Sensing voltage across lipid membranes

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Abstract

The detection of electrical potentials across lipid bilayers by specialized membrane proteins is required for many fundamental cellular processes such as the generation and propagation of nerve impulses. These membrane proteins possess modular voltage-sensing domains, a notable example being the S1-S4 domains of voltage-activated ion channels. Ground-breaking structural studies on these domains explain how voltage sensors are designed and reveal important interactions with the surrounding lipid membrane. Although further structures are needed to fully understand the conformational changes that occur during voltage sensing, the available data help to frame several key concepts that are fundamental to the mechanism of voltage sensing.

Most of us appreciate the importance of electricity at a young age. It powers many useful and entertaining man-made devices; a light for seeing the world, a computer for searching the internet, or an omnipresent iPod for listening to music. The concept that living organisms use electricity as a fundamental mechanism for signaling across membranes and between cells is less widely appreciated, even though scientists have been thinking about the biological roles of electricity since the late 18th century. Indeed, Galvani’s experiments showing that spark generators can elicit contraction of frog muscles were contemporary with key developments in understanding the phenomenon of electricity itself. Today we know a great deal about the biological roles of electrical signals and it would be impossible to overstate their profound importance. In humans, electrical signals are used to complete a computation within the cerebral cortex, to secrete insulin after a meal, or to signal that a sperm has entered an egg and for embryogenesis to commence.

In cells, electrical forces arise from the separation of Na+, K+, Ca2+ and Cl− across a lipid membrane that is intrinsically highly impermeable to these ions. ATP is consumed by pumps to produce concentration gradients across the membrane for these ions, creating a chemical potential that can generate an electrical potential when ion-selective channel proteins open and provide pathways for these ions to move down their concentration gradients. In a quiescent neuron, for example, K+ channels most commonly establish an electrical potential, or voltage, that is negative inside relative to outside because the concentration of K+ is high inside the cell and low outside. A cell membrane can be depolarized (less negative inside) when neurotransmitters open ion channels that are permeable to Na+ or Ca2+ because the concentrations of these ions are high outside and low inside. For such electrical potentials to generate action potentials1, to trigger neurotransmitter secretion or initiate muscle contraction, voltage-activated ion channels must sense the voltage across the cell membrane and react by opening or closing an ion conduction pore. In these instances, changes in membrane voltage are detected by voltage sensors that are intrinsic to the ion channel; in most instances the voltage sensors move between ‘resting’ and ‘activated’ states to open the pore when membrane voltage becomes depolarized, but there are varieties that trigger pore opening when the membrane

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becomes hyperpolarized (more negative inside). Investigation into the mechanism of voltage sensing has progressed rapidly since the classical studies of Hodgkin and Huxley on the Squid giant axon\(^1\), with the cloning of the voltage-activated Na\(^+\) (Nav), K\(^+\) (Kv) and Ca\(^{2+}\) (Cav) channels in the late 1980's\(^2-5\) and the first X-ray crystal structure of a Kv channel in 2003\(^5\) representing historical landmarks. Although many mechanistic aspects of voltage sensing are debated, this review focuses on the emerging mechanistic concepts for which there is growing consensus. I will begin by discussing the architecture and structure of voltage-sensing domains in Kv channels, where our understanding is most advanced, and outline four key concepts that emerge from both functional and structural studies. I will end by discussing how voltage sensors might move within the membrane and highlight exciting directions for future research.

### Voltage-sensing domains in membrane proteins

The best understood voltage sensors are found in the voltage-activated ion channels that are so crucial for electrical signaling in the nervous system. When these channels were first cloned, the fourth putative membrane-spanning segment, termed S4, stood out because it contains many basic Arg or Lys residues that are capable of carrying a positive charge\(^2-4\), \(6, 7\). The S4 segment is always found together with three other transmembrane segments (S1-S3) that contain negatively charged acidic residues, and the four segments together collectively form an S1-S4 voltage-sensing domain (Fig 1a). The concept that S1-S4 is a domain in the formal sense grew out of comparisons of Kv channels and other simpler K\(^+\) channels that lack the S1-S4 region\(^8\), experiments identifying important acidic residues in S2 and S3\(^9\), and the discovery that tarantula toxins can bind to S1-S4 domains in different types of voltage-activated ion channels\(^10\). That S1-S4 voltage sensors are functionally independent protein domains was demonstrated by Lu and colleagues when they found that the S1-S4 region of a Kv channel could endow the KcsA K\(^+\) channel with strong voltage-sensitivity\(^11, 12\). Bacteria can synthesize, properly fold and insert isolated S1-S4 voltage sensing domains into their membranes\(^5, 13\), further supporting the idea that S1-S4 is an independent domain.

For S1-S4 voltage sensing domains to sense voltage across membranes, they must change conformation between states where charged residues or dipoles have different relative free energies. In principle, this conformational change could result from charges moving through a stationary electric field or the electric field reshaping around fixed charges. An important advance in studying this conformational change was the discovery that the movement of charges relative to the field could be detected electrically as a non-linear capacitive current, or gating current\(^14-16\). Measurements of this sort led to two key conclusions about the conformational change occurring when voltage sensors activate. First, the conformational change involves the movement of 3 to 3.5 elementary charges relative to the full membrane electric field for each S1-S4 voltage sensor\(^17-21\). Second, the external four Arg residues in S4 (Fig1b) carry most of the measured gating charge\(^18, 19, 21\). Although reshaping of the electric field may play a role, it is clear that S4 moves outward in channels that activate with membrane depolarization\(^22-37\) (see Voltage sensor motions).

While S1-S4 voltage-sensing domains were originally thought to only exist in voltage-activated ion channels, Okamura's group made the seminal discovery that S1-S4 domains exist in proteins that lack associated ion conducting pores\(^38\). The Ciona intestinalis voltage-sensitive phosphatase (Ci-VSP), for example, uses an S1-S4 domain to control the hydrolysis of membrane phosphoinositides by an associated phosphatase domain\(^38, 39\) (Fig1a). Another S1-S4 domain protein without a separate pore domain was shown to function as a voltage-activated proton channel (Hv1)\(^40-42\) (Fig1a). In this case, the S1-S4 domain appears to provide the proton permeation pathway (see below). One of the interesting differences between voltage-activated ion channels and the other voltage-sensing proteins is the number of S1-S4 domains they contain. There are four in Kv, Nav and Cav channels,\(^2-5\) two in the proton channel\(^42\).
and only one in the phosphatase. Although much remains to be learned about these fascinating new voltage sensing proteins, key regions can be swapped with Kv channels without disrupting function or pharmacological sensitivities, suggesting that their basic voltage sensing mechanisms are related.

Structures of voltage-sensing domains

For many years, the lack of three dimensional structures of voltage sensors was a barrier to understanding the mechanism of voltage sensing. At present, four X-ray structures have been solved which contain S1-S4 voltage sensing domains. As will be discussed below, these structures are probably most relevant for thinking about the activated state of voltage sensors that exist at depolarized membrane voltages. The first two structures reported in 2003 by MacKinnon and his colleagues were of the isolated S1-S4 domain and full-length channel of KvAP, an archaeabacterial Kv channel from Aeropyrum pernix. The structure of the isolated S1-S4 domain comprises five helices, with S1, S2 and S4 forming long uninterrupted helices and S3 containing two shorter helices, consistent with helix scanning studies in eukaryotic Kv channels. Because this domain was crystallized without the pore domain, it wasn't obvious how it is oriented with respect to the pore or the surrounding membrane.

The structure of the full-length KvAP channel didn't answer this question because the voltage-sensing domains in this structure are distorted. However, comparisons of the structures of the isolated S1-S4 domain and the full-length channel led MacKinnon and colleagues to three important conclusions. First, both structures contain a conserved helix-turn-helix motif that is composed of the S3b and S4 helices, termed the voltage sensor paddle. This motif contains the most crucial S4 Arg residues and is widely targeted by tarantula toxins that inhibit voltage-activated ion channels by binding within the membrane. Second, the distortions evident in the KvAP full-length structure suggest that S1-S4 domains are inherently flexible and lack tight attachments to the surface of the pore domain. In other words, the 'malleability' of voltage-sensors reveals an underlying structural flexibility that may be related to how these domains function. Third, the paddle motif moves at the periphery of the channel protein in contact with the surrounding lipid membrane. As wonderful as it was to 'see' the first structures of voltage sensors and to ponder these new ideas, many questions remained unanswered. How distorted is the structure of the S1-S4 domain in the full-length KvAP channel? How representative is the structure of the isolated S1-S4 domain of KvAP, and how is this domain oriented with respect to the central pore domain?

The X-ray structure of the eukaryotic Kv1.2 channel provided the first clear perspective of S1-S4 domains in a full-length channel and revealed how they are oriented with respect to the central pore domain and the surrounding lipid membrane. Even though the quality of electron density within the S1-S4 domain is relatively weak (mean B factor = 162 Å²), leaving many residues unresolved, the S1 through S4 helices clearly adopt transmembrane orientations and appear loosely attached to the central pore domain. The most recent X-ray structure of the Kv1.2 channel containing the paddle motif from the Kv2.1 channel (paddle-chimera channel) currently provides the clearest picture of voltage sensor structure in a full-length channel (Fig2). The close apposition of the S3b-S4 helices are similar to what is seen for the paddle motif in the KvAP structures, S1 and S4 are positioned adjacent to the pore domain and both S2 and S3 are positioned peripherally. Electron densities for many lipid molecules are also present in the maps for the paddle-chimera channel, revealing that lipids interact intimately with the channel. Lipids are present between the S1-S4 voltage-sensing domain and the pore domain, between adjacent S1-S4 domains and even intercalating between helices (Fig2a,b; teal molecules). Superposition of the structures of the paddle-chimera and the isolated S1-S4 domain of KvAP (Fig3a,b) reveals that the structures of voltage sensors are well conserved.
from archaebacteria to humans. In addition, two additional structural features can be seen in the comparison. First, both structures reveal a widespread network of electrostatic interactions between basic residues in S4 and acidic residues in S1, S2 and S3, and in many cases the side chains of these key residues virtually superimpose (Fig 3a,b). In the activated conformation of the paddle-chimera there are two clusters of acidic residues, an external acidic cluster formed by Glu residues in S1 and S2, and an internal acidic cluster consisting of a Glu in S2 and an Asp in S3a. The residues in the external cluster lie within a large crevice exposed to the external aqueous environment (Fig 3a; green arrow), while the internal cluster is largely buried within the protein interior. Based on their earlier mutagenic rescue experiments, Papazian and colleagues had concluded that the inner acidic residues in S2 and S3 form a stabilizing network with the fifth basic residue in S4 when the voltage sensor is in an activated state9, in agreement with the inner network shown in the structure of the paddle-chimera. A second structural feature evident in the comparison is that the outermost Arg residues do not interact with acidic residues, but project out from the domain towards the surrounding membrane (red arrow in Fig 2a,b and Fig 3a,b). Both of these structures were solved in the absence of an electric field, or 0 mV, a voltage where functional studies on KvAP and the paddle-chimera indicate that their voltage sensors are in activated conformations46, 57, implying that these two structures are representative of the activated state. The internal S6 gate region within the pore of the paddle-chimera is open, consistent with the voltage sensors being activated, and there are no structural changes in the selectivity filter to suggest that the channel is inactivated58. The paddle-chimera structure is compatible with constraints imposed by metal bridges between S4 and S5 in the activated state35, further suggesting that it is representative of the activated conformation.

Emerging concepts

The available structures of activated voltage sensors represent a remarkable advance and, when considered in the context of other biophysical studies, constrain our thinking about the mechanism of voltage sensing. While it is clear that structures of voltage sensors in other conformations are needed, such as the resting state that predominates at negative voltages, the available data allow us to frame several key concepts that are fundamental to the mechanism of voltage sensing. Some of these concepts have been central to our thinking for some time, while others have their roots in the more recent X-ray structures.

Charged residues have two important roles

The external four Arg residues in S4 appear to carry the majority of gating charge, thus serving to drive conformational changes in response to changes in membrane voltage18, 19, 21. Although this concept is also supported by extensive evidence that S4 actually moves22-37, an untested assumption in many of these studies is that neutralization of charged residues does not alter the position of the S4 helix. If this were to occur, the contribution of residues to gating charge could go undetected. The second important role of charged residues is to stabilize the basic Arg residues in the low dielectric of the membrane interior. Glu and Asp residues in S1-S3 form the external and internal clusters seen in the paddle-chimera structure and contribute to a network of electrostatic interactions that stabilize some of the S4 Arg and Lys residues within the membrane9, 40(Fig 3a,b). The outermost Arg residues do not appear to interact with acidic residues in the external cluster, but project towards the surrounding lipid membrane (see Fig 3a,b and below).

Voltage sensors contain a water-filled permeation pathway

The first hints that water might penetrate voltage sensors and that the charged Arg residues move within an aqueous pathway came from studies by Horn and others showing extensive reactivity of water soluble MTS reagents with Cys residues introduced within S422-25. In effect, the Arg residues in S4 can be viewed as tethered ions that move within S1-S4 domains.
Remarkably, untethered ions can actually permeate through voltage sensors. Bezanilla and colleagues found that His residues introduced within S4 create a proton pore\textsuperscript{27, 33}, and Isacoff’s group made the astonishing discovery that ions as large as guanidinium can permeate S1-S4 domains when the outermost Arg in S4 is truncated\textsuperscript{59, 60}. The recent discoveries of voltage-activated proton channels containing only an S1-S4 domain\textsuperscript{40-42} suggests that this permeation pathway can serve an important biological function. The paddle-chimera structure shows a large crevice between S1-S4 helices that would be contiguous with the external aqueous phase when the channel is embedded in a membrane. This region of the voltage sensor houses many of the basic and acidic residues (Fig 3a; green arrow), presumably identifying the pathway for ions to permeate voltage sensors\textsuperscript{61}. One important implication of water intercalation into voltage sensors is that it would be expected to focus the membrane electric field, allowing movements smaller than the full thickness of the bilayer to translocate charge across the full extent of the electric field. Indeed, focusing of the electric field has been inferred from studies with electrochromic fluorophores\textsuperscript{62}, proton permeation mediated by introduced His residues\textsuperscript{63} and charged MTS compounds with variable length tethers\textsuperscript{64}.

### The paddle motif flexes within S1-S4 domains

The S3b-S4 paddle motif contains the outer four S4 Arg residues that carry most of the gating charge in Kv channels\textsuperscript{18, 19, 21}, and in the X-ray structures this motif is connected to other regions of the protein via flexible linkers\textsuperscript{5, 34}. The accessibility of biotinylated positions in the paddle motif to avidin suggests that it is a uniquely mobile motif within S1-S4 domains\textsuperscript{34, 36, 65}, a feature that also can be seen in electron paramagnetic resonance (EPR) measurements of spin label mobility\textsuperscript{13, 66}. The paddle motif can be transplanted between proteins with S1-S4 domains, whether they are found in archaeobacterial or eukaryotic Kv channels\textsuperscript{44}, Nav channels\textsuperscript{67}, or voltage-sensing proteins like Ci-VSP or Hv\textsuperscript{144}, suggesting that this motif resides in a relatively unconstrained environment and that its function is conserved throughout voltage sensors. Another indicator that the paddle motif is a crucial flexing motif within S1-S4 voltage sensors is that it forms the receptor for tarantula toxins, such as hanatoxin\textsuperscript{44}, \textsuperscript{50-52}, which inhibit voltage sensor activation\textsuperscript{51, 54}. Many other toxins from spiders and scorpions alter the activity of voltage-activated ion channels by binding to paddle motifs\textsuperscript{67}, indicating that nature has targeted this motif throughout evolution.

### Voltage sensors interact intimately with lipids

The structures of KvAP and Kv1.2 channels discussed above suggest that the paddle motif is exposure to the surrounding lipids and the paddle-chimera structure provides a glimpse of what these interactions might look like because lipid molecules actually crystallize with the channel (Fig 2). EPR studies on the S1-S4 domain and full-length KvAP channel demonstrate extensive exposure of the voltage sensors to the surrounding lipid membrane\textsuperscript{13, 66, 68}. The interaction of tarantula toxins with paddle motifs also strongly support the idea that the paddle moves at the protein-lipid interface because these toxins bind to it within the membrane\textsuperscript{44, 53-55} and the toxin binding surface projects out towards the surrounding lipid membrane\textsuperscript{44}.

Although we are only beginning to learn about the nature of the interactions between lipids and voltage sensors, several recent studies suggest that these lipid-protein interactions may be fundamentally important for voltage sensor function. MacKinnon and colleagues showed that lipids with phosphate-containing headgroups are required for channel function and that they stabilize the voltage sensor in the activated state\textsuperscript{69}. One possibility is that the outer S4 Arg residues projecting out towards the surrounding membrane in the structure of the activated state of the paddle-chimera channel (red arrows in Figs 2a, b, 3a, b) actually interact directly with phosphate headgroups. The outer Arg residues exhibit exposure to the lipid membrane in EPR studies\textsuperscript{13, 66, 68} and stabilizing interactions between Arg residues and phosphate headgroups can be seen in molecular dynamic simulations\textsuperscript{70-72}.
A fascinating series of recent studies by Lu and coworkers points to intimate interactions between specific lipids and voltage sensors. They found that extracellular application of sphingomyelinase D, an enzyme that hydrolyzes the zwitterionic membrane phospholipid sphingomyelin to the anionic ceramide-1-phosphate, produces a dramatic stabilization of the activated state of the voltage sensors. The implication is that ionic interactions between positively charged S4 Arg residues and the negatively charged phosphate headgroup are strengthened when the positively charged choline group is removed. The effects of sphingomyelinase D can only be seen on certain types of Kv channels, suggesting that sphingomyelin may be interacting intimately with those channels that are sensitive to the lipase. In addition, sphingomyelinase D treatment dramatically alters the stability of hanatoxin bound to voltage-sensor paddles, consistent with an intimate interaction between toxins, lipids and paddle motifs. Treatment of membranes with sphingomyelinase C, which removes the entire headgroup of the lipid, immobilizes the voltage sensors in a resting state, consistent with the possibility that phosphate-containing lipids help to stabilize the voltage sensors in the activated state. Remarkably, binding of tarantula toxins to the voltage sensors appears to diminish access of sphingomyelinase C to the channel, suggesting that the interaction between sphingomyelin and voltage sensors are remarkably intimate. All of these studies raise the fascinating possibility that lipid interactions with voltage sensors may play crucial roles in the mechanism of voltage sensing.

Voltage sensor motions

To proceed from these concepts to a more refined understanding of the mechanisms of voltage sensing, we will need to see an actual structure of a voltage-sensing domain in the resting state. The rearrangements within the voltage sensors that occur as the protein moves from an activated state, of which we now know quite a bit about the structure, to the resting state, where we know relatively little, remains poorly constrained. If we are willing to use broad strokes, however, the emerging concepts can be put together to paint an impressionistic image of voltage sensor motions. It seems likely that the activated state of the voltage sensor is the stable conformation because this is the state found in the absence of a membrane electric field (0 mV). The domain is organized so that the S3b and S4 helices of the paddle motif reside in a relatively unconstrained and flexible environment, with much of the surface of the motif exposed to the surrounding lipid bilayer (Fig 2,3). This includes the outermost Arg residues that likely interact with phosphate lipid head groups (Fig 4). The deeper Arg residues are stabilized by interactions with acidic residues in S1, S2 and S3 (Fig 3,4), which line a water-filled crevice projecting into the protein from the external aqueous phase. In the case of Kv channels studied thus far, the S1-S4 domains have few interactions with the pore domain within the outer half of the bilayer that are required for voltage sensing. The existence of Ci-VSP and Hv1 seem to confirm this, in particular because both of these voltage sensing proteins can function as monomers. When membrane voltage becomes negative (inside relative to outside), the paddle motif moves the most, with charged Arg residues in S4 moving towards the internal side of the membrane (Fig 4-5). The S3b helix may move in association with S4, or the interface between the two helices may change somewhat, but in either case it seems likely that the S3b helix does not become deeply submerged in the membrane. The outer most Arg residues probably exchange their interactions with phosphate head groups for acidic residues in the external cluster in an ion-exchange type mechanism, the middle Arg residues exchange interactions with the external cluster for those in the internal cluster (Fig 4), and the inner most Arg residues might exchange interactions with the internal cluster for those with phosphate head groups of lipids within the inner leaflet of the bilayer. It's tempting to speculate that the prominent external facing crevice seen in the paddle-chimera structure (Fig 3a; green arrow) may largely vanish when the domain adopts a resting conformation and an inward facing water-filled crevice might emerge.
Range of motion

While this picture captures the key concepts for which there is experimental evidence, we haven’t said much about the nature of the motions within the voltage-sensing domains, and this is where things remain somewhat dicey. Viewed collectively, most of the evidence is compatible with the notion that the Arg residues in S4 move about 15 Å inward along a trajectory that is roughly perpendicular to the membrane plane as the voltage sensors move from activated to resting states. Tarantula toxins can partition into the outer leaflet of the membrane and bind to the paddle motif in both resting and activated states in eukaryotic Kv channels, a constraint suggesting that the motions of the paddle are confined to the outer half of the bilayer. Similarly, fluorescence resonance energy transfer between fluorophores attached near S4 and lipophilic acceptors that flip back and forth across the membrane suggest that motions of S4 are considerably less than the full thickness of the bilayer. One informative estimate of the allowed motions of S4 comes from experiments measuring the reactivity of biotinylated residues in the S1-S4 domain of KvAP with avidin applied to external or internal solutions. When the accessibility data are mapped onto the structure of the S1-S4 domain of KvAP, it is evident that the S4 helix must be capable of motions of at least 15 Å (Fig 5a). Because the structure represents the activated conformation of the voltage sensor, the positions towards the external side of the membrane that are accessible to internal avidin (Fig 5a; yellow and blue spheres) must move to within 10 Å of the internal side of the membrane in the resting state (in the vicinity of the blue spheres towards the internal side of the structure, which are marked by a dashed line). Such a motion is consistent with MTS accessibility experiments, His-mediated proton transfer studies and mutagenic rescue experiments. Another constraint for S4 motions comes from the recent discovery by Bezanilla and colleagues in the Shaker Kv channel that Cys residues introduced at the outermost Arg in S4 can form disulphide and metal bridges with Cys residues introduced into S1 or S2 when the voltage sensors are in the resting state. The distance between Cβ atoms in the X-ray structure of the paddle-chimera is 19 Å for the S4 (green sphere) to S1 (light pink sphere) bridge, and 18.4 Å for the S4 to S2 (magenta sphere) bridge. Although S4 might move somewhat more or less than the 18-19 Å separating these residues (depending on the orientation of the Cys side chains), these bridges would seem to indicate that S4 motions in Shaker Kv channels are somewhat less than those proposed for KvAP based on the biotin-avidin studies. In contrast to these estimates, most fluorescence distance measurements point to motions on the order of a few angstroms. A recent fluorescence study on the Shaker Kv channel reporting an extensive number of distance constraints supports larger displacements of S4, on the order of about 10 Å. The question of how far particular regions within S1-S4 move will require further study and it will be important to explore the possibility that the range of motions can vary for different types of voltage-activated ion channels.

Possible rotational motions

In addition to translational motions, the S4 helix has been proposed to rotate, perhaps as much as 180°. The bridges already discussed between S4 and either S1 or S2 offer an informative perspective on the question of S4 rotations. If we consider these bridges in the context of the X-ray structure of the activated state of the paddle-chimera, it seems that S4 wouldn’t need to rotate much for the relevant position (green sphere) to form a bridge with S1 (light pink sphere), but would need to rotate significantly (< 90°) to bridge with S2 (magenta sphere). These bridges seem to rule out large rotations of S4, but would be consistent with the possibility that S4 undergoes a modest rigid body rotation relative to S1 and S2. Another scenario put forward based on the X-ray structures of the paddle-chimera and the MloTiK channel is that S4 undergoes a transition from a standard α-helix with 3.6 residues per turn in the activated state to a more tightly wound 3_10 helix with 3 residues per turn in the resting state. Transitioning from a standard α-helix in the activated state to a 3_10 helix in the resting state would allow the outermost Arg residues to rotate moderately into the S1-S4 domain.
without necessitating the type of rigid body motion that would require extensive adjustments in packing between S4 and the other helices in the voltage sensor. In the structure of the activated paddle-chimera channel, the internal region of S4 is also a \( 3_{10} \) helix, leading to the suggestion of a wave-like transition where the zone of \( 3_{10} \) secondary structure propagates from the internal to external regions of S4 as the voltage sensor moves from activated to resting states\(^{46} \). \( 3_{10} \) helices are energetically unstable, but in the case of voltage sensors one can imagine that the energy necessary to adopt a \( 3_{10} \) helix could come from the membrane electric field. In this way, the electric field would hold the voltage sensor in an unstable or 'cocked' position at negative membrane voltages, which would readily relax to an energetically more stable conformation once the field dissipates with depolarization of the membrane towards 0 mV.

**Prospects for the future**

The present picture of voltage sensor structure and motions has evolved considerably since the cloning of the first membrane proteins with S1-S4 voltage sensors, yet many exciting and open questions remain. What does the structure of a voltage sensor look like in the resting conformation? How does the voltage sensor move during activation? Where exactly is the permeation pathway for ions to permeate S1-S4 domains? How do the interactions of lipids with voltage sensors influence their structure and motions? What does the structure of the lipid membrane look like around voltage sensors? The majority of structural and biophysical experiments have been done on a relatively small number of voltage-activated ion channels, mostly Shaker, Kv1.2 and KvAP, and at this point we don't know how similar the mechanisms will be for the large family of ion channels and enzymes that contain S1-S4 domains. Indeed, there is precedence from the work of Aldrich and colleagues that gating in some types of Kv channels involves only a subset of the conformational changes thought to occur in Shaker Kv channels\(^{86} \). In addition, very little is known about the S1-S4 domains for ion channels such as cyclic nucleotide gated channels or transient receptor potential channels. Do the S1-S4 domains in these channels have any voltage sensing functions? Do they undergo structural changes upon binding ligands that might be related to those that occur in voltage sensors? In many respects, we have only recently developed a solid foundation for thinking about how S1-S4 voltage-sensing domains work, and there are many fascinating experiments to be done.

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**References**

42. Tombola F, Ulbrich MH, Isacoff EY. The voltage-gated proton channel Hv1 has two pores, each controlled by one voltage sensor. Neuron 2008;58:546–56. [PubMed: 18498736]
50. Li-Smerin Y, Swartz KJ. Helical structure of the COOH terminus of S3 and its contribution to the gating modifier toxin receptor in voltage-gated ion channels. J Gen Physiol 2001;117:205–18. [PubMed: 11222625]
Figure 1. Types of membrane proteins that contain S1-S4 voltage-sensing domains and structure of an S1-S4 domain
a) Cartoon illustration of S1-S4 voltage-sensing domains in different types of membrane proteins. S1-S4 helices are labeled with the paddle motif colored purple. In voltage-activated ion channels the S1-S4 domains couple to an ion selective pore domain (yellow); only one of four S1-S4 domains is shown for clarity. In enzymes like the voltage-sensitive phosphates (Ci-VSP) the S1-S4 domain couples to a soluble phosphatase domain (yellow). In voltage-activated proton channels, protons are thought to permeate the S1-S4 domain directly. b) Ribbon representations of the X-ray structure of the S1-S4 domain of KvAP (PDB accession code 1ORS) with the paddle motif colored purple. The outer four Arg residues in S4 are shown as stick representations with carbon colored yellow and nitrogen colored blue. Structural representations in all figures were generated using PyMOL (DeLano Scientific).
Figure 2. Structure of the paddle-chimera Kv channel
a) Ribbon representation of the X-ray structure of the paddle-chimera channel viewed from the external side of the membrane. The paddle motif is colored purple, the pore domain is colored yellow, lipids are colored teal and basic residues in S4 are shown as stick representations with carbon atoms colored yellow, oxygen atoms colored red and nitrogen atoms colored blue. PDB accession code is 2R9R. Red arrow identifies outer S4 Arg residues projecting towards the lipid membrane. b) Side view of the paddle-chimera channel focusing on the S1-S4 voltage-sensing domain and its interface with the pore domain.
Figure 3. Charged amino acids in S1-S4 voltage-sensing domains
Superposition of the structures of the S1-S4 domains of KvAP (yellow ribbon) and the paddle-chimera channel (white ribbon) shown as a stereo pair and viewed from the side. The structures were aligned to minimize deviations between the S1 and S2 helices. Basic and acidic residues are shown as stick representations with carbon colored yellow or white, oxygen colored red and nitrogen colored blue. The green arrow identifies a aqueous crevice within the S1-S4 domain that would be continuous with the external solution and the red arrow identifies outer S4 Arg residues projecting towards the lipid membrane. PDB accession code for S1-S4 of KvAP is 1ORS and that for the paddle-chimera is 2R9R. b) Superimposed structures viewed from the external side of the membrane.
Figure 4. Changes in charge interactions during movements of voltage sensors

The cartoons illustrate shifting interactions between positively charged Arg residues in S4 and either negatively charged phosphate lipid headgroups (orange) or acidic residues (red) in the external and internal acidic residue clusters. Cartoon to the left is for the activated state that occurs at positive membrane voltages and that on the right is for the resting state that occurs at negative voltages. The illustrated motions of helices are not meant to imply anything about the structural changes occurring during voltage sensor movement.
Figure 5. Inferring motions from accessibility and bridging experiments

(a) Ribbon representation of the structure of the S1-S4 domain of KvAP showing accessibility of biotinylated positions to avidin with 9 and 10 Å tethers. Positions marked by red spheres (α-carbon) are accessible to only external avidin, those marked by blue spheres are accessible to only internal avidin, and those marked by yellow spheres are accessible to either external or internal avidin. Black spheres mark inaccessible positions. The structure is of an activated voltage sensor, so yellow and blue positions located towards the external side of the membrane must move to within ~10 Å of the internal side of the membrane (dashed line) in the resting state, as indicated by the black arrow. Basic and acidic residues are shown as stick representations.

(b) Ribbon representation of the structure of the S1-S4 domain of the paddle-chimera channel showing positions that form disulphide or metal bridges in the resting state. The structures shown in a and b are aligned as in Fig 3. In the Shaker Kv channel, Cys substituted at R362 in S4 (equivalent to R294 in the paddle-chimera; green sphere) can bridge with either I241C in S1 (I177 in the paddle-chimera; light pink sphere) or I287C in S2 (I230 in the paddle-chimera; magenta sphere) when the voltage sensor are in a resting state.

c) Bridging positions from b viewed from the external side of the membrane.