performed the converse experiment by testing the effect of GDPβS, which inhibits G protein signaling by locking G proteins in an inactive state. Blocking G protein signaling with GDPβS prevented ASH neurons from being activated by high osmolality (Figures S1B and S1D). Thus, G protein signaling is necessary for high-osmolality-induced activation of OSM-9 in ASH neurons. Similar results were also obtained with some pungent chemicals such as octanol (X.W. and X.Z.S.X., unpublished data). These data suggest that the TRP channel OSM-9 can be activated through G protein signaling.

**Alkaline pH Induces an Inward Current in ASH Neurons Independently of G Protein Signaling**

As TRPV channels are also known to be directly activated by noxious stimuli (Caterina et al., 1997), we sought to search for such a stimulus for OSM-9. A prior study demonstrated by calcium imaging that alkaline pH can excite ASH neurons through OSM-9 (Sassa et al., 2013). Alkaline pH can also activate the mammalian TRP channels TRPV1 and TRPA1 (Dhaka et al., 2009; Fujita et al., 2008). We thus tested how ASH neurons react to alkaline pH by patch-clamp. Perfusion of alkaline pH solution to the nose tip activated an inward current in ASH neurons (Figures 1A and 1C). Unlike high osmolality, alkaline pH activated an inward current even in the presence of GTPγS (Figures 1B and 1C), suggesting that alkali can excite ASH independently of G protein signaling. Additional evidence came from GDPβS, which failed to block alkali-activated currents in ASH neurons (Figures 1B and 1C), though it completely abolished glycerol-evoked currents in these neurons (Figures S1B and S1D). Similarly, alkali-activated currents were nearly normal in odr-3 mutant worms (Figures 1A and 1C), yet glycerol-evoked currents were absent in this mutant (Figures S1A and S1D). Much to our surprise, alkali-activated currents remained in ASH of osm-9 mutant worms (see below), in which glycerol failed to elicit any current (Figures S1A and S1D). These observations demonstrate that alkali can excite ASH neurons independently of G protein signaling and OSM-9/TRPV channel.

**TMC-1 Plays a Critical Role in Mediating Alkali-Activated Currents in ASH Neurons**

We then assessed how tmc-1 and osm-9 genes may affect alkali-activated currents in ASH neurons by whole-cell recording. Unlike the severe behavioral deficit, mutations in osm-9 only led to a moderate reduction in alkali-activated currents in ASH (Figures 4A and 4B). This could be explained by the relative
low resolution of the behavioral assay, in which case a mild deficit in neuronal activity might lead to a severe phenotype in behavioral output. Alternatively but not mutually exclusively, OSM-9 might also act in additional neurons to mediate alkaline pH-induced avoidance behavior. By contrast, alkali-activated currents in ASH were severely diminished in \textit{tmc-1} mutant worms (Figures 4A and 4B), revealing an important role for TMC-1 in mediating this type of currents. Transgenic expression of wild-type \textit{tmc-1} cDNA in ASH rescued alkali-activated currents in \textit{tmc-1} mutant worms (Figures S3A and S3B). Consistent with the behavioral data, no defect was detected in \textit{tmc-2} mutant worms (Figures 4A and 4B), and no additive effect was observed in \textit{tmc-1};\textit{tmc-2} double mutant worms (Figures 4A and 4B), indicating the lack of a role for TMC-2 in alkali-activated currents in ASH neurons. Notably, in \textit{tmc-1};\textit{osm-9} double mutant worms, very little if any current was detected (Figures 4A and 4B), suggesting that TMC-1 and OSM-9 together are required for alkali-activated currents in ASH neurons. These data demonstrate that though TMC-1 and OSM-9 both contribute to alkali-activated currents in ASH neurons, TMC-1 apparently plays a more important role.

TMC-1 is expressed in some other sensory neurons in the amphid, such as ADL and ADF, which also send their sensory endings to the nose tip (Chatzigeorgiou et al., 2013). Like ASH, ADL is also considered a nociceptive neuron (Trommel et al., 1995). However, perfusion of alkaline pH solution to the nose tip evoked very little if any current in ADL and ADF (Figures S3C and S3E). This is consistent with the view that ASH is the primary sensory neuron detecting noxious alkaline pH and mediating avoidance responses to such noxious environment in the worm nose. Interestingly, perfusion of alkaline pH solution toward the cell body of ADL and ADF evoked robust currents (Figures S3D and S3E). These alkali-activated currents were severely diminished in \textit{tmc-1} mutant worms but were only slightly reduced in \textit{osm-9} mutants (Figures S3D and S3E), which are very similar to those in ASH neurons evoked at the nose tip. The finding that we can detect alkali-activated currents in ADL and ADF soma rather than their sensory endings suggests that TMC-1 is indeed functionally expressed in these neurons but probably not dedicated to detecting alkaline pH in the environment. This is also consistent with the notion that TMC-1 may have other functions (Zhang et al., 2015). Future effort is needed to determine the physiological roles of TMC-1 in these neurons.

**TMC-1 Is Not Required for Acidic pH-Induced Avoidance Behavior or Electrical Currents in ASH**

As TRPV channels can be activated by both acidic and basic pH (Dhaka et al., 2009; Tominaga et al., 1998), we asked whether this is also the case for TMC-1. We first tested low-pH-induced
avoidance behavior. In line with previous work (Hilliard et al., 2004), we found that worms showed a robust avoidance response to acidic pH, which saturated at pH 3 to 4 (Figure 5A). Interestingly, while osm-9 mutant worms exhibited a deficit in this behavior, tmc mutants did not (Figure 5A). In addition, tmc-1;osm-9 double mutant displayed a behavioral phenotype similar to that of osm-9 single mutant (Figure 5A), indicating the lack of a role for TMC-1 in sensing acidic pH.

We also examined acidic pH-activated currents in ASH neurons by whole-cell recording and obtained results similar to those collected by behavioral assays (Figures 5B and 5C). We conclude that while OSM-9 contributes to both acid- and alkali-activated currents in ASH, TMC-1 appears to be specifically involved in mediating alkali-evoked responses.

**NaCl-Evoked Responses in ASH Neurons Require OSM-9 and G Protein Signaling**

TMC-1 has been suggested to form a Na+-sensitive cation channel in ASH neurons, mediating salt chemosensation in C. elegans (Chatzigeorgiou et al., 2013). We thus tested the possibility whether TMC-1 can be activated by both NaCl and alkali. As previously reported (Chatzigeorgiou et al., 2013), we also found that C. elegans can detect high concentrations of NaCl, as well as Na-glucosinate by initiating reversals (Figures S4A and S5A). However, we found that such a behavioral response did not require TMC-1 or TMC-2 (Figures S4A and S5A). We tested a number of experimental conditions using different types of buffers to dissolve NaCl, including the commonly used M9 and S basal buffers and the one described previously (Chatzigeorgiou et al., 2013), as well as other types of buffers (Figures S4A and S5B–S5D). No significant defect was detected under these conditions (Figures S4A and S5B–S5D). A slight behavioral deficit was observed only when one concentration of NaCl was dissolved in pure water (Figure S4D). Thus, Na+-induced behavioral responses do not require TMCs under most conditions.
High concentrations of NaCl increase osmolarity. As OSM-9 channel mediates osmosensation, we tested osm-9 mutant worms. Loss of OSM-9 also gave rise to a strong defect in NaCl- and Na-glucolate-induced behavioral responses (Figures S4A and S5A–S5E). As osmotic activation of OSM-9 is mediated by G protein signaling, this prompted us to test odr-3 mutants. Indeed, worms lacking ODR-3 were severely defective in responding to NaCl (Figures S4A, S5D, and S5E), suggesting that Na+–induced behavioral response is primarily mediated by G protein signaling-dependent activation of the TRPV channel OSM-9.

To provide additional evidence, we recorded ASH neurons for their responses to NaCl by calcium imaging. We applied the condition described previously by perfusing bath solution containing 500 mM NaCl or Na-glucolate toward the cilia of ASH (Chatzi-georgiou et al., 2013) and found that it evoked calcium transients in ASH (Figures S4B, S4C, and S5F–S5I). However, such calcium transients did not depend on TMC-1 or TMC-2, but instead required OSM-9 and ODR-3 (Figures S4B, S4C, and S5F–S5I). To provide further evidence, we directly recorded ASH neurons for their electrical responses to NaCl by patch-clamp. As was the case with calcium imaging, alkaline pH evoked little if any current in ASI neurons (Figures 7A and 7D). In animals carrying a tmc-1 transgene in ASI neurons, alkaline pH then elicited a robust inward current in these cells (Figures 7B and 7D). By contrast, the same tmc-1 transgene did not promote acidic-pH-evoked currents in ASI neurons, demonstrating a specificity of TMC-1 toward alkaline pH (Figures S6A and S6B). We also recorded the activity of ASI in response to 500 mM NaCl (Figures S6C and S6D). Though this high concentration of NaCl induced a small current in ASI neurons, the current density was similar between wild-type and tmc-1 transgenic animals, indicating that this current was endogenous to ASI (Figures S6C and S6D). Alkali-activated inward current in TMC-1-expressing ASI neurons was nearly eliminated in NMDG-Cl solution (Figures 7C and 7D), indicating that the current was primarily carried by a cation channel. Notably, this current displayed a nearly linear I-V relationship (slightly outwardly rectifying) with a reversal potential close to zero (Figure 7E), suggesting that it was carried by a non-selective cation channel. These results demonstrate that ectopic expression of TMC-1 is sufficient to promote alkali-activated currents.

Ectopic Expression of TMC-1 Can Confer Alkaline Sensitivity to Alkali-Insensitive Cells

Having demonstrated that TMC-1 is necessary for alkaline sensitivity, we then wondered if TMC-1 is also sufficient to mediate such sensitivity. However, similar to the case with mammalian TMCs (Kawashima et al., 2011; Kawashima et al., 2015), we found that C. elegans TMC-1 was also trapped intracellularly when transfected into cell lines such as HEK293 and CHO cells (X.W. and X.Z.S.X., unpublished data). In ASH neurons, TMC-1 is localized to the plasma membrane in the sensory cilia (Chatzi-georgiou et al., 2013), suggesting that mammalian cells may lack some factors needed to transport TMC-1 to the cell surface. To overcome this difficulty, we sought to ectopically express TMC-1 in worm cells. ASI neurons came to our attention, as calcium imaging revealed that alkaline pH did not induce calcium transients in these cells (Figures 6A and 6B). We then expressed tmc-1 as a transgene in ASI neurons and found that alkaline pH evoked calcium transients in these neurons (Figures 6A and 6B). As a control, bath solution containing 500 mM NaCl failed to trigger a notable calcium response in ASI of wild-type or tmc-1 transgenic animals, though it was able to evoke such a response in ASH neurons (Figures 6C and 6D). Thus, ectopic expression of TMC-1 can confer alkaline sensitivity to alkali-insensitive cells.

**DISCUSSION**

Environmental pH is closely monitored by animals through taste and somatosensation (Holzer, 2009; Liman et al., 2014). Acidic pH induces sour taste and pain (Holzer, 2009; Liman et al., 2014). Similarly, alkaline pH also induces gustatory responses and nociception. For example, it has been known for decades that application of alkaline solutions of pH 11–12.5 to the tongue induces profound taste responses in mammals, though the cellular and molecular nature of such taste responses is not well understood.

**Figure 4. TMC-1 Plays a Critical Role in Mediating Alkali-Activated Currents in ASH Neurons**

(A) Alkali-activated currents in ASH are severely diminished in tmc-1(ok1859) mutant and nearly absent in tmc-1(ok1859);osm-9(ky10) double mutant worms. Recordings were done as described in Figure 1A. Shown are sample traces. Voltage: –60 mV. (B) Bar graph summarizing the data in (A). n ≥ 8. Error bars: SEM.
well understood (Liljestrand and Zotterman, 1956). As worms may encounter noxious pH environment in the wild (e.g., alkali soils), alkaline sensation would provide a protective mechanism for these animals to avoid such a harsh environment. While the molecular mechanisms by which animals detect acidic pH have been extensively characterized, little effort has been directed to understand how animals sense alkali in the environment. In the current study, we showed that TMC-1 plays a key role in alkaline sensation in C. elegans. Specifically, it is required for alkaline pH-triggered nocifensive behavior. Calcium imaging experiments show that ectopic expression of TMC-1 can confer alkaline sensitivity to alkali-insensitive cells, suggesting that it has the potential to function as an alkali sensor to detect noxious alkaline pH insults in the environment. In addition, TMC-1 is both necessary and sufficient to mediate alkali-activated currents, indicating that it is an essential component of a novel type of alkali-activated channel.

TMC-1 has been suggested to form a Na⁺-selective channel activated by high concentrations of NaCl (Chatzigeorgiou et al., 2013). It has also been demonstrated to be critical for salt chemosensory behavior (Chatzigeorgiou et al., 2013). It is a bit surprising that we did not detect a major role for TMC-1 in sensing high concentrations of Na⁺ or salt chemosensory behavior. On the other hand, we did identify a critical role for the TRPV channel OSM-9 in NaCl sensation. Notably, the G protein ODR-3 is also required for NaCl sensation by ASH neurons, suggesting that the NaCl sensor in ASH is probably a GPCR that is coupled to the OSM-9/TRPV channel through G protein signaling. Alternatively, as high concentrations of NaCl yield high osmolarity, the responses may result from osmosensation, consistent with the model that TMC-1-dependent currents might be directly activated by alkali. Notably, unlike TRP channels, TMC-1 is specifically required for alkali- but not proton-activated currents, demonstrating a specificity toward alkali. Ectopic expression of TMC-1 can promote alkali-evoked currents, suggesting that TMC-1 has the potential to form an alkali-activated channel. These currents are primarily carried by cations and exhibit a reversal potential close to zero, indicating that TMC-1-dependent channel is a non-selective cation channel. Nevertheless, it should be noted that TMC-1 may be incapable of forming an alkali-activated channel on its own and might instead function as an auxiliary subunit of a channel complex. Future effort is needed to address this question, and given the difficulty of expressing TMCs in heterologous cells, it may be necessary to consider alternative approaches—for example, functional reconstitution of purified TMC proteins in vitro in a lipid bilayer.

TMC-1 is a member of the TMC family of channel-like proteins, which are evolutionarily conserved from worms to humans (Keresztes et al., 2003; Kurima et al., 2003). The function and regulation of TMCs are poorly understood. Mammalian TMC1 and TMC2 are enriched in hair cells and are required for hearing, but their exact role in auditory transduction is not well understood (Beurg et al., 2014; Pan et al., 2013). Little is known about the other six mammalian TMC genes. As they are not restricted to the auditory system and are more widely expressed in different tissues, they may regulate other forms of physiological processes (Keresztes et al., 2003; Kurima et al., 2003). Our finding that C. elegans TMC-1 plays a critical role in sensing noxious alkaline environment uncovers a new function for
TMcs. We suggest that some mammalian TMcs may also function in alkaline sensation and/or other forms of nociception.

EXPERIMENTAL PROCEDURES

Strains

Wild-type: N2, TQ1784: xuEx631[Psra-6::DsRed + Prst-3::yfp2], TQ5709: osm-9(ky10) null allele. TQ6602: odr-3(n2150) null allele. TQ3367: tmc-1(ok1859) deletion allele, x4 outcrossed. TQ3369: tmc-2(ok1302) deletion allele, x4 outcrossed. TQ3369: tmc-1(ok1859); tmc-2(ok1302) x4 outcrossed. TQ548: ocr-2(ak47) deletion allele. TQ5664: osm-9(ky10); xuEx631[Psra-6::DsRed + Prst-3::yfp2]. TQ5918: odr-3(n2150); xuEx631 x4 outcrossed. TQ6111: tmc-1(ok1859); xuEx631. TQ6112: tmc-2(ok1302); xuEx631. TQ6113: tmc-1(ok1859); tmc-2(ok1302); xuEx631. TQ6122: tmc-1(ok1859); osm-9(ky10); xuEx631. TQ6855: ocr-2(ak47); xuEx631. TQ6382: tmc-1(ok1859); xuEx2201 [Psra-6::Tmc-1::DNA]; osm-9(ky10)); xuEx31. TQ6835: osm-9(ky10); xuEx2205 [Psra-6::osm-9(cDNA)]; osm-9(ky10); xuEx31. TQ5856: xuEx17976[Psra-6; Gcamp6f]; + Psra-6::DsRed]. TQ6021: osm-9(ky10); xuEx17978. TQ6114: tmc-1(ok1859); xuEx17978. TQ6115: tmc-2(ok1302); xuEx17978. TQ6857: odr-3(n2150); xuEx17978. TQ6116: tmc-1(ok1859); tmc-2(ok1302); xuEx17978. TQ6125: tmc-1(ok1859); osm-9(ky10); xuEx17978. TQ574: ocr-1(ok132) deletion allele. TQ575: ocr-3(ok1559) deletion allele. TQ684: ocr-4(e137) insertion/deletion allele. TQ223: trpa-1(ok999) deletion allele. TQ4946: trpa-2(tm3085) deletion allele. TQ17: trp-1(ok323) deletion allele. TQ194: trp-2(e901) deletion allele. TQ51: trp-3(e969) deletion allele. TQ296: trp-4(e9695) deletion allele. TQ5729: cng-1(jh111) insertion/deletion allele. TQ6003: cng-2(tm4267) insertion/deletion allele. TQ5730: cng-3(jh113) deletion allele. TQ718: tmc-2(ok869) deletion allele. TQ725: tmc-1(ok1859). TQ7259: tmc-1(ok1859); osm-9(ky10). TQ7258: xuEx22625[Psra-220::yfp]; osm-9(ky10); tmc-1(ok1859). TQ6483: xuEx2304[Pth-1(BC);::c2::mCherry2 + Punc-122:: s2::GFP]. TQ7182: xuEx2304; tmc-1(ok1859). TQ7261: xuEx2304; osm-9(ky10). TQ7260: xuEx2304; tmc-1(ok1859); osm-9(ky10).

Behavioral Assays

Avoidance behavior was performed using a drop test assay at 20°C as previously described (Mellem et al., 2002; Piggott et al., 2011). Briefly, a small drop of solution was placed on the path of a forward-moving animal. If the animal stops forward movement and also initiates a reversal that lasts at least half a head swing, it is scored as a positive response. OP50 was not included on the assay plate (NGM plate). In general, M9 buffer was made and used freshly at room temperature to avoid precipitation. To make NaCl solutions, varying amounts of NaCl were dissolved in M9, S basal, standard bath solution (see below), water, or the same buffer described previously in Chatzigeorgiou et al. (2013): 1 mM MgSO4, 1 mM CaCl2, and 5 mM KPO4. The final concentration values of NaCl shown in Figures S4 and S5 include both newly added NaCl and that pre-existing in the original buffers. To make Na-glucuronate solutions, NaCl was replaced with Na-glucuronate. Each animal was tested five times, and a response rate was tabulated for each animal.

Figure 6. Ectopic Expression of TMC-1 Can Confer Alkaline Sensitivity to Alkali-Insensitive Cells

(A and B) Ectopic expression of TMC-1 in ASH confers alkaline sensitivity to these neurons. ASH was labeled with GCaMP6 and DsRed (internal reference), which were co-expressed as a transgene using the sra-6 promoter. Bath solution (pH 12) was perfused toward the nose tip using a microfluidic system. As ASH neurons are alkali-sensitive and are also functionally connected to ASI neurons (Guo et al., 2015), imaging experiments were performed on worms with ASH ablated using a laser microbeam. tmc-1 cDNA was expressed in ASH under the sra-6 promoter. Shown in (A) are calcium imaging traces. Shades along the traces denote error bars. Bar graph in (B) summarizes the data. n ≥ 8. Error bars: SEM. **p < 0.0005 (t test).

(C and D) TMC-1 ectopically expressed in ASI does not promote NaCl sensitivity. Bath solution containing 500 mM NaCl was perfused toward the nose tip. As no notable response was detected in ASI in response to NaCl (500 mM), we imaged ASH of wild-type worms as a positive control and observed NaCl-induced calcium transients. (C) Calcium imaging traces. (D) Bar graph. n ≥ 8. Error bars: SEM. n.s.: not significant (t test).
described previously in Chatzigeorgiou et al. (2013): 1 mM MgSO4, 1 mM adjusted to 7.3). We also used the same bath solution for calcium imaging as CaCl2, and 5 mM KPO4. To make alkaline solution, the pH value of standard 1 CaCl2, 1 MgCl2, 20 glucose, and 5 HEPES (320 mOsm; pH adjusted to 9) was adjusted with Tris base to 9 and then to 12 with NaOH, and the osmolarity of the alkaline solution was also adjusted to 320 mOsm. This solution was made and used freshly.

Electrophysiology

Patch-clamp recordings were performed on an Olympus upright microscope with an EPC-10 amplifier as previously described (Kang et al., 2010; Li et al., 2011; Li et al., 2014). Worms were dissected on a sylgard-coated cover-glass. Standard bath solution was as follows (in mM): 145 NaCl, 2.5 KCl, 1 CaCl2, 1 MgCl2, 20 glucose, and 5 HEPES (320 mOsm; pH adjusted to 7.3). To make alkaline pH bath solutions, we used Tris base to adjust the pH to 9 and then to 12 with NaOH, and the osmolarity of the alkaline solution was also adjusted to 320 mOsm. This solution was made and used freshly to avoid precipitation. NMDG-Cl and Na-glucocinate bath solution was made by replacing NaCl with the same concentration of NMDG-Cl and Na-glucocinate, respectively. Normal pipette solution was as follows (in mM): 115 K-gluconate, 15 KCl, 1 MgCl2, 10 HEPES, 0.25 CaCl2, 20 sucrose, 5 BAPTA, 5 Na2ATP, and 0.5 NaGTP. To derive I-V relations, 115 mM CsCl was used to replace K-gluconate in the pipette solution, and 20 mM TEA and 5 mM 4-AP were also included to block voltage-gated potassium channels. To identify ASH and ASI neurons for recording, we labeled them with DsRed and YFP fluorescence markers expressed from a transgene xuEx631[Psra-6::DsRed + Pstr-3::yfp2]. To record ASH currents, we perfused the solution toward the nose tip where the sensory endings of ASH are localized. To record ASI neurons ectopically expressing TMC-1, we perfused alkaline pH solution toward the cell body, as this induced more robust currents. It is possible that the ectopically expressed TMC-1 was enriched in the cell body. A similar phenomenon was observed with ADL and ADF neuron recordings, in which case we detected robust currents when stimulating the cell body rather than the nose tip. Series resistance and membrane capacitance were both compensated during recording.

SUPPLEMENTAL INFORMATION

Supplemental Information includes six figures and can be found with this article online at http://dx.doi.org/10.1016/j.neuron.2016.05.023.

AUTHOR CONTRIBUTIONS

X.W. performed patch-clamp recordings and behavioral experiments and analyzed the data. G.L. carried out molecular biology, genetics, and calcium imaging experiments and analyzed the data. Jie Liu provided technical assistance on patch-clamp. X.W., G.L., Jianfeng Liu., and Z.Z.S.X. wrote the paper.

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