A uniquely adaptable pore is consistent with NALCN being an ion sensor

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NALCN is an intriguing, orphan ion channel among the 4x6TM family of related voltage-gated cation channels, sharing a common architecture of four homologous domains consisting of six transmembrane helices, separated by three cytoplasmic linkers and delimited by N- and C-terminal ends. NALCN is one of the shortest 4x6TM family members, lacking much of the variation that provides the diverse palate of gating features, and tissue specific adaptations of sodium and calcium channels. NALCN’s most distinctive feature is that it possesses a highly adaptable pore with a calcium-like EEEE selectivity filter in radially symmetrical animals and a more sodium-like EEKE or EKEE selectivity filter in bilaterally symmetrical animals including vertebrates. Two lineages of animals evolved alternative calcium-like EEEE and sodium-like EEKE/EKEE pores, spliced to regulate NALCN functions in differing cellular environments, such as muscle (heart and skeletal) and secretory tissue (brain and glands), respectively. A highly adaptable pore in an otherwise conserved ion channel in the 4x6TM channel family is not consistent with a role for NALCN in directly gating a significant ion conductance that can be either sodium ions or calcium ions. NALCN was proposed to be an expressible Gd³⁺-sensitive, NMDG⁺-impermeant, non-selective and ohmic leak conductance in HEK-293T cells, but we were unable to distinguish these reported currents from leaky patch currents (I_LP) in control HEK-293T cells. We suggest that NALCN functions as a sensor for the much larger UNC80/UNC79 complex, in a manner consistent with the coupling mechanism known for other weakly or non-conducting 4x6TM channel sensor proteins such as Na⁺_v or Ca₁.1. We propose that NALCN serves as a variable sensor that responds to calcium or sodium ion flux, depending on whether the total cellular current density is generated more from calcium-selective or sodium-selective channels.

A Brief History of NALCN Research

Hermann J. Muller, in the 1930s, came across the mutant “narrow abdomen” or “na” allele of what has been dubbed NALCN or sodium leak channel, non-selective¹ while cataloguing phials of fruit fly progeny derived from males subject to relatively high doses of radiation crossed with virgin females.² Six decades later, Howard A. Nash rediscovered Muller’s narrow abdomen phenotype in mutagenized flies that were especially sluggish in climbing up the side of the fly phial in response to halothane anaesthetic.³ The hypersensitive fly mutants mapped to an unusual four repeat channel, whose rat homolog had been cloned by Dr Edward Perez-Reyes’ lab at the University of Virginia, a year earlier in 1999.⁴ NALCN has been labeled also as “Voltage-Independent Sodium Channel 2.1” (NaVI₂.1)⁵ and “Voltage-Gated Channel-Like Protein 1” (VGCNL1) in mammals,⁶ “Unknown- or U-type” (Dmα₁U) in Drosophila⁷ and nematode cation channel (nca-1/nca-2) in C. elegans.⁸ NALCN (NA Leak Channel) received its official

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designated when one research group reported its function as a non-selective, sodium leak conductance channel, after countless others tried similar experiments for more than a decade. The features of the reported sodium leak currents are equally present in native HEK-293T cells without expressing a NALCN gene and its associated UNC79 or UNC80 subunits or SRC kinase. The reported ohmic, linear current is identical to a leak patch current from an imperfect whole cell patch clamp seal1 that can be corrected for by online P/N leak subtraction.10 The inward sodium current of a leaky patch has NALCN’s reported characteristic of disappearing when NMDG+ replaces external sodium ions and blockade of NALCN with 10 μM Gd3+. Ten micromolars Gd3+ dramatically improves the membrane seals of native HEK-293T and other patched cells, mimicking the apparent block of a linear, non-selective conductance.11 If 10 μM Gd3+ targets a specific linear conductance through a membrane channel protein, it would be hard to distinguish this current from membrane seal changes in a whole cell patch.

NALCN is attractive as a sodium leak conductance because structurally NALCN appears as a cation channel, which is voltage-independent and has the appearance of a non-selective type pore that is in between sodium and calcium channels. Furthermore, a missing sodium conductance is the simplest explanation for the hyperpolarized plasma membranes in animal mutants that lack NALCN channels such as C. elegans,3,11,12 Drosophila,3,13 snails40 and mammals.1 Many publications now have ascribed 10 μM Gd3+-sensitive sodium leak currents as confirmation of the presence of NALCN, on the assumption that 10 μM Gd3+ is a blocker of a NALCN conductance. We propose an alternative possibility, that NALCN is a weakly conducting or non-conducting channel. We glean clues as to the function of NALCN from the structural and evolutionary history of this intriguing ion channel and also collate all the common observations of NALCN mutants in C. elegans, Drosophila and mice. We come to a conclusion that NALCN might function as a variable calcium and sodium sensor protein.

Structural Features of NALCN, Calcium Channels and Sodium Channels

The origin of 4x6TM channels. The orphan NALCN and the 20 calcium and sodium channel mammalian genes belong to a structurally related superfamily of voltage-gated channel proteins with 24 membrane spanning α helices. Each cation channel is a large single polypeptide of ~1,700 to ~2,900 amino acids of four homologous domains (DI, DII, DIII, DIV) chained together by cytoplasmic linkers, with each domain containing six transmembrane helices (4x6TM) (Fig. 1) that is typified to a subunit of a “Shaker”-type voltage-gated potassium channel gene (1x6TM), which forms tetramers of four different expressed subunits. The close kinship of Domains II and Domains IV, and Domains I and III, is evidence that all members of the 4x6TM family converge from a likely common single domain ancestor, followed by a period of divergence of the domains and followed by another repeated duplication and period of divergence before speciation into the 21 4x6TM channels consisting of NALCN, sodium and calcium channels.44 4x6TM cation channels are found in photosynthetic green algae,45 and there is also a single calcium transporter in yeast16 but not found in bacteria, suggesting that the 4x6TM family may have arose in basal eukaryotes, perhaps more than once. Single-celled organisms can possess both an L-type channel and a sodium channel,17 but the earliest NALCN is found in multicellular organisms without a tissue level organization like sponges.18

The voltage sensor. The 6TM domain contributes to one quadrant of the 4x6TM cation channel pore and consists of two interdependent but separate modules, a voltage-sensor domain (transmembrane segments S1 to S4) and a pore domain (segments S5 and S6). The voltage-sensing S4 helix has repeating positive charges (of Lys+ and Arg+ amino acid residues) and an amphipathic S4-S5 linker that couples the S4 helix to gating (opening/closing) of the pore domain in response to membrane voltage changes.19 All NALCN channels from simple multicellular organisms without a tissue level organization (sponge) to humans have a similar distribution of charges in S4 helices, albeit reduced compared with most sodium and calcium channels. The fewer positive changes in the S4 helix of NALCN resemble non-voltage gated channels such as yeast calcium transporter Chc1p46 and the non-conducting salt sensor, Na+.16 (Fig. S1). It should be noted that not all S4 helices of all four domains contribute equally to voltage-sensitivity activation gating, and the consequent dependence on both the domain and position of the missing charges in the S4 helices. NALCN also has conserved counter-charges in S2 and S3 segments of the voltage-sensor domain, and an amphipathic S4-S5 linker which is critical for the mobility of the voltage-sensor of calcium and sodium channels (Fig. S1). NALCN and salt sensor Na+ may have altered voltage-sensitivity, but the level of conservation of cationic charges and countercharges in the voltage sensor domain suggest that NALCN and Na+ possess a mobile S4 helix like conventional voltage-gated sodium and calcium channels. Overexpression of a gain-of-function NALCN mutation causes overexcitability, with the mutation located in a hotspot location for gain-of-function alleles associated with altered gating of voltage-gated channels, such as slowing of inactivation.8,11,20 The NALCN gain-of-function mutation is in the proximal I-II linker adjacent to the IS6 transmembrane domain (Domain I, segment 6), where gating modifiers such as β subunits of Ca1.1 and Ca2.2 calcium channels are coupled to a rigid post 1S6 helix,21 and where a “gating brake” of Ca3.3 calcium channels is located.22 The existence of this NALCN mutation within a hotspot region associated with gating behavior, which produces an over-excitability phenotype, leads us to speculate that NALCN may have a mobile voltage-sensor to mediate its functional activation.

Models for pore selectivity. The pore domain is formed by re-entrant cytoplasmic loops spanning the S5 and S6 membrane helices of all four domains, containing a signature selectivity filter that dictates the relative permeability to sodium, calcium or potassium ions. Contributing S6 helices of each domain line the inner pore below the selectivity...
filter and form a C-terminal helical bundle that meet like an inverted teepee serving as a channel gate to occlude ion passage at their cytoplasmic ends.23

X-ray structures from Roderick Mackinnon elegantly illustrate the concept of potassium channel permeation.24 The potassium selectivity filter is made up of four rigid amino acid backbones that together mimic the hydration shell oxygen atoms surrounding potassium ions in solution.24 Optimized arrangement of surrogate oxygen groups lining the pore selectivity filter provides passage that is only energetically favorable for dehydrated potassium ions to permeate.24 The selectivity of sodium and calcium channels and NALCN is much more ambiguous because it is governed by protruding and flexible, amino acid side chains of selectivity filter residues in the pore, which produce a wider and shorter pore than the potassium channel, according to the X-ray structures of NaAb, a 1x6TM bacterial sodium channel crystalized by Bill Catterall and colleagues.23 Also notably different from the potassium permeation through potassium channels is that sodium ions are expected to permeate the pore of sodium channels in a semi-hydrated23 or hydrated state.25

**Selectivity filter residues of calcium and sodium pores.** The selectivity filter residues for ion permeation are located at the most constrictive point of the “hourglass” re-entrant (Pore-) P-loops between segments 5 and segments 6 of all four domains (Fig. 1). The three lineages of 4x6TM channels (Ca, Na, and NALCN) appear to follow universal rules, almost without exception, in eight established pore configurations, including Ca2+-selective channels with negatively-charged glutamates and aspartates (EEEE, EDEE, EEDD, DEEA) and Na+-selective channels with a positively-charged lysine (K) in either the 2nd or 3rd domain (EKEE, EEKE, DKEA, DEKA) (Fig. 2). While these four residues play critical roles in governing sodium and calcium selectivity of 4x6TM channels, each re-entrant pore is not equal in its contribution as in symmetrical (homo-multimeric) potassium channels, and furthermore the divergent residues of the re-entrant P-loops outside of the selectivity filter residues also can influence ion selectivity.26,27

The selectivity filter consists of dipole pairs of characteristic negatively charged glutamate residues projecting into pore, providing an EEEE selectivity filter generating the highly calcium-selective filter ubiquitously featured in all Ca1 and Ca2 calcium channels.28,29 Invertebrates express a unique, calcium-selective Na2 channel with a DEEA pore.30 A more sodium selective channel is generated by replacing the negatively charged glutamate residue with a lysine residue in either Domain II or Domain III.31 The lysine residue governs the highly sodium selective, DEKA configuration of the classical Nav1 sodium channels that generates the upstroke of the action potential,28,31 and what is considered a slightly less sodium-selective, DKEA configuration found in Na2 channels of basal cnidarians.30,26 NALCN has indeterminate pores that can both resemble calcium channels or sodium channels, or both.9 Basal NALCN channels in radially symmetrical animals (sponge/Trichoplax/Cnidaria) all have calcium-like EEEE pores.9,18

**Variability in calcium and sodium pores of 4x6TM channels.** Different invertebrates groups appear to have independently evolved NALCN channels with alternative calcium-like EEEE and sodium-like EKEE or EEKE pores.9 Alternative EEEE and EKEE pores

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**Figure 1.** Alternative splicing in NALCN generates selectivity filters that resemble a calcium channel pore (EEEE) or sodium channel pore (EKEE/EEKE), by alternative splicing of either exon 15 or exon 31 in different animal lineages.
Notable NALCN Features

NALCN evolved with the advent of multicellularity in animals. A NALCN gene is a required gene in mammals and present in every known animal species, beginning with the simplest multicellular organisms, such as sponge and...
the placozoan *Trichoplax*. Single-celled organisms closely related to animals lack NALCN, such as the ciliate paramecium, which has an EEEE L-type calcium channel that functions in turning movements, and protozoans that have a DEEA Na+1 type sodium channel. NALCN is in between sodium and calcium channels in overall structure and may have evolved from an ancestral channel resembling the 4x6TM calcium transporter Cch1 found in the sister group to the animals, the fungi.

NALCN is linked to the interconnectivity between cells in multicellular animals. The emergence of NALCN in primitive multicellular organisms without organized tissues and not single-celled organisms may relate to a role for NALCN in the interconnectivity of cells. Interestingly, NALCN in invertebrates at least is functionally linked to a gap junction hemichannel, innexin (UNC7). NALCN evolved in a parallel lineage from ancestral multicellular organisms and did not emerge from a specialized branch of sodium channels or calcium channels. NALCN first appears in extant relatives of the most basal multicellular organisms without organs, and thus, evolved differently from the highly-specialized vertebrate-specific genes like the non-conducting Na+ salt sensor for the subfornical (a brain circumventricular) organ and weakly conducting Ca+1.1 channel for muscle. NALCN adapted as sensor proteins for functions in vertebrates, losing a primary purpose in ion conduction that was present in their forbears that were likely both voltage-gated and ion conducting, Na+1 and Ca+1 channels in basal organisms respectively. NALCN is highly conserved in the simple multicellular organisms which lack organs, so its fundamental role likely is not associated within specialized intracellular signaling associated with the organ function.

NALCN has resisted gene duplication compared with other members of the 4x6TM family and is uniquely, a single copy gene in most animals. Although NALCN is required in all multicellular organisms, and it is noteworthy that NALCN is mostly a single copy gene in every animal, with exceptions such as sponge (*Amphimedon*), anthozoan (*Nematostella*) and nematode (*Caenorhabditis*), which contain two mostly redundant NALCN gene copies. While NALCN has largely resisted increases in gene numbers, sodium and calcium channels expanded from five genes in invertebrates to 20 different mammalian genes. Many of the mammalian sodium and calcium channels are selectively expressed for different functions in different cells such as skeletal muscle (Ca+1.1, Na+1.4), neurons (Ca+2.1 and Ca+2.2, Na+1.1, Na+1.2, Na+1.3, Na+1.6, Na+1.7, Na+1.8, and Na+1.9), retinal and leukocytes (Ca+1.4), heart muscle (Na+1.5), inner hair cells (Ca+1.3) and subfornical organ (Na+). NALCN is a highly conserved ion channel between animal groups. NALCN is more highly conserved across the whole protein from end to end from sponges to humans than are other 4x6TM channels. Conversely, sodium and calcium channels have undergone an accelerated rate of evolution, associated with the rising complexity of cellular signaling in higher invertebrates and vertebrates. NALCN’s lack of variability suggests its general functions have little changed from its origins in simple multicellular organisms without significant tissue level organization such as sponge or the placozoan, *Trichoplax*.

NALCN protein is short and lacking of the typical variation in cytoplasmic regions common in other 4x6TM channels. 4x6TM channels differ from the tetrameric 1x6TM channels in having, single N- and C-termini and inter-Domain II-II and II-III cytoplasmic linkers serving as protein interaction platforms adapted for different intracellular environments. NALCN is the second shortest 4x6TM channel in total size (1738 amino acids) because of the shortness of its cytoplasmic N- and C-terminal ends and the I-II and II-III linkers. The shortest full-length 4x6TM channel (1682 amino acids) is the highly specialized, non-conducting sodium channel Na+ that serves as a sodium sensor in glial cells and couples to the α subunit of the Na+/-K+/ATPase, to transduce a signal that regulates salt appetite in the subfornical organ. The difference between the largest (LCa+3, 2,886 amino acids) and smallest (Na+ or NALCN) 4x6TM channels is an extra ~70% of the whole channel length. The shortness and lack of variability in NALCN channel, specifically in the cytoplasmic regions, may suggest that NALCN has a more limited function like the sensor protein Na+.

Large UNC80/UNC79 proteins coupled to NALCN are unconventional compared with the small accessory proteins of most voltage-gated channels. Most of the 4x6TM channels outside of Ca+3 T-type channels are known to be associated with accessory subunits. These subunits in vertebrate calcium channels (αδ, β, γ) and sodium channels (β1, β2, β3, β4) are smaller than the conducting channel α subunits. These accessory subunits are promiscuous in being able to bind to several different, related cation channel types. They serve to facilitate membrane expression (and can prevent protein degradation) and affect the biophysical features of the ion conducting channels in a manner similar to 1x6TM potassium channel β subunits. NALCN channels require accessory subunits for function too, but these are unlike the sodium and calcium channel accessory subunits. NALCN (1738 aa) is associated with cytoplasmic protein, UNC80, which is twice its size (3258 aa). UNC80 itself is coupled to UNC79, which is also larger than NALCN (2635 aa). NALCN, UNC80 and UNC79 are contiguous proteins, and the mutant phenotypes of these three proteins overlap. NALCN constitutes less than 1/4 of the size of the NALCN/UNC80/UNC79 complex. The majority or 3/4 of the size of the NALCN complex embodies large interconnected, cytoplasmic proteins UNC80 and UNC79. A typical sodium channel β subunit in comparison is a miniscule, 1/10 of the size of a voltage-gated sodium channel. It seems more probable that NALCN is the external calcium or sodium ion sensor for the intracellular signaling mechanism performed by the much larger UNC80/UNC79 complex, than to conceive of a model where NALCN functions mostly as an independently conducting and signaling unit.

NALCN coupling to intracellular UNC80/UNC79 complex is reminiscent of known ion sensor proteins, Cav1.1 and Nax. The small NALCN channel
intimately coupled to a larger UNC80-UNC79 intracellular protein complex is reminiscent to calcium channel Ca1.1, for example, which is adapted as a specialized voltage-sensing receptor of the T-tubule junction membrane.\textsuperscript{34} Ca1.1 (1873 aa) is bi-directionally coupled to a much larger intracellular sarcoplasmic reticulum's ryanodine receptor subunit (~5,000 aa).\textsuperscript{34} The calcium entry through Ca1.1 is minor, and the channel's primary purpose is signaling voltage changes to the massive web of ryanodine receptor-gated, intracellular calcium, surrounding muscle fibers.\textsuperscript{34} Of all the 4x6TM channels, NALCN appears to have more in common with the non-conducting Na\textsuperscript{+} salt sensor in its small size, lack of variability and reduced voltage sensor,\textsuperscript{35} and also similar to the highly specialized and weakly-conducting voltage sensor protein, Ca1.1 whose function is inseparable from the much larger calcium conducting ryanodine receptor subunit.\textsuperscript{34}

UNC80 and UNC79 have not provided insights into how NALCN works. So far, no functions are ascribed to NALCN’s partners UNC80 and UNC79. UNC80 and UNC79 are highly disordered proteins that likely bear flexible docking sites for multiple protein interactions. Neither UNC80 nor UNC79 are classified within known protein families.

A variable sodium and calcium pore is not consistent with NALCN being an ion conducting channel. The argument for NALCN gating a significant ion conductance on its own (i.e., ionotropic effect) is difficult to reconcile, when it has a variable pore for ion selectivity that can be calcium-selective or sodium-selective in different animal species.\textsuperscript{9} NALCN cannot be a ubiquitous sodium leak channel, when NALCN likely evolved in basal animals with a calcium-selective EEEE pore. These simple animals do not possess a NALCN pore that resembles a sodium channel EKEE or EEKE.\textsuperscript{9} While both sodium and calcium ions can depolarize membranes, these are not interchangeable conductances, where the impact of the relatively inert and highly abundant Na\textsuperscript{+} ion serves mostly an electrogenic role, while Ca\textsuperscript{2+} influx at the same levels is highly toxic to cells, and serves as an exquisitely-sensitive signaling molecule.

Instead, variability in the pore suggests that ion permeation is somewhat inconsequential for NALCN, which has permitted unprecedented levels of variability in the selectivity filter without dramatically impacting the physiology of cells.

**NALCN lacks the abundant alternative-splicing associated with most other 4x6TM channels.** Each 4x6TM channel usually possesses significant (sometimes > 100 different) alternatively-spliced isoforms,\textsuperscript{46} and many of these are designed to fine-tune the ion channel gating and expression characteristics. Sodium and calcium channels are limited to specified voltage ranges, which shape their signal specificity. NALCN lacks the extensive alternative-splicing of other 4x6TM channels.

A key feature of NALCN is its variable and adaptive pore. Invertebrate NALCN from snail *Lymnaea stagnalis* possess no significant gene splicing other than the alternative exon that splices in a novel pore to change ion selectivity.\textsuperscript{9} The variable and adaptive pore is the distinctive feature of NALCN, as is the variable-splicing that generates channel gating changes in voltage-sensitive 4x6TM channels. Mutually-exclusive splicing to generate alternative EEEE calcium and EEKE or EKEE sodium pores evolved twice independently within the two major lineages of invertebrates, the lophotrochozoans-deuterostomes (mollusks/annelids; echinoderms/hemichordates) and the ecdysozoa (centipedes, arachnids) (Fig. 1).\textsuperscript{9} Evolutionary pressure was present to duplicate exons for generating alternatively-spliced pores but to also retain both alternatives pores in many animal groups including deuterostomes.

**Dual alternative NALCN calcium and sodium pores are adapted for differing tissue functions in invertebrates.** Expression patterns of the EKEE sodium pore and the EEEE calcium pore are equally abundantly-expressed overall in the snail *Lymnaea stagnalis*.\textsuperscript{9} Snail NALCN with a sodium pore is most abundant in secretory brain and glandular tissue, which correlate with cells containing Na\textsubscript{1} channels and sodium-dependent membrane fluxes generating action potentials.\textsuperscript{9} Snail calcium pore is associated with contractile heart and skeletal muscle, where there are greater requirements for general calcium-sensing.\textsuperscript{9} The heart of snails, for example, completely lack sodium-dependent action potentials and expression of Na\textsubscript{1} sodium channels.\textsuperscript{27} NALCN is thus likely working mostly as a calcium sensor in muscle, to sense the ion fluxes during contraction that are mostly carried by calcium ions.

NALCN could still be a regulated channel even if it were non-conducting or weakly-conducting. Structurally, NALCN has reduced S4 charges in the voltage-sensor relative to standard voltage-gated sodium and calcium channels, but this is as true for NALCN from the simplest basal animals, such as sponge, as it is for human NALCN. An overall level of conservation of cationic charges and countercharges in the voltage sensor domain, and conservation of an amphipathic S4-S5 linker (Fig. S1) suggests that NALCN is like other 4x6TM channels in containing mobile elements that are triggered upon activation.

Other known leak conductance channels are regulated leak channels. It is hard to imagine NALCN, the singleton sodium leak channel, as the depolarizing counterpart to the two-pore, KCNK potassium leak conductance channels. There are more than 50 different KCNK genes in animals, and at least 18 human KCNK genes.\textsuperscript{45} KCNK leak conductance channels are governed by fundamental cellular conditions such as membrane stretch, external pH and temperature, are variable in their conductances and are often rectifying channels.\textsuperscript{57} NALCN was first described as a linear, non-selective, ohmic conductance at all voltage ranges,\textsuperscript{27} but it has also been asserted that NALCN can be receptor-operated.\textsuperscript{42,48} A lack of gating mechanism would mean in essence, that NALCN is more like an unregulated hole in the membrane when open, draining membrane gradients at high metabolic cost to ATP driven pumps and dampening activity through shunting inhibition because of the membrane’s lowered input resistance.

NALCN with a calcium or sodium pore correlates with animals that rely on calcium or sodium ion flux, respectively. It appears that NALCN’s changeable calcium and sodium pores correlate
with the changing roles of calcium and sodium ions as signaling molecules in animal evolution (Fig. 2). The first appearance of a 4x6TM channel with a lysine in the pore (DKEA), is in cnidarians, which are the most primitive animal group with a nervous system that contains sodium-dependent action potentials.\(^9\) Prior to the cnidarians, all calcium channels, sodium channels and NALCN have calcium pores (EEEE or DEEA). Invertebrate 4xTM channels then evolved both calcium pores (EEEE/EEDD/DEEA) and sodium pores (DEKA/EKEE/DKEA/EEKE).\(^9\) Invertebrates have greater promiscuity between sodium and calcium selectivity within the major 4x6TM channels. They have both sodium-selective Na\(_1\) (DEKA)\(^9\) and an invertebrate-specific calcium-selective sodium channel gene Na\(_2\) (DEEA).\(^8\) Ca\(_3\) T-type channels in invertebrates have a differing sodium and calcium ion permeability through alternative splicing, which is not present in the more exclusively calcium-selective Ca\(_3\) channels in vertebrates.\(^27\) The greater flexibility in ion selectivity may relate to the presence of \(-4\times\) fewer numbers of 4x6TM channels to draw upon in invertebrate genomes for structural and functional diversification.

Many animal species have lost NALCN with a calcium pore. The calcium pore is lost in insects and nematodes as well as many parasitic species of flatworms, arachnids and annelids. Free-living forms, closely related to the parasites that have lost the calcium pore, maintain dual calcium and sodium pores. The loss of selective ion sensing in endoparasites may reflect that they live within the regulated environment of their host with a more limited requirement for NALCN sensing.\(^9\) In higher vertebrates, NALCN is mostly a brain protein but is also can be found in heart tissues (particularly atria) and glandular tissues, including the thyroid gland, the adrenal gland and the pancreas.\(^7,48\) Vertebrates lost the more muscle-specific, NALCN spliced calcium-sensing pore, which may relate to the decreasing reliance of plasma membrane calcium flux through L-type calcium channels in the progressive evolution of muscle excitation-contraction coupling.\(^49\)

NALCN is concentrated in cells with rapid ion flux rates. NALCN loss of function phenotypes are prominently associated with defective pacemaking in rhythmically oscillating neurons,\(^1,8,12,13,44\) and there are defects evident in secretion especially in cells with fast or continuous vesicle turnover.\(^8,11\) A potential adaptable sodium and calcium sensing pore of NALCN may play a role in particular cells with rapid ion fluxes such as pacemaker tissue, where there would be the greatest need to monitor and respond to changing calcium and sodium concentrations across cell membrane and between cells through gap junctions.

There are links between NALCN and energy metabolism. NALCN/UNC80/UNC79 knockouts do not result in drastic developmental defects in the embryo,\(^52,54\) suggesting that the NALCN complex does not have a critical role in development. NALCN appears to have an association with metabolism and mitochondrial function. Defects in the NALCN complex produce animals that are smaller,\(^5,45\) consume more energy\(^45\) and are more hypersensitive to alcohol and volatile anesthetics.\(^55\) It is tempting to speculate that NALCN coupled to UNC80/UNC79 relates to a defect in mitochondrial signaling, linking a sensing of rapid turnover of membrane cation flux with energy usage and generation.

**Conclusions**

Many different channel conductances active at low membrane potentials near rest are known to contribute to pacemaking (e.g., T-type channels and HCN channels), but these are highly-regulated and diverse and do not resemble a non-specific, linear, ohmic leak conductance. NALCN-UNC80 complexes influence membrane conductances, suggesting a potential leak channel role,\(^56,57\) but the difficulties and elusiveness of locating expressed NALCN currents are more suggestive that its major function might not be its ion conductance. Rather, NALCN’s distinctiveness is in its changeable pore, which mimics the pores that differentiate the classes of calcium- and sodium-selective channels. Appropriately, NALCN is in between sodium and calcium channels in overall structure and evolved after them in likely ancestors of extant multicellular organisms. NALCN’s adaptable pore switches with the changing dominance of calcium and sodium fluxes in primitive to more advanced animals and evolves a dual pore in two different lineages of animals to serve differing proportions of calcium and sodium fluxes required in different tissues of many animals. NALCN is a ubiquitous and likely critical gene in all multicellular animals, but it has resisted changes in gene numbers and diversity of form that is the hallmark of voltage-gated sodium and calcium channels. NALCN, with a similar overall structure as other 4x6TM channels, evolved from a calcium sensor in primitive, radially symmetrical animals, to dual calcium and sodium sensor in different tissues of more advanced bilaterally symmetrical animals\(^8\) to mostly serving as a sodium sensor in the mammalian brain.\(^53\) The animal knockdown of NALCN is perinatal lethal, not embryonic lethal in mammals, which may relate directly to a reduced electrical activity in spinal nerves innervating the diaphragm.

A future model of NALCN function will have to tie all the known attributes of NALCN together. NALCN possesses an adaptable calcium and sodium sensing pore and is enriched in pacemaking cells with rapid ion fluxes. NALCN’s activation likely involves a mobile voltage-sensor domain that is coupled through a large,
cytoplasmic UNC80/UNC79 complex to possible functions, which may include a homeostatic regulation of sodium or calcium ions, and energy utilization. Overall, there is no definitive proof that NALCN is an ion sensor, but at least one should consider this as an alternative possibility to the prevailing viewpoint that NALCN is a non-selective, ohmic leak conductance to the prevailing viewpoint that NALCN is calcium ions, and energy utilization. Overall, there is no definitive proof that NALCN is an ion sensor, but at least one should consider this as an alternative possibility to the prevailing viewpoint that NALCN is a non-selective, ohmic leak conductance to the prevailing viewpoint that NALCN is calcium ions, and energy utilization. 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40. Altier C, Garcia-Caballero A, Simms B, You H, Chen L, Walcher J, et al. The Cavβ subunit prevents RFP2-mediated ubiquitination and proteasomal degradation of L-type channels. Nat Neurosci 2011; 14:733-80; PMID:21186355; http://dx.doi.org/10.1038/nn.2712
41. Pong O, Schwarz JR. Ancillary subunits associated with voltage-dependent K+ channels. Physiol Rev 2010; 90:755-96; PMID:20393197; http://dx.doi.org/10.1152/physrev.00020.2009
42. Lu B, Su Y, Das S, Wang H, Wang Y, Liu J, et al. Peptide neurotransmitters activate a cation channel complex of NALCN and UNC-80. Nature 2009; 457:741-4; PMID:19092807; http://dx.doi.org/10.1038/nature07579
43. Lu B, Zhang Q, Wang H, Wang Y, Nakayama M, Ren D. Extracellular calcium controls background current and neuronal excitability via an UNC79-UNC80-NALCN cation channel complex. Neuron 2010; 68:488-99; PMID:20408499; http://dx.doi.org/10.1016/j.neuron.2010.09.014
44. Pierce-Shimomura JT, Chen BL, Mun JJ, Ho R, Sarkis R, McIntire SL. Genetic analysis of crawling and swimming locomotory patterns in C. elegans. Proc Natl Acad Sci U S A 2008; 105:20982-7; PMID:19074276; http://dx.doi.org/10.1073/pnas.0810359105
45. Specia DJ, Chihara D, Ashique AM, Bowers MS, Pierce-Shimomura JT, Lee J, et al. Conserved role of unc-79 in ethanol responses in lightweight mutant mice. PLoS Genet 2010; 6:6; PMID:20714347; http://dx.doi.org/10.1371/journal.pgen.1001057
46. Lipscombe D, Andrade A, Allen SE. Alternative splicing: Functional diversity among voltage-gated calcium channels and behavioral consequences. Biochim Biophys Acta 2012; pii: S0005-2736(12)00539-2; PMID:23022282; 10.1016/j.bbamem.2012.09.018
47. Enyedi P, Czirják G. Molecular background of leak K+ currents: two-pore domain potassium channels. Physiol Rev 2010; 90:559-605; PMID:20393194; http://dx.doi.org/10.1152/physrev.00029.2009
48. Swayne LA, Mezghrani A, Varrault A, Chemin J, Bertrand G, Dalle S, et al. The NALCN ion channel is activated by M3 muscarinic receptors in a pancreatic beta-cell line. EMBO Rep 2009; 10:873-80; PMID:19575010; http://dx.doi.org/10.1038/embor.2009.125
49. Spafford JD, Spencer AN, Gallin WJ. A putative voltage-gated sodium channel alpha subunit (PpSCN1) from the hydrozoan jellyfish, Polyorchis penicillatus: structural comparisons and evolutionary considerations. Biochem Biophys Res Commun 1998; 244:772-80; PMID:9593741; http://dx.doi.org/10.1006/bbrc.1998.8332
50. Schredelseker J, Shrivastav M, Dayal A, Grabner M. Non-Ca2+-conducting Ca2+ channels in fish skeletal muscle excitation-contraction coupling. Proc Natl Acad Sci U S A 2010; 107:5658-63; PMID:20212109; http://dx.doi.org/10.1073/pnas.0911531105
51. Sinko AP, Caputo C, Tsaih SW, Yuan R, Ren D, Deen PM, et al. Genetic analysis of mouse strains with variable sodium concentrations identifies the Nalcn sodium channel as a novel player in osmoregulation. Physiol Genomics 2011; 43:205-70; PMID:2177381; http://dx.doi.org/10.1152/physiolgenomics.00018.2010
52. Lu B, Su Y, Das S, Liu J, Xia J, Ren D. The neuronal channel NALCN contributes resting sodium permeability and is required for normal respiratory rhythm. Cell 2007; 129:371-83; PMID:17448995; http://dx.doi.org/10.1016/j.cell.2007.02.041
53. Nakayama M, Iida M, Koseki H, Ohara O. A gene-targeting approach for functional characterization of KIAA genes encoding extremely large proteins. FASEB J 2006; 20:1718-20; PMID:16807365; http://dx.doi.org/10.1096/fj.06-5952fje
54. Specia DJ, Chihara D, Ashique AM, Bowers MS, Pierce-Shimomura JT, Lee J, et al. Conserved role of unc-79 in ethanol responses in lightweight mutant mice. PLoS Genet 2010; 6:6; PMID:20714347; http://dx.doi.org/10.1371/journal.pgen.1001057
55. Humphrey JA, Hamming KS, Thacker CM, Scott RL, Sedensky MM, Snutch TP, et al. A putative cation channel and its novel regulator: cross-species conservation of effects on general anesthesia. Curr Biol 2007; 17:624-9; PMID:17350263; http://dx.doi.org/10.1016/j.cub.2007.02.037
56. Lu TZ, Feng ZP. NALCN: a regulator of pacemaker activity. Mol Neurobiol 2012; 45:415-23; PMID:22476981; http://dx.doi.org/10.1007/s12035-012-8260-2
57. Lu TZ. Sodium leak channels in neuronal excitability and rhythmic behaviors. Neuron 2011; 72:899-911; PMID:22196327; http://dx.doi.org/10.1016/j.neuron.2011.12.007