I. INTRODUCTION

The earth is 4.6 billion years old (Fig. 1A). Since it became a living planet some 3.8 billion years ago, microbes represent the largest group of living organisms on it (394). Microbes are much more diverse than animals and plants (393) and outweigh all other living organisms by far in terms of their biomass. They include prokaryotes (bacteria and archaea) and eukaryotes (protozoa, algae, and fungi), occupying numerous evolutionary branches
on the universal phylogenetic tree (Fig. 1B). Their diversity, number, and adaptability to environments characterized by extreme temperatures, salinity, or acidity (270) have enabled them to inhabit all the niches of life on the planet and will in all likelihood allow them to outlive any future global cataclysm.

Why study ion channels in microbes? Microbes are not only very interesting study objects in their own right, but they also offer specific experimental advantages for studies of structure and function of ion channels. Highlighted by the 2003 Nobel Prize in Chemistry “for structural and mechanistic studies of ion channels” in which bacterial ion channels played an essential role (229), microbes have in the past often been used to advance our understanding of basic biochemical or physiological cellular processes. They are uniquely suited for genetic manipulation and are the prime subject of recombinant DNA technology. They have very short replication times and some are simple to grow, allowing for large yields in laboratory cultures, which is a prerequisite for a large-scale production of milligram amounts of ion channel proteins required for structural studies by X-ray crystallography or magnetic resonance spectroscopy. Microbes also provide a unique opportunity to study evolutionary origins and primordial design of ion channels. Channel prototypes may be deduced from the commonalities of channels of extant samples from all three domains.

A. A Brief History

Beginning with Galvani’s discovery, bioelectricity is largely the concern of animal physiology, with the excitation of nerve and muscle being the centerpiece. This animalcentric view is further accentuated in recent years because of human health concerns. Ion channels, the molecular bases of excitation and beyond, are therefore continued to be studied mainly in the human, mouse, and fly. Two developments broadened our horizon, revealing the existence of ion channels in all three domains of life (Fig. 1). First, patch-clamp survey of the model bacterium *Escherichia coli* (238; see sect. II A1) and the model unicellular eukaryote *Saccharomyces cerevisiae* (114, 115; see sect. IV C1) revealed unitary ion conductances on their membranes. These findings not only showed that ion channels exist in microbes but also showed that they are not necessarily associated with motility, an impression previously reinforced by the study of *Paramecium* behavior (see sect. IV A). Second, the ever-increasing number of microbial genome sequences revealed gene products akin to animal channels. Because of the relative ease in microbial culture, milligrams of bacterial and archaeal channel proteins can be purified, yielding diffractable crystals (80; see sect. III). The results are microbial ion-channel structures of atomic resolutions, leading to insights that revolutionized our understanding of ion channels.

Structural and functional studies of ion channels in prokaryotic cells are some of the best examples of how microbes as model systems have contributed towards our understanding of general principles underlying the structure and function relationship in biological molecules. A multidisciplinary approach that can be employed in studies of bacterial mechanosensitive channels (13, 22, 27, 44, 202, 238, 300–302, 366, 367, 401), K⁺ channels (11, 80, 298, 299, 340), Na⁺ channels (294, 321), Cl⁻ channels (83), cNMP-gated channels (53), and glutamate receptor channels (47) as well as archaeal mechanosensitive channels.
and Ca\textsuperscript{2+}-gated K\textsuperscript{+} channel (206) enhanced our understanding of the function of ion channels in living cells in ways that could not have been done by studying animal channels only.

Not only for historical reasons but also for keeping an accurate record of achievements of the pioneers in electrophysiology, it is worth mentioning here that the first in vivo cellular electrical recordings from a living organism were obtained from a microbial cell. It was a protozoan *Paramecium* that in 1934 was impaled by a glass microelectrode to register the resting membrane potential of this microbe (150). This work was later overshadowed by the great work of Alan L. Hodgkin and Andrew F. Huxley on neuronal excitation and conduction in squid giant axon (134, 139).

### B. Scope of This Review

The focus of this review is on the structural and functional characterization of ion channels in prokaryotic (bacteria and archaea) and eukaryotic microbes (algae, ciliates, and fungi). Some of these microorganisms have been excellent experimental objects amenable to modern electrophysiological techniques. Probing their cellular membranes by glass microelectrodes connected to amplifiers, oscilloscopes, and computers has made a great contribution towards our understanding of the workings, functional role, and evolutionary origins of ion channels in living cells. Conceptually, this review consists of two major parts: the first one focusing on ion channels in prokaryotic cells and the second one describing ion channels in eukaryotic microbes. For completeness, we also included in the review two sections that are briefly touching upon the channels formed by microbial toxins and ion channels found in viruses.\(^1\) Space concern does not allow us to cover other channels such as porins, aquaporins, organelle channels such as VDAC, nucleapore, etc. Comprehensive reviews on these subjects are available elsewhere.

### C. Voltage-Clamp and Patch-Clamp Investigation of Microbial Cells

With the exception of some large protozoa and algae, the classical electrophysiological techniques of the whole cell current clamp and voltage clamp using two glass microelectrodes were for a long time not applicable to microorganisms because of their very small size. The patch-clamp method (118) provided means to overcome this shortcoming. Prior to the advent of the patch-clamp technique, electrophysiological measurements in microbial cells could only be carried out in large protozoan or algal cells in vivo or in vitro by reconstituting ion channels into planar lipid bilayers (87, 263, 395). The patch-clamp technique allowed, for the first time, the study of electrical properties of very small cells by examining ionic currents flowing through individual ion channels in their cellular membranes in situ. Whereas little modification of recording solutions, glass pipettes, and electrophysiological equipment is required to record ionic currents from microbial cells, appropriate preparations of microbial cell membranes amenable for giga-seal formation and patch-clamp recording present a major challenge. The following examples may illustrate the variety and architectural complexity of microbial cell envelopes.

A single *Paramecium* cell is covered by $\sim$5,000 cilia, and the cell membrane is covered by a thick extracellular matrix consisting mainly of the “immobilization antigen” protein (228). In yeast (single-celled fungi), the cell membrane is covered by a cell wall made of glucans, mannans, proteins, and lipids (269). The cytoplasmic membrane of Gram-positive bacteria is enclosed by a thick cell wall made of peptidoglycan that provides the rigidity necessary to maintain cell integrity, whereas Gram-negative bacteria have in addition a second chemically distinct outer membrane attached to the peptidoglycan layer on its external side (385). The situation with archaea is further complicated by the large diversity of extreme habitats to which archaea have adapted, which greatly influenced the chemical composition and architecture of their cell membranes (162). Not only the cell wall structure, which varies considerably among archaeal species, but also the composition of cell membranes of archaea is very different from bacterial membranes. Archaeal membranes are not made of phospholipids, but consist of diphytanyl-glycerol-dieither or -tetraether or both (79). Consequently, there is no single method that can be applied to all microbes to expose their membranes for the purpose of patch-clamping. Each type of microbial cell has to be approached individually. For example, the examination of a bacterial cell membrane by the patch-clamp technique became possible some 20 years ago with development of a “giant spheroplast” preparation in *E. coli*, which has led to a discovery of MS channels in these microbes (238, 328) (Fig. 2). Briefly, a culture of *E. coli* cells is grown in media containing antibiotic cephalxin until strings of bacteria, called “snakes,” of the appropriate length are formed as observed under the microscope. Giant spheroplasts are then obtained by addition of EDTA and lysozyme to perforate the outer membrane and break glycosidic linkages in the cell wall structure causing the “snakes” to round up into spheroplasts of 5–10 $\mu$m in diameter. This giant spheroplast preparation has recently been optimized for studies of MmaK, a putative cyclic nucleotide-gated K\textsuperscript{+} channel cloned from *Magnetospirillum*.

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\(^1\)Although viruses are not microbes, but encapsulated parasitic fragments of DNA or RNA, we have also included in this review a brief account of viral ion channels, because of their minimalistic structure matching the simplicity of microbial channel-forming toxins and of some of the extant channels.
lum magnetotacticum (194). A diagram summarizing several of the methods used for patch-clamp recording from microbial cells is shown in Figure 2. Details of these methods (334) are described in references cited under individual organisms detailed below.

II. LARGE-CONDUCTANCE MECHANOSENSITIVE CHANNELS OF PROKARYOTES

A. Mechanosensitive Channels of Bacteria: MscL, MscS, and MscK

Although they conduct ions and are usually characterized by their ionic conductance and selectivity, prokaryotic MS ion channels are often referred to as MS channels rather than MS ion channels. This is because in bacteria (and possibly also in Archaea) they primarily serve to transport cellular osmoprotectants other than ions by acting as osmosensors that regulate the cellular turgor (see sect. II C). Bacterial MscL and MscS channels have large conductances of ~3 and ~1 nS, respectively, and lack ionic specificity by passing anything smaller than ~1,000 molecular weight including proline, potassium glutamate, trehalose, and ATP (117). MscL is nonselective for both anions and cations (62, 368), whereas MscS exhibits a slight preference for anions over cations with a permeability ratio $P_{Cl}:P_{K} = 1.5-3.0:1$ (238, 354, 364, 368).

1. MscL (mechanosensitive channel of large conductance)

The technical advance of the “giant spheroplast” preparation in E. coli (238, 328) opened a window of
opportunities for structure and function studies of mechanosensitive (MS) class of membrane proteins in prokaryotes. Prokaryotic MS channel currents have most extensively been studied in *E. coli* (236, 238, 239, 297, 362, 416), which harbors several MS channels in its cell envelope: MscL (large), MscS/MscK (small/kalium, i.e., potassium), and MscM (mini), named for their single-channel conductances of ~3, ~1, and ~0.3 nS, respectively (19). Given that bacteria can be grown in large quantities delivering the milligram amounts of channel proteins required for structural studies, three-dimensional structures of MscL from *Mycobacterium tuberculosis* (44) (Fig. 3A) and MscS from *Escherichia coli* (13) (Fig. 3C) were solved by X-ray crystallography only a few years after identification and cloning of both channels in *E. coli* (202, 366). MscK was cloned simultaneously with MscS (202), whereas MscM (19) still remains to be identified at the molecular level.

The strategy used for cloning of MscL consisted of detergent solubilization and fractionation of *E. coli* membrane constituents by column chromatography combined with the patch-clamp examination of the individual protein fractions reconstituted into artificial liposomes (366, 367). Given that mechanosensitivity of MscL is fully preserved upon the reconstitution of the channel protein into liposomes (367, 368), the protein fraction containing MscL activity could be identified, which enabled the isolation of the MscL protein comprising 136 amino acid residues followed by the cloning of its corresponding *mscl* gene (366). The expression of the gene in trans as well as with an in vitro transcription/translation system confirmed that the *mscl* gene alone was required and sufficient for the MscL activity.

Three-dimensional structure of the MscL homolog from *M. tuberculosis* obtained at 3.5 Å resolution shows a homopentameric channel, most likely in a closed state (Fig. 3A) (44). The channel monomer consists of two α-helical transmembrane (TM) domains, TM1 and TM2, cytoplasmic NH₂- and COOH-terminal domains and a central periplasmic domain. The channel pore is formed by five transmembrane TM1 helices organized as a tightly packed bundle. At the cytoplasmic side, the five helices funnel into a hydrophobic constriction of 2 Å functioning as the channel gate. The diameter of the channel pore at the gate expands from 2 to 30 Å during the channel opening as determined by electrophysiological perme-
oration studies using large organic cations (62) and electron paramagnetic resonance (EPR) spectroscopy combined with cysteine scanning mutagenesis and site-directed spin labeling (SDSL) (300–302). FRET spectroscopic analysis confirmed these findings by showing that the overall change in diameter of the MscL pentamer between its closed and open state is 16 Å (59). This result is consistent with the very large conformational changes that MscL undergoes during gating, which are necessary to accommodate the opening of the large pore of the channel. Mutational studies that led to isolation of either gain-of-function (GOF) or loss-of-function (LOF) mutant strains helped further in identifying functionally important residues and structural regions in MscL (208, 242, 283, 400, 401).

Since the publication of a three-dimensional structure of MscL (44), a large number of computer simulation studies aimed at understanding the gating mechanism of the channel by changes in membrane tension or composition. Simulations examining the stability of the protein structure when embedded in the lipid bilayer membrane indicated large conformational changes occurring when the MscL protein is influenced by an external force, in agreement with the results of the experimental studies. The simulations that have been carried out to investigate the nature and sequence of these conformational changes differ mostly in the way that the external force is applied to the MscL protein and conditions that affect the interactions between the lipid and the protein (25, 54, 90, 112, 113, 170) (for review on computational studies of bacterial MS channels, see also Ref. 58).

The TM1 and TM2 helices of MscL are connected by a structurally not well defined periplasmic loop, which is thought to function as a spring, resisting the channel opening (2, 290). This view has been supported by a molecular dynamics study of MscL embedded in a curved bilayer (246). According to this study, the periplasmic loop is the first domain of MscL to change its structure leading to channel opening. At the cytoplasmic end of MscL, the short NH2-terminal domain comprising approximately a dozen amino acids has been proposed to function as a second channel gate working in accord with the hydrophobic gate formed by the TM1 helix bundle (22, 365). Although results from a recent study, which utilized cysteine trapping of single-cysteine mutated channels, do not support the “second-gate” concept (144), the NH2 terminus is nevertheless an important structural domain of MscL, as demonstrated by amino acid deletion and substitution studies showing that structural changes in the NH2 terminus severely affect MscL function (30, 122). The precise role of the NH2 terminus for the MscL gating thus remains to be determined. The COOH-terminal domain, which forms an α-helical bundle at the cytoplasmic end of the channel (44) (Fig. 3A), appears to be the least important structural domain for the channel gating, since a deletion of 27 COOH-terminal residues constituting a large portion of the COOH terminus was shown not to significantly affect mechanosensitivity of the channel (22, 29, 365). Although the precise physiological role of the COOH-terminal domain is unclear at present, a current view of its role is that of a size-exclusion filter, which should prevent a loss of essential metabolites at the cytoplasmic side of the MscL channel pore (5). Recently, it has been shown that the structural stability of the COOH-terminal bundle is pH dependent (160), suggesting that in addition to its role of a size-exclusion filter, COOH terminus may also influence the channel gating in a pH-dependent manner. Furthermore, it has recently been shown that the COOH terminus may actively participate in the channel gating, since the opening of MscL is accompanied by the dissociation of the COOH-terminal helix bundle and pore formation (402).

2. MscS and MscK (mechanosensitive channel of smaller conductances)

MscS and MscK were cloned several years after MscL (202). Two genes, yggB and kefA, corresponding to MscS and MscK respectively, were identified on the E. coli chromosome. Deletion of these genes led to the abolishment of the MS currents corresponding to the MS channel of small conductance, leaving the activity of MscL intact in E. coli giant spheroplasts. Originally, MscS and MscK appeared indistinguishable in the patch-clamp experiments (238). However, MscS was later shown to differ functionally from MscK by exhibiting a rapid inactivation upon sustained application of pressure (173). A distinguishing property of MscK on the other hand is its sensitivity to the extracellular ionic environment (207). Structurally, MscS is a small membrane protein of 286 amino acids, whereas MscK is a much larger, multidomain membrane protein comprised of 1,120 amino acid residues. Both proteins share a common structural feature such that the primary amino acid sequence of MscS closely resembles the sequence of the last two domains of the MscK protein (202).

A three-dimensional crystal structure of MscS of E. coli obtained at 3.9 Å resolution (13) shows a channel folded as a homoheptamer (Fig. 3C). Each of the seven subunits is composed of three transmembrane domains with NH2 termini facing the periplasm and COOH termini facing the cytoplasm. Initially, the three-dimensional structure of MscS was thought to depict an open channel (13). However, recent computer simulation studies have cast doubt on this assertion (58). The TM1 and TM2 transmembrane domains, which are in direct contact with the surrounding lipids, are thought to constitute the sensor for membrane tension as well as a voltage sensor because of the presence of three arginine residues in their structure (13, 23). Since the precise contribution of these charged residues to the channel voltage dependence is
not known, their role in the channel gating remains to be established experimentally. TM3 helices line the channel pore. As a distinguishing feature, the channel has a large, cytoplasmic domain (Fig. 3C), which is in contact with the cytoplasm through multiple openings. Its major characteristic is a large interior chamber of 40 Å in diameter. The cytoplasmic domain is a dynamic structure that undergoes significant conformational changes upon the channel gating (172). It passively follows the movement of the TM1-TM2 “hairpin” and the membrane portion of the TM3 helix by expanding during the channel gating. In addition to its role in the channel gating, this domain has been suggested to also play a role in the channel desensitization and stability (248, 342) as well as in ion transport and selectivity (86).

Most of the computational studies that have appeared since the publication of the crystal structure of the MscS channel (13) have attempted to answer the question on the conduction state of the crystallized structure. The crystal structure of the protein showing a large nonoccluded pore passing across the membrane has suggested that the protein had most likely been imaged in an open state. Closer inspection of the structure, however, indicates that the narrowest region of the pore has a diameter of only ~3.5 Å and is surrounded by two rings of nonpolar leucine residues (L105 and L109). Such narrow hydrophobic pores can, however, prevent the passage of ions without presenting a physical occlusion as indicated by a number of molecular dynamic investigations of model pores (17, 18) and the nicotinic acetylcholine receptor (16, 57), which have shown that water tends to evacuate the hydrophobic regions of such pores under certain conditions. If the radius of a pore is increased to 4.0 – 4.5 Å or a more polar surface is introduced, then water will fully hydrate the channel. As the radius of the constriction in the MscS pore is below the critical radius for hydration seen for other pores, it is likely that a similar hydrophobic gating mechanism could play a role in the MscS gating. Consequently, the MscS crystal structure could represent a closed or nonconductive state rather than an open conformation as originally proposed. Molecular dynamics studies of the MscS pore indicated that water does indeed evacuate the constricted region of the pore and provided some of the first evidence for a hydrophobic gating mechanism in a real channel rather than model (6, 353).

3. Force transmission from the lipid bilayer

Both channels, MscL and MscS, directly sense membrane tension developed in the lipid bilayer alone (123, 234, 237, 300, 301, 364, 383) (Fig. 3, B and D). The bilayer mechanism, as this mechanism of the MS channel gating has been named (119), has been well documented for MS channels of prokaryotes (236, 303), since the prokaryotic MS channels preserve their mechanosensitivity after reconstitution into artificial liposomes. The fact that MscL and MscS can be activated by amphipaths, known to insert preferentially in one leaflet of the lipid bilayer (237, 346), provided a basis for a recent spectroscopic and patch-clamp studies. In these studies, structural changes in MscL induced by either hydrophobic mismatch or curving the bilayer by insertion of the amphipath lysophosphatidylcholine (LPC) were examined by combining cysteine-scanning mutagenesis with SDSL, EPR spectroscopy, and patch-clamp functional analysis of MscL reconstituted into liposomes (300, 301). The studies demonstrated that hydrophobic surface matching could stabilize intermediate conformations of MscL, requiring less tension to open the channel in thin bilayers (<18 hydrocarbons/acyl chain) compared with thick bilayers (>18 hydrocarbons/acyl chain), but was insufficient to fully open the channel. However, curving the bilayer by asymmetric insertion of LPC opened MscL without applying membrane tension, indicating that the mechanism of mechanotransduction in MS channels is defined by both local and global asymmetries in the transbilayer tension profile at the lipid protein interface. A recent study indicated the asparagine substitution of the residues near both ends of the transmembrane helices impaired the function of MscS (275). Similarly, it has been shown that introduction of Tyr, Trp, Asn, or Asp residues at the periplasmic end of the transmembrane domains impairs MscL mechanosensitivity (401). Thus MscS shares the same characteristics with MscL in that the interaction with lipids near the polar-apolar boundary is essential to the channel function (36). These findings have general implications for MS channel gating, since they seem to suggest the role of the lipid bilayer in actively modulating the activity of MS ion channels in all types of living cells (184).

4. MscL and MscS homologs

The accessibility of a large number of genome sequences of different evolutionary groups has made possible the analysis of the phylogenetic distribution of MscL and MscS type of channels in these microorganisms. Multiple sequence alignments of homologs of MscL and MscS revealed that they form separate families of prokaryotic MS channels (240, 307). Furthermore, a recent comprehensive analysis of nucleotide composition and codon usage from six phylogenetic groups of prokaryotic MS channel genes indicates that the evolution of the prokaryotic MscL and MscS genes followed separate evolutionary pathways (49). A subfamily group of MscL-type proteins encompasses MS channels of gram-negative and gram-positive bacteria. It also includes MscL-like proteins of a single archaeon, Methanosarcina acetivorans, and a fungus, Neurospora crassa (163, 181, 307). The archaeal and fungal proteins are the most divergent members of the MscL family, which most closely resemble those of gram-
positive bacteria (307). Compared with the MscL subfamily, the MscS family varies much more in size by including a number of representatives from bacteria, archaea, fission yeast *Schizosaccharomyces pombe*, single-celled alga *Chlamydomonas*, and plant *Arabidopsis thaliana* (266, 307). Although the MscS subfamily is more diverse, it is not ubiquitous. Several organisms with fully sequenced genomes including gram-negative chlamydias, the gram-positive clostridia, mycoplasmas, and ureaplasmas, do not encode recognizable MscS homologs (307). Furthermore, neither of the two subfamilies of MS channels include representatives from animal cells (163, 240, 307).

B. Mechanosensitive Channels of Archaea: MscMJ, MscMJLR, and MscTA

MS channel activities have also been reported in several species of Archaea (Fig. 1B) including halophilic *Haloferax volcanii*, thermophilic *Thermoplasma acidophilum*, and methanogenic *Methanococcus jannashii* (162, 240). Like MscL and MscS, archaeal MS channels are gated by tension in the lipid bilayer and are activated by amphipaths. They have large conductance and low selectivity for ions and are weakly voltage dependent. They can also be blocked by submillimolar concentrations of gadolinium, which is a common inhibitor of MS type of ion channels.

MscMJ and MscMJLR are the two types of MS channels found in *M. jannashii* (164, 165). These two channels differ functionally from each other despite sharing 44% sequence identity (240). Although they have similar ion selectivity with preference for cations over anions ($P_K/P_{Cl} \sim 5.0$), the conductance of MscMJ of 0.3 nS is much smaller compared with 2.0 nS conductance of MscMJLR (165). Both channels were identified during inspection of the *M. jannashii* genome database using the DNA corresponding to the TM1 helix of MscL as a probe. MscMJ comprises 350 residues, whereas MscMJLR is a slightly bigger protein comprising 361 residues (165). Both channels are members of the MscS subfamily of MS channels, with both proteins exhibiting ~28% sequence identity with overlapping residues of the MscS sequence (163, 165) (Fig. 4).

MscTA is a 15-kDa membrane protein underlying the activity of the MS channel of *Thermoplasma volcanium* (161). It was identified using a functional approach similar to the one used for molecular identification of MscL (366, 367). Like MscL, the channel has a large conductance of 1.5 nS and is nonselective for ions. MscTA seems also structurally related to MscL, since secondary structure analysis revealed two putative α-helical membrane-spanning regions despite that its primary amino acid sequence contains an unusually high proportion of charged residues for a membrane protein.

C. Physiological Role of MS Channels in Prokaryotes

Multiple adaptation mechanisms enable bacteria to grow in a wide range of external osmolarities (351, 396). Located in the cytoplasmic membrane (20, 124, 202, 276), MS channels have evolved specifically to serve as emergency valves for fast release of osmolytes in osmotically challenged bacterial cells. The opening of MS channels upon a hyposmotic shock enables rapid loss of cytoplasmic solutes and excess water, which helps to restore a normal cellular turgor and prevent cell lysis. As unambiguously demonstrated for MscL and MscS, these MS channels thus function as “emergency valves” for rapid and nonspecific release of solutes (28, 37, 202). As sensors and regulators of the cellular turgor, they provide a safeguard without which the bacterial cells would lyse. Double knock-out mutants of *E. coli* lacking both MscL and MscS die upon transfer from a medium of high to a medium of...
low osmolarity (38, 202). MscM, the third bacterial MS channel whose molecular identity remains unknown to date, is insufficient alone to protect the cells from lysis. Further evidence for the MS channel “safety valve” function has been provided by a study on marine bacterium *Vibrio alginolyticus*, in which introduction of an *mscL* gene was found to alleviate cell lysis by hyposmotic shock (265). Except in the extreme conditions of a hyposmotic shock, knockout bacterial cells lacking MscS, MscL, or both are otherwise fully viable. The multiplicity of MS channels is likely required to provide a safeguard against the deleterious effects during sudden changes in external osmolarity occurring in the living habitats in which prokaryotes exist.

Given that cell turgor is essential for growth and cell wall synthesis, prokaryotic MS channels may also be designed to detect changes in turgor pressure during cell division and cell growth, which are accompanied by an increase in cell volume required for the synthesis and assembly of cell wall components (63). Indeed, it has been demonstrated that the expression of MscS and MscL is stimulated in cells entering into stationary growth phase when they undergo cell wall remodeling and need to relieve the turgor pressure (361). Although the physiological role of MS channels in Archaea has not clearly been established, the archaeal MS channels, however, could be expected to have functions similar to those of their bacterial counterparts.

### III. ION-SPECIFIC CHANNELS OF PROKARYOTES

Unlike the study of mechanosensitive channels above, studies of prokaryotic ion-specific channels are not systematic. Historically, they are largely the extensions of crystallographic studies and not originated with microbial-physiological concerns. The recent decade saw the three-dimensional structures at atomic resolutions of prokaryotic K⁺ channels, Na⁺ channels, glutamate receptors, etc. These structures provided unprecedented insights into the molecular mechanisms of ion channels and revolutionized the field of ion-channel research. While these high-resolution structures now serve as the foundation of our ion-channel understanding and its biomedical extensions, these channels have not been studied extensively or systematically in the bacteria and archaea themselves. The roles they play in the various prokaryotic species are largely unknown. The narrative below will therefore concentrate on the knowledge gained through these structures. The deepest knowledge came from the studies of K⁺ channels. We will therefore begin with a simpler K⁺ channel, KcsA, from the gram-positive bacterium *Streptomyces lividans*. We will then build on the KcsA core other gating modules to describe Ca²⁺-, cNMP-, or voltage-gated K⁺ channel. We will then discuss glutamate receptor, a type of ligand-gated K⁺ channel in bacteria. This will then be followed with prokaryotic Na⁺ channel, akin to K⁺ channels, before describing mechanistically different types of channels including NH₄⁺ channels and Cl⁻ “channels.” In this narrative, examples from both bacterial and archaeal channels will be used, largely depending on the availability of crystal structures. Animal ion channel pores have associated partners that play auxiliary or regulatory roles. Often, the channel proper is called the α or the pore-forming subunit, and the other are called β₁, β₂, γ, etc. Some homologs of the non-pore-forming subunits are found in microbes, but are not reviewed here because of space consideration.

#### A. K⁺ Channels

K⁺ is the major cation in any cytoplasm. Correspondingly, K⁺ channel genes are found in nearly all genomes. These include those of bacteria and archaea, except the greatly reduced genomes of parasites and organelles (190, 192). Because of the crystal structures of prokaryotic channels, far more is known about prokaryotic K⁺ channels than most eukaryotic channels or other microbial channels, except MscL and MscS (above). This knowledge has tremendous impact on ion-channel biology and is extensively reviewed and discussed in the literature of animal physiology. For current discussions on K⁺-channel structures and functions in general, see References 107 and 278. For more on microbial K⁺ channels, see Kuo and co-workers (190, 192).

The study of ion channels is deeply rooted in neurophysiology. Excitation of nerves or muscles entails the sequential rise and fall of Na⁺ (Ca²⁺) and K⁺ permeability of the membrane. Permeation to a specific ion implies selectivity; rise and fall imply portal control. These functional characteristics have their conceptual structural counterparts: the “filter” and the “gate,” formulated long before we have clear structural information. These two key structures became “crystal clear” with the arrival of prokaryotic channels and are described below.

The puzzling feature of the ion filter is its very key properties: permeation and selectivity. A K⁺ channel can pass some 10⁷ K⁺ per second and discriminate against the smaller Na⁺ at the same time (Pₖ/Pₙ > 1,000:1). These properties challenge our common intuition about binding energy and binding specificity. The structure of the bacterial K⁺ channel, KcsA, readily explains these properties. The puzzling feature of the gate is how it can be controlled, and, in different channels, apparently controlled by all the known stimuli including ligands, second messengers, voltage, heat, and mechanical force. To more clearly summarize our current knowledge of cation channels, this review will first give a global view of the KcsA
channel and then proceed with the description of its filter, its gate, and its lipid environment.

1. KcsA at a glance

Most cation-specific channels, including K\textsuperscript{+}, Na\textsuperscript{+}, Ca\textsuperscript{2+}, and the cation-nonspecific TRP channels, have a similar structural motif. Each channel has four subunits, converging to enclose the ion pathway. The first K\textsuperscript{+} channel with high-resolution crystal structure is KcsA of Streptomyces lividans (80). The primary structure predicts each subunit of the tetramer to have, at its core, a TM1-P-TM2 arrangement, comprising two transmembrane helices (TM1, TM2), flanking a short "pore helix" and the filter sequence (together referred to as P) (Fig. 5a). Note that the well-known crystal structure of KcsA was that of its core (80) but not the entire KcsA peptide. Even in this simple channel, there is an additional helix upstream of TM1 and one downstream from TM2 (60). The simplest K\textsuperscript{+} channels with only TM1-P-TM2 are found in certain viruses. As shown below and in Figure 5, a common motif of K\textsuperscript{+} channel subunits is TM1-TM2-TM3-TM4-TM5-P-TM6, where TM5-P-TM6 is structurally and functionally homologous to the TM1-P-TM2 of KcsA. Genes predicting TM1-TM2-TM3-P-TM4 are found in prokaryotes (190). Two pore-domain K\textsuperscript{+} channels each with two TM1-P1-TM2-TM3-P2-TM4 subunits are found in animals (292, 293); those with two TM1- TM2-TM3-TM4-TM5-P1-TM6-TM7-P2-TM8 are found in fungi. Genes predicting K\textsuperscript{+} channel subunits with 12 TMs and 2Ps are also found in ciliates (see below) (187).

The KcsA tetramer is shaped like an inverted teepee (Fig. 6, a–c). The periplasmic side has a diameter of ~40 Å, the cytoplasmic side ~25 Å. The portion embedded in the lipid is ~35 Å long. TM1 transmembrane domains are the outer helices facing the lipids. TM2 domains are the inner helices, the COOH-terminal ends of which form the gate at the cytoplasmic side. Between TM1 and TM2 is the short "pore helix" cradling the ion-selective filter located towards the periplasmic side of the channel. The K\textalpha protein has two belts of aromatic amino acids, positioned to extend into the lipid bilayer near the lipid-water interface (Fig. 6e). The one towards the outside includes a cuff that surrounds the selectivity filter (80).

\textbf{A) THE K\textsuperscript{+} FILTER OF KCSA.} To restate, the channel resembles an inverted teepee, with the gate closing towards the cytoplasmic end, and the filter located at the other end. The K\textsuperscript{+}-filter sequence TXGY(F)GD is the most evolutionarily conserved sequence in the protein and is used as the diagnostic feature for K\textsuperscript{+} channels from all sources. In the tetramer, the four successive layers of carbonyl oxygens from the residues in the signature GYG sequence (residue 77–79 of KcsA) face inward, lining the filter (Fig. 6D). In KcsA, these oxygens are held in positions to provide the coordinates for the naked K\textsuperscript{+} in transit (80, 414). Five sites of K\textsuperscript{+} coordinations can be recognized, site 0 to site 4, counting from the outside. Since this coordination is essentially the same as that by water, these oxygens act as surrogate waters allowing the passage of dehydrated K\textsuperscript{+} through the filter to be nearly energetically costless, much like K\textsuperscript{+} diffusing through water, exchanging its hydration shell. The spatial geome-

\begin{figure}[h]
\centering
\includegraphics[width=0.8\textwidth]{figure5}
\caption{Topologies of K\textsuperscript{+} channel subunits. Each channel is a tetramer in forms of a, b, or c, or a dimer of d. a: A simple form of the subunit comprises two α-helices (TM1 and TM2, left and right rods) that traverse the membrane (marked with broken lines, separating the outside above and the cytoplasm below). Between the two are the pore helix (short rod) and the filter sequence (line between the pore helix and TM2). Certain viral K\textsuperscript{+} channels, KcsA, Kir, MthK, TvoK, have this topology, though helices preceding TM1 or trailing TM2 exist (not shown) in some of these channels forming structures in the cytoplasm. b: A common form of subunit comprising 6 transmembrane helices, in which the last two helices and the pore helix (TM5-TM6, right portion) form the permeation and filtration core, similar to the structure in a. The preceding four helices (TM1–TM4, left portion) form a separate domain, housing other structures, such as the voltage sensor. This 6-TM motif is found in KvAP, MVP, KvLqn, Kch, MloK, and MmaK. The subunit of the bacterial Na\textsuperscript{+} channels, NaChBac, also has this structure, although it has a slightly altered filter sequence. c: The subunit structure of GluR0, a prokaryotic glutamate receptor. The pore helix and the filter sequence are arranged in the direction opposite that of a. The portion outside the membrane forms the glutamate-binding site. d: The subunit structure of TO1K1, the K\textsuperscript{+} channel of budding yeast. Here the first four TM helices are followed by two core structures, each similar to that in a.}
\end{figure}
try of these coordinates does not fit other cations such as Na\(^{+}\) or Ca\(^{2+}\), let alone anions, the passage of which is therefore energetically costly. Thus the crystal structure of this K\(^{+}\) filter neatly solves the age-old puzzle of how the filter can be specific and efficient at the same time. In the three-dimensional structure, the ion filter is located near the outer surface of the membrane and is held in place by bonds between the amino acid residues in the filter sequence and those in the pore helices, including a network of aromatic-aromatic bonding (80). The conceptual separation of specific permeability to K\(^{+}\) and Na\(^{+}\) that form the foundation of membrane excitability and therefore neurophysiology has been formulated for more than half a century. It is ironic that the puzzling basis of ion specificity is finally solved in concrete terms with a channel from a “lowly” bacterium, and its true function in the life of the bacterium is unknown.

Compared with the gate that makes major movement (below), the filter is often viewed as a fixture. However, the filter is far from static. Its conformations are determined by the number and nature of the occupying ions (77) as well as the bonds between its constituent amino acid residues and those of the surrounding pore helices (56, 278). For example, the activation of KcsA is naturally followed by an inactivation (56), akin to the so-called “C inactivation” of other K\(^{+}\) channels. (Inactivation means the loss of channel current while the conditions of activation are maintained.) This inactivation occurs when the gate remains open (298) and explains why the observed open probability of KcsA is so low. Apparently, in the

**FIG. 6.** The crystal structure of KcsA. KcsA is a homotetramer of subunits each with a TM1-P-TM2 core as diagrammed in Fig. 5a. a: Viewed from the outside, with each subunit represented in one color and a K\(^{+}\) at the center. b: Side view of the membrane-embedded portion of the channel in form of an inverted teepee. Each subunit comprises an outer transmembrane helix (TM1), a short pore helix (P), and an inner transmembrane helix (TM2). Between TM1 and P are residues facing the outside (up), forming a turret. Between P and TM2 is the filter sequence. c: Same view as b, but with the outer TM1 helices removed to show more clearly the inner TM2 helices (red) and pore helices (white). d: The tetramer with the front and back subunit removed to show clearly the turret (white), the filter (GYG, red), and gate region (green). e: Side view of the KcsA tetramer, giving dimension, and showing the two belts of aromatic amino acids facing the membrane (black). [From Doyle et al. (80). Reprinted with permission from AAAS.]
conducting state, the network of hydrogen bonds between the filter sequence and the pore helix has the tendency to drive the filter into a new and nonconducting conformation. Chakrapani et al. (2007) models this network of bonds as a spring, set on the filter to balance the activated and inactivated conformations (42). The network includes bonds between charged residues, explaining why the entry into the inactivated state is voltage dependent (42, 43). A nonconductive conformation of the filter is observed when KcsA was crystallized in a low K+ concentration. This conformation likely corresponds to a vivo state when the bottom gate is closed (below) and the filter is facing a lumen of low [K+] (414). When there is no K+ to occupy the filter, it collapses. This property explains C-inactivation, the voltage dependence of KcsA channel (149, 324), and the dependence on K+ driving force in the case of the yeast TOK1 K+ channel (219) (below).

Many variants of the filter were found in a survey of 270 K+-channel sequences (190, 192). Instead of the usual TVGYG found in the KcsA filter, genes predicting channel subunit with TVGDG and TVGDA filter sequence, but otherwise similar to KcsA were found in Bacillus cereus and B. anthracis, respectively. The former channel protein has been crystallized and its structure solved (347). It shows that K+ coordination sites 3 and 4 are like those of KcsA, but sites 1 and 2 have become a vestibule where cations cannot bind specifically. Competition experiments of 86Rb+ flux showed that K+ and Na+ compete, but Li+ or N-methyl-D-glucamine+ does not. Such channels are now called NaK channels. A molecular dynamics simulation indicates that the lack of selectivity of NaK reflects the differences in hydration of the ions in the filter (278).

B) THE KCsA GATE. Except in the filter, the K+ in transit is surrounded by shells of polarized water molecules. Correspondingly, except in the filter, the pathway is lined largely with hydrophobic amino acids of four TM2 inner helices, preventing the lingering of the hydrated K+. The ion pathway of the filter is connected to a gate near the opposite surface facing the cytoplasm. The gate of KcsA is formed by the convergence of the four TM2 helices that are slightly tilted and kinked (80). These helices come into contact at the point near the bundle crossing to occlude the ion pathway in the closed conformation. In KcsA, the contact is at residue 112–119, at the COOH-terminal end of TM2 (31, 80). Near the center of KcsA's TM2 is glycine, which acts as the “gating hinge,” allowing the COOH-terminal half to move. The gate likely opens at the hinge and by rotating the TM2 helices counterclockwise away from the central axis. There are concerted movements of the TM1 helices as well (31, 298). As described below, various “gating principles” (physiologically relevant ligands, voltage, stretch force, etc.) are believed to dilate the gate by moving the COOH-terminal ends of the inner helices outward. Note that parts of the NH2- and the COOH-terminal sequence are not represented in the KcsA crystal structure. EPR spectroscopy shows that the NH2-terminal peptide forms a helix lying at the membrane-water interface, and the COOH-terminal peptide forms a helical bundle in the cytoplasm (60). The functions of these helices are unclear.

The movement of the gate and the dynamic filter, which is often referred to as the “second gate,” accounts for complex kinetic behavior, even for a relatively simple channel like KcsA (11, 42, 43). In general, a channel rarely visits its open state without a stimulus, which is referred to as the “gating principle” in channel biology. In the case of KcsA, proton opens the channel in the laboratory, although the physiological significance of this pH gating in vivo is unknown. In this case, pH apparently affects both the inner gate as well as the filter (42, 43, 55). Opening and closing of the gate entail a concerted movement of the inner helix (TM2) and the outer helix (TM1) that interact in a hand-and-handle manner (see below).

C) THE LIPIDS AROUND KCsA. Unlike soluble proteins, ion channels reside in the membrane and function to ferry ions across the lipid bilayer, a special environment. There is obviously an intimate relation between channels and their surrounding lipids. The KcsA tetramer is more stable in lipid bilayer compared with detergent micelles. In NMR studies, KcsA dissociating organic solvent has a large effect on the acyl-chain ordering, suggesting that the bilayer stabilizes KcsA by the lateral pressure in the acyl chain region (377, 378).

Two types of protein-lipid interactions can be distinguished: annular or nonannular. Annular interactions occur between the embedded portion of protein surface and the surrounding lipids. Nonannular interactions entail tighter association of the lipid into crevices between transmembrane helices. Such a tightly bound lipid often acts like a subunit or cofactor of the protein. Vallyaveetil et al. (376) showed in a KcsA crystal that there are four lipid molecules, each located between two subunits at the part of the channel that faces the periplasmic side. This is surprising because the KcsA protein examined here has been purified and crystallized in detergent with no added lipids. Note that the conditions required for crystallization are extremely harsh. Cocystalization indicates specific and strong binding. The head group of the bound lipid was not resolved, but biochemical analyses showed that the lipids are likely phosphatidylglycerol (PG), even though the E. coli membrane, in which the KcsA is produced, has more of the neutral (zwitterionic) phosphatidylethanolamine (PE) than the anionic PG. The head group is apparently held in place by nearby arginines and its fatty acid tails wedged into crevices formed by helices from adjacent subunits. Fluorescence quenching experiments largely agree with these findings. They showed that charged interaction is crucial in this nonannular binding.
The nonannular binding constant of PG was determined to be $\sim 3$ mole fraction $^{-1}$, corresponding to $\sim 40\%$ occupancy by PG in the E. coli membrane (233). The details of arginine-lipid binding are revealed in a molecular dynamics simulation (73). Using $^{86}\text{Rb}^+$-uptake to assess KcsA function in reconstituted lipid vesicles, Valiyaveetil et al. (376) showed that uptake occurs in vesicles containing negatively charged lipids (3PE:1PG) but not in those with only neutral lipids (3PE:1PC). It seems likely that this functional requirement of PG is at least in part to fulfill the binding seen in the crystal structure (376). Mass spectrometry by others showed that KcsA forms noncovalent complexes preferentially with PG, less so with PE but little with PC (71). In a separate assay, Valiyaveetil et al. (376) investigated the requirement for KcsA refolding. KcsA is a very stable tetramer, requiring organic solvents to unfold into monomers. They found that lipids are required to refold the monomers into functional tetramers (376). Here the lipids apparently do not act as cofactors, but provide the bilayer environment. Negatively charged lipids are not required for the refolding.

Unlike KcsA, purification and crystallization of Kv1.2, an animal voltage-gated K$^+$ channel (below), required a mixture of phospholipids and detergents (210). When channels are crystallized in a lipid-detergent mixture, many lipids surrounding the channel can be resolved or partially resolved. A hybrid voltage-gated K$^+$ channel has been shown to be surrounded by 64 lipids, 16 per subunit. These lipids are organized in two layers, reminiscent of the membrane. The outer layer, especially, resembles the outer leaflet of the bilayer. The lipid heads are not at the same level, and the tails wedge between proteins filling up space. It appears that the protein-lipid interaction displaces the lipids from their position in the pure lipid bilayer (212).

The more general interactions between the channel protein and the surrounding annular lipids are more difficult to investigate. Such lipid-protein interactions are less likely to survive the harsh condition of crystallization and cannot be revealed by crystallography, especially not in proteins purified through detergents. In fluorescence quenching experiments, annular binding at the periplasmic side of KcsA was found to be nonspecific with respect to the lipid’s head group charge. Binding of anionic lipids are stronger at the cytoplasmic side, however, reflecting the pattern of surface charge distribution of KcsA itself (233). In a molecular dynamics simulation (73), the channel protein reduces the lateral mobility of the lipids that dynamically interact with its surface, reducing their diffusion to half the rate of the bulk lipids in the bilayer. About 30 lipid molecules contact KcsA with their head groups, which include the glycerol and the acyl oxygens. About 40 make contact with their acyl tails. Interactions, defined as protein-lipid interatomic distance being $\leq 3.5$ Å, are concentrated in two broad bands ($\sim 10$ Å) that level with the two interfaces of the bilayer. Few contacts are made near the bilayer’s center, reflecting the mobility of the acyl tails.

Most membrane proteins of known 3-D structures have aromatic belts at the interface, because aromatic amino acids favor this location (390, 397). Such belts apparently stabilize the protein structure and may play roles in membrane-protein folding (390) and possibly in conformation changes (78). Tryptophan and tyrosine, amphipathic aromatic residues, are found to form bands on membrane proteins at levels corresponding to the interfaces (390, 397). Experiments showed that, when properly placed, they help transmembrane peptides to resist dislocation and are thought to anchor membrane proteins to the bilayer. As summarized above, KcsA has two such belts of aromatic amino acids (Fig. 6F). In simulation, the aromatic rings of KcsA are found to be roughly normal to the bilayer plane, with the polar moieties (the ring nitrogen of Trp and the hydroxyl of Tyr) nearest the membrane interfacial region. So oriented, these aromatics apparently have their polar regions interacting with the lipid head groups and their hydrophobic region with the lipid tails. Basic residues, arginines in the case of KcsA, are also seen to form hydrogen bonds with the lipid phosphates, mostly at the intracellular interface (73).

2. K$^+$ channels with additional modules

A tetrameric assembly of subunits each with an inner transmembrane helix, a short pore helix, filter sequence, and an outer transmembrane helix is apparently the core of all K$^+$ channels. This core motif is found in other cation channels including Na$^+$ channels, Ca$^{2+}$ channels, cyclic nucleotide-gated (CNG) channels, and TRP channels. It is likely an early evolutionary design, onto which later embellishments can be added. Such embellishments include voltage-sensing domain, glutamate-, Ca$^{2+}$-, or cyclic nucleotide-binding domains, etc. It illustrates that, like other proteins, channels are modular, and the permutation of modules yields the diversity we see in the biological world. The modularity of channel’s construction is not only evident by surveying existing channel variations, but has been shown in two experiments.

KcsA with the TM1-P-TM2 structure is activated by a proton. On the other hand, the subunit of the depolarization-activated Shaker channel comprises six TM helices by 10.220.32.247 on April 18, 2017 http://physrev.physiology.org/ Downloaded from http://physrev.physiology.org/
linker acts as the “hand” and the lower end of S6 receptor acts as the “handle” of the gate. Because the hand-and-handle interaction is stereochemically specific, they need to be homologous. This set of experiments illustrates the modular principle of channel construction and anticipated some of the conclusions derived from crystal structures.

The modularity of K+ channel construct is also evident in the chimera experiment by Ohndorf and Mackinnon (282). They added cyclic nucleotide-binding domain (CNBD) of Rhodopsseudomonas palustris to the COOH-terminal ends of KcsA’s inner helices and showed that the construct opens upon the addition of cAMP. As described below, CNBD is a module in the cytosol and is attached to the membrane-embedded core at the end of its inner helix directly by peptide linkage. Here the force that opens the membrane-gated channels (282). They added cyclic nucleotide-binding domain (CNBD) of MloK1 continued to be analyzed functionally and not by a hand-and-handle mechanism of the voltage-gated channels. 

A) CNMP-GATED K+ CHANNELS (MIOK, MMAK). Several K+-channel cores trail by sequences of CNBD can be found in α-proteobacteria and a cyanobacterium (190, 192). A 6-TM K+ channel (Fig. 5B) with CNBD from Mesorhizobium loti, MloK1, has been expressed, and its 86Rb+ flux shown to be activated by cyclic nucleotide (274). The crystal structure of MloK1’s CNBD has been solved (53). In this structure, the CNBD domains are in forms of dimers, leading to the proposal that the gate operates possibly as a dimer of dimers. A recent electron microscopy structure of the entire MloK1 protein, however, found that each CNBD in the tetramer to be independent, supporting the mode of gate opening by cyclic nucleotide (cNMP) similar to that in the animal CNG or HNG channels (50). MloK1 continued to be analyzed functionally with 86Rb+ flux but not by electrophysiology (273). A CNMP-gated channel, MmA K, from the gram-positive bacterium Methanobacterium thermoautotrophicum, has recently been cloned and expressed in E. coli (191). Here, the authors generated giant E. coli spheroplasts (see sect. iC), excised membrane patches from them, and examined the electric activities of MmA K therein. Both macroscopic and microscopic currents can be observed. Activation by CNMP and inactivation by acidic pH are evident. The growth of E. coli expressing mmaK is suppressed by the external addition of K+ but not Na+. This K+-specific phenotype should make future genetic dissection of MmA K promising. B) RCK-OPERATED K+ CHANNELS (MTHK, KCH, TVOK). In eu- karyotes, Ca2+ actuates the changes in actin-myosin and dynemein-tubulin based motility. This key second messenger also controls secretion, fertilization, and many important animal functions. Like many enzymes, some ion channels are also directly controlled by Ca2+. Ca2+ binding inactivates some channels but activates others. Channels that are opened by Ca2+ are found in archaea and bacteria, although they seem to require much higher concentrations of Ca2+ than the eukaryotic channels do.

MthK, a Ca2+-activated K+ channel of the archaeon Methanobacterium thermoautotrophicum, has been crystallized and its structure in an open conformation solved (147). In its primary structure, each MthK subunit has the TM1-P-TM2 sequence (Fig. 5A) followed by an extension of some 200 amino acid residues. These residues form a module that contains the Rossmann fold, which has a scaffold of α-helices and β-sheets in a dimeric arrangement. This fold has a key sequence, GXGXXG, indicative of NAD binding. Such modules are found in many K+ transporters or K+ channels in a variety of prokaryotes and is called the KTN (K channel transport and NAD-binding) domain (324, 325). A related structure, called the RCK (for regulator of the conductance of K+) domain, retains the αβ-scaffold and the dimeric arrangement without the GXGXXG. Instead of dinucleotides, RCK of MthK binds divalent cations, such as Ca2+ (147, 149). KTN or RCK assembles in pairs, and the ligand (dinucleotides or Ca2+) is coordinated at the dimer interface. Surprisingly, besides the full-length 200 kDa mthk gene product, purified MthK channel protein contains a 26-kDa peptide, indicative of translation from an internal AUG. This smaller peptide corresponds to a free RCK domain. The crystal structures (147, 149) and other studies led to a model in which the channel is a hetero-octamer, with the MthK transmembrane tetramer assembles with eight RCK domains: four covalently linked to the four TM2 helices and four additional ones assembled onto them at the cytoplasmic side. The crystal was formed in the presence of 200 mM Ca2+ and eight Ca2+ found coordinated in clefs between adjacent RCK domains. MthK reconstituted into lipid bilayers can indeed be activated by Ca2+. It was proposed that the eight RCKs form a preassembled gating ring. Ca2+ binding changes the conformation of the “gating ring” generating the outward force to pull apart the converged ends of the four TM2 helices, through a 17-residue linker not resolved in the crystal structure. However, there are reasons to question the need of the free RCK domains in gating from the studies of other RCK-bearing bacterial channels (193, 314). Recently, the MthK activities were examined in patches excised from mthk-expressing giant E. coli spheroplasts (191). Besides Ca2+, Cd2+ also activates this channel. MthK desensitizes in the continued presence of these ligands and inactivates in acidic pH. Unlike the “gating-ring” model, biochemical studies of isolated RCK domains showed that they are dimeric in acid pH, regardless of Ca2+. In basic pH, however, they convert from monomers to multimers upon the addition of Ca2+. These findings led to another model in which multimerization is a part of the gating mechanism (191). The physiological meaning of this activation with millimolar Ca2+ is unclear.
since Ca\(^{2+}\) as a second messenger operates at micromolar concentrations in eukaryotes, and the signaling roles of Ca\(^{2+}\) in prokaryotes, if any, are uncertain.

An RCK-containing K\(^{+}\) channel from the archaeon *Thermoplasma volcanium*, TvoK, has recently been cloned and reconstituted into planar lipid bilayers for functional analyses (289). The 160-pS K\(^{+}\) conductance was found to be activated by millimolar Ca\(^{2+}\) or Mg\(^{2+}\). The molecular mass and subunit composition of TvoK are consistent with the hetero-octameric structures proposed for MthK.

In 1994, Milkman (247) recognized a gene in *E. coli* that corresponds to a K\(^{+}\) channel, called Kch, because it predicts six transmembrane helices (Fig. 5B) similar to the *Shaker* channel of *Drosophila*. Its TM6 is attached to an RCK domain, which was later isolated and crystallized, solving the first RCK structure (149), though its potential ligand is yet to be identified. The electric activities of Kch have never been reported, despite various attempts. However, genetic experiments indicate the Kch does function in vivo. Kuo et al. (2003) isolated GOF *kch* mutants that failed to grow in millimolar added K\(^{+}\) but not Na\(^{+}\) (193). This K\(^{+}\) susceptibility phenotype vanished when the canonical K\(^{+}\) filter sequence is further mutated, showing that it is the passage of K\(^{+}\) that kills cells. All seven GOF mutations selected with this phenotype after a random mutagenesis were found to be in the RCK domain. Removing the internal ATG of the GOF mutants does not change the phenotype, indicating that the corresponding free RCK is not needed for K\(^{+}\) passage in vivo. External H\(^{+}\) suppresses the GOF phenotype. This observation supports a model that Kch is not for bulk K\(^{+}\) uptake, but functions to regulate membrane potential under certain stress (189, 190, 193).

C) VOLTAGE-GATED CHANNELS (KVAP, MVP, KVLM). The intellectual history of ion channel is rooted in the study of the excitable membrane. A big step forward was the separation of the regulated rise and fall of Na\(^{+}\) and K\(^{+}\) permeability that underlie the generation of action potential of axons in the 1950s. Voltage-clamp experiments show that the specific channels involved are governed by membrane voltage. The material basis of the permeability was not understood. Much of our knowledge on how voltage drives open the gate is derived from the crystal structure of archaeal KvAP and the animal Kv1.2 solved by the MacKinnon laboratory (148).

KvAP is a depolarization-activated K\(^{+}\) channel of the hyperthermophilic archaeon *Aeropyrum pernix*. Like the *Shaker* channel, each KvAP subunit comprises six transmembrane helices (Fig. 5B), with TM4 bearing the gating charges and the canonical K\(^{+}\)-filter sequence. The TM5 and TM6 flank the pore helix and the filter sequence. In the tetramer, they converge to enclose the pore and form the filter, much like the TM1 and TM2 of KcsA (148). This portion is therefore the core, or the permeation domain of the protein. TM1 through TM4 of each subunit forms a separate domain, surrounding this core. In this structure, TM1 and TM2 are near the TM5-P-TM6 core with the TM3 and TM4 at the periphery. There is surprisingly little direct contact between the core and the four peripheral domains, considered the voltage-sensing domains. This modular arrangement has been anticipated by a chimera experiment described above, in which a precise grafting of *Shaker*’s voltage-sensing domain onto the core of KcsA confers voltage sensitivity (221). Purified KvAP has been incorporated into planar lipid bilayers and found to correspond to a K\(^{+}\)-specific unitary conductance of 170 pS. Its macroscopic current is found to be activated by depolarization (327).

The current understanding of voltage sensing, based on the structures of KvAP and Kv1.2, is as follows. The voltage sensor is an electromechanical coupling device, allowing the transmembrane voltage to bias the conformation of the domain. The sensor consists of the voltage paddle and the S1-S2 region with which the paddle interacts. (“S,” same as “TM,” is used here to make easier the narrative and reference to the original papers) The voltage paddle is a helix-turn-helix structure comprising S3b (the COOH-terminal portion of S3) and S4 helices running antiparallel (3). S4 bears positively charged arginines spaced 3 residues apart, named R0, R1, etc. counting from the NH\(_2\) terminus. Seen in the open state, the paddle is located near the outer surface, with one side facing the lipids, and the other side resting on the outer half of S1 and S2 helices, with R0, R1, and R2 interacting with the phosphoester head groups of lipids, and R3 and R4 with a cluster of glutamates of surrounding helices, including S2 (212). The position of the paddle in the closed state is uncertain, since there is no clear closed state structure as yet. Accessibility and other studies suggest that the paddle translates inward, moving all the gating charges beneath a S2 phenylalanine at the midpoint (212).

In the KvAP structure, the voltage-sensing paddle lies along the membrane plane positioned at the inner lipid-plasma interface. The detergent used apparently distorted the native structure (200). The resolution of Kv1.2,
a *Shaker* homolog from rat brain, amended the KvAP structure (210, 211). Here, the TM1-TM4 module remains outside the core, but with the TM3-TM4 voltage-sensor paddle erected, crossing the bilayer. TM4 is seen as a long helix with its NH$_2$-terminal side bearing the regularly spaced arginines. At its COOH-terminal end, it connects to a helical S4-S5 linker through a bend. Raising the paddle from the position in the KvAP crystal fits the Kv1.2 structure almost perfectly. Long et al. (212) recently showed in the crystal of an open-state Kv2.1-Kv1.2 chimera, formed in lipids and detergent, that the voltage paddle tilts outward, with one entire face exposed to the lipids. It is postulated that the paddle moves between this outward and an inward position upon voltage changes to drive the conformation (state) changes of the protein. The S4-S5 linker is an amphipathic helix running parallel to the membrane plane on the cytoplasmic side. It crosses over TM6 and makes many amino acid contacts with TM6’s cytoplasmic end. In the open state, TM6 bends at a hinge near the region of contact to be nearly parallel with the membrane, forming the “handle” for the S4-S5 linker. The presumed flexible hinge allows the lower end of TM6 to dilate or constrict the pathway. In many voltage-gated K$^+$-channels, the hinge is a glycine (e.g., KvAP) as in KcsA and KirBac; in others, it is the sequence Pro-X-Pro (X being any residue, e.g., Kv1.2). By analogy, the S4-S5 linker can be viewed as the “hand” and the COOH-terminal end of TM6 the “handle,” such that the gating force from the movement of TM4 is transmitted to the TM6 bundle crossing.

Alabi et al. (3) recently showed that voltage paddle is itself a portable domain. When the portion from the juncture between S3a and S3b to just past the first four critical Arg residues of S4 from KvAP replaces the comparable portion in Kv2.1, the chimera channel retains voltage sensitivity. Different tarantula toxins attack different voltage-sensitive channels specifically. The chimera shows sensitivity to the toxin that is specific for KvAP. Note that the voltage paddles of KvAP and Kv2.1 are not very similar in sequence. Therefore, this work not only shows that the voltage sensor is modular, but also that the paddle is not packed tightly with other protein domains forming extensive bonds. It appears to be in a relatively unconstrained environment, most likely the lipids. Nature uses this voltage-sensing module in a different context. Besides these K$^+$ channels, they are found in the voltage-activated proton channel (Hv1) (319, 336) and the voltage-sensitive phosphatase of *Ciona* (Ci-VSP) (258).

A gene homologous to *Shaker* and relatives is found to encode MVP, a hyperpolarization activated 37-pS K$^+$ channel from the archaeon *Methanococcus jannaschii* (344). *MVP* transgene complements the K$^+$-uptake defects of both mutant *E. coli* (133) and yeast (344). Although MVP responds to hyper- instead of depolarization, it is modeled to have the same S4 movement, but S4’s coupling with the gate is such that the S4 position corresponding to open in *Shaker* is close in MVP.

A survey shows that there is a far greater variation in the S4 gating charges among prokaryotes than in animals. S4’s with as little as one and as many as eight charges can be found (192). KvLm, the *Shaker*-type channel from the gram-positive bacterium *Listeria monocytogenes*, with only the three of the positively charged S4 residues has been investigated. Expressed and examined in giant *E. coli* spheroplasts, the open probability of KvLm nonetheless shows a steep dependence on depolarization (335). Further mutational investigation shows that the voltage sensor of KvLm likely has a structure similar to that of animal *Shaker* channels (224).

**D) INWARD RECTIFIERS (KIRBAC).** Inward rectifiers are K$^+$ channels that behave like diodes. They open when the membrane is polarized or hyperpolarized, allowing K$^+$ to enter the cell, but the open channel is blocked by internal Mg$^{2+}$ or polyamines when the polarity is reversed to prevent K$^+$ exit. Durell and Guy (82) recognized a family of prokaryotic inward-rectifier K$^+$ channels (KirBacs) by sequence similarity to eukaryotic homologs, including the sequence of the large COOH-terminal region unique to inward rectifiers (82). KirBacs are found in many species of *Burkholderia, Magnetospirillum magnetotacticum,* *Nostoc punctiforme,* (82) *Synechocystis sp., Ralstonia solanacearum,* *Solibacter usitatus,* *Chromobacterium violaceum* (369), and others. The crystal structure of KirBac1.1, the inward rectifier channel from the bacterium *Burkholderia pseudomallei,* has been solved in a closed state by Kuo et al. (189). Like the full-length KcsA, a “slide helix” at the NH$_2$ terminus positioned at the level of the lipid-water interface, precedes a TM1-P-TM2 core (Fig. 5A). The slide helix apparently interacts with the phospholipid head groups (76). In this crystal structure, the core of KirBac1.1 is similar to that of KcsA, except that the pore helices do not point at the center of the channel, a feature considered a key to the closed channel here. The structure of the filter is largely the same as KcsA. Similar dynamics of the filter are seen in a molecular dynamics (MD) simulation (73). This core is trailed by a large COOH-terminal cytoplasmic domain, which is made of $\beta$-sheet structures and contains the putative Mg$^{2+}$-biding glutamate residues. Although the core is in a fourfold radial symmetry, the cytoplasmic domains are in forms of a dimer of dimers. The channel closes at the end of TM2, forming a hydrophobic seal by the convergence of phenylalalines. The inner helix (TM2) of KirBac1.1 has a hinge glycine like in KcsA. Channel opening is modeled to be a concerted outward movement of the slide helices and the TM1 helices, making room for the TM2 helices to bend at their glycine hinges. This notion is further supported by the electron crystallography of two-dimensional crystals of KirBac3.1 of *M. magnetotacticum* in two conformations at 9-Å resolution (188). In a molecular dynamics
simulation, aromatic residues appear to be more evenly distributed in the closed state but shift toward lipid-water interface, suggesting that aromatics preference to be at the interface is being used in gating or conformational stabilization (78).

Sun et al. (369) tested a large number of KirBacs for their ability to complement the K$^+$ auxotrophy of a TK strain of E. coli lacking the three major K$^+$ uptake mechanisms (Kdp, Trk, Kup). Some of them complement; some complement less well at a higher temperature; some do not complement. The results show differences among KirBacs in their ability to take up bulk K$^+$ in this heterologous setting.

A current topic in animal channel physiology is the modulation by phosphatidylinositol 4,5-bisphosphate (PI(4,5)P$_2$). Pure KirBac1.1 and KcsA have been reconstituted into liposomes of defined lipid composition and assayed for $^{86}$Rb$^+$ uptake. KcsA is not affected but Kir. Bac1.1 is inhibited by PI(4,5)P$_2$. This result shows that there is a direct interaction between PI(4,5)P$_2$ and KirBac, excluding modulation through cytoskeletal or other intermediary elements (91). The physiological significance needs to be addressed, since phosphatidylinositol has limited distribution and is in small amounts among bacteria.

E) GLUTAMATE RECEPTORS (GLUR), GluR, glutamate receptors, bind glutamate or related neurotransmitters to open a cation conductance in excitatory postsynaptic membranes. Animal GluR are sorted into subfamilies depending on their ligands (NMDA, AMPA, or kainate). Each GluR is a tetramer. The core of each subunit of animal GluR consists of an extracellular region called S1 followed by the first transmembrane TM helix called M1, the pore P (with the filter sequence), the second TM helix M2, the second extracellular region S2, and a third TM helix M3. The external S1 and S2 form the glutamate-binding site. There are additional NH$_2$- and COOH-terminal domains up- and downstream of this core. A sequence homolog was found in Synechocystis PCC6803 that has the S1-M1-P-M2-S2 core, but lacks M3 and the additional NH$_2$- or COOH-terminal domains. It was named GluR0 (47). S1 is followed by TM1 beginning from the outside instead of the cytoplasmic side, thus placing the TM1-P-TM2 permeation and filtration core in an orientation opposite that of KcsA (Fig. 5C). Key residues for glutamate binding in animal GluRs are conserved in the S1 and S2 region of GluR0, and the corresponding protein overexpressed in E. coli has the expected ligand-binding properties. The pore region contains the perfect K$^+$-selectivity sequence TVGYGD identical to that of KcsA. GluR0 expressed in HEK cells or Xenopus oocytes indeed produce K$^+$ currents, found to flow through animal ionotropic glutamate-receptor channels (47). The ligand-binding core of GluR0 was crystallized and its structure solved. This core consists of two domains separated by a deep cleft. The larger Domain 1 includes $\alpha$ helices A, B, C, from S1 and J, K, from S2. Domain 2 includes $\alpha$ helices F,C, H of S2. L-Glutamate, the ligand, binds in the cleft between the domains. The symmetry of the ligand-binding core suggests that the tetrameric channel is assembled from two dimers (243). A molecular dynamics simulation of GluR0 shows that concerted K$^+$-water move along the selectivity filter, as in the simulations of KcsA and the K$^+$ exit is associated with the opening of the hydrophobic gate formed by the M2 termini (8). GluR0 homologs can be found in other bacteria. Interestingly, those from Prochlorococcus marinus, Trichodesmium erythraeum, and two species of Synechococcus have the canonical K$^+$-filter sequence, and those from Silicibacter pomeroyi, Magnetospirillum magnetotacticum, and two sequences of Nostoc punctiforme have the filter and pore sequences more like those of the cation nonspecific eukaryotic glutamate receptors (182). A preliminary X-ray crystallographic analysis of the one from Nostoc punctiforme has been reported (199).

B. Na$^+$ Channels (NaChBac)

The animal depolarization-activated Na$^+$ or Ca$^{2+}$ channels are similar to K$^+$ channels in structure. However, each Na$^+$ or Ca$^{2+}$ channel comprises 24 TM domains that are covalently linked, with each 6-TM repeat being comparable to a Shaker-type 6TM K$^+$-channel subunit. Although high-resolution structures are not yet available, the Na$^+$/Ca$^{2+}$ channel filter is likely lined with four glutamic or aspartic residues. A 6-TM gene product from Bacillus halodurans, an extremophile that lives in high salt and high pH, has been expressed in mammalian cells and found to form a 12-pS channel that is activated by depolarization, followed by in- and deactivation. It is similar to L-type Ca$^{2+}$ channel in being blocked by dihydropyridines, nifedipine, and nimodipine. However, this channel is selective for Na$^+$, not Ca$^{2+}$, and is named NaChBac (321). It is speculated that the Ca$^{2+}$ selectivity may require a flexible tertiary structure around the acidic residues provided by the four nonidentical repeats in the animal channels, but are not provided by the NaChBac homotetramer (321). Mutational analyses showed that increasing the number of aspartic substitutions in the filter sequence (LESWAS) increases the Ca$^{2+}$ selectivity. In the series of mutations LESWAD, LEDWAS, LEDWAD, and LDDWAD, the first has the lowest and the last the highest Ca$^{2+}$ selectivity (404). The TM4 of NaChBac subunit bears arginine residues spaced every third residue apart, as in KvAP and Kv1.2 (321). The gating-charge movement of NaChBac has been studied electrically (197) and by real-time fluorescence spectroscopy (32). The acidic residues on TM2 and TM3 of NaChBac that likely neutralize the arginines of TM4 have also been mutated and analyzed (26). NaChBac has homologs in at least 11 other bacteria,

Physiol Rev • VOL 88 • OCTOBER 2008 • www.prv.org
forming a family called Na,Bac. Two members, Na,PZ from Paracoccus zeaxanthinifaciens and Na,SP from Silicibacter pomeroyi, have also been expressed and examined in mammalian cells (169). A role in controlling the Na⁺-driven flagellar motility for some Na,Bac channels has been proposed (321). Na,Bac channels appear to be a good source for material for crystallography towards a better understanding of Na⁺ and Ca²⁺ filtration.

C. Ammonium Channel (Amt, Mep)

Ammonium channels have no relation to the K⁺ or Na⁺ channels described above. Transport of nitrogen is important to most organisms, since it is a major element in life and available nitrogen is often limited in the environment. Though toxic to animals, ammonium is used by prokaryotes, fungi, and plants as the preferred nitrogen source. Most cellular organisms have AMT/Rh family proteins to transport ammonium, except some parasites and those that use specialized nitrogen source, such as urea. Ammonium channels are called Amts (ammonium transporters) in the bacteria literature and Meps (methylamine permeases) in the fungal literature. They have some sequence homology to Rh, the Rhesus blood group antigens (294). The ones from Escherichia, Salmonella, Shigella, Vibrio, and Yersinia (145) have been crystallized and their structures solved at high resolution (83). As predicted from the double-barrel gating, it exists as a homodimer. The two subunits meet at an extensive contact surface, but each has its own pore. Each subunit comprises 18 α-helices (A through R) arranged in two structurally similar halves. Helix A–G and helix H–R have only weak sequence homology, but are arranged in two structurally similar halves. Helix A–G and helix H–R have only weak sequence homology, but are packed similarly. The two halves ran antiparallel, creating a pseudo-twofold symmetry, and wrap around a common center forming the Cl⁻-binding sites. The helices vary in length and are all tilted with respect to the membrane normal. It is so arranged that the NH₂ termini of helix D, F, N, and R, i.e., the positive ends of their helical dipoles, point into the central Cl⁻-binding site. In the crystals, a Cl⁻ or Br⁻ occupies this site coordinated by main-chain nitrogens and side-chain oxygens of nearby helices. This central binding site is connected through tunnels to the wider, water-filled vestibules, exposed to the intra- and extracellular solutions. Several crystal structures taken together show that the permeant anion likely passes through three binding sites, the central one being the one described above. All three sites have been seen occupied in various crystals with different anions (271). In the first set of crystals, just above the pseudo-twofold axis, Glu148 of helix F projects into the pore, blocking the presumed external Cl⁻-binding site. The crystallized form appears to be that of a “closed” channel. This glutamate residue likely swings out of the way in the open conformation to allow Cl⁻ passage. Mutation analysis indeed shows that the carboxyl group of this glutamate acts as a pseudosubstrate, occupying the external site to block passage (84).

Accardi et al. (1) succeeded in reconstituting pure CIC-ec1 into planar bilayer and made the surprising discovery that it acts like an exchanger rather than a channel. Only the macroscopic current of CIC-ec1 can be registered, with noise properties consistent only with very small unitary conductance (<0.1 pS). The current has no

D. Cl⁻ “Channels” (CIC-ec1)

Cl⁻ channels are entirely different from the cation channels described above. As with the cation channels, Cl⁻ channels were first studied electrophysiologically in animal cells. These studies showed that the Cl⁻ channel behaves as “double barreled”, i.e., two ~8-pS units that can gate independently and rapidly (ms) as well as together and slowly (s). There is also a strong coupling between Cl⁻ conduction and gating. First cloned from the fish Torpedo, the animal CIC-0 has many homologs including those in bacteria, such as the enteric species of Escherichia, Salmonella, Shigella, Vibrio, and Yersinia (145). The ones from Salmonella typhimurium and E. coli (CIC-ec1) have been crystallized and their structures solved at high resolution (83). As predicted from the double-barrel gating, it exists as a homodimer. The two subunits meet at an extensive contact surface, but each has its own pore. Each subunit comprises 18 α-helices (A through R) arranged in two structurally similar halves. Helix A–G and helix H–R have only weak sequence homology, but are packed similarly. The two halves ran antiparallel, creating a pseudo-twofold symmetry, and wrap around a common center forming the Cl⁻-binding sites. The helices vary in length and are all tilted with respect to the membrane normal. It is so arranged that the NH₂ termini of helix D, F, N, and R, i.e., the positive ends of their helical dipoles, point into the central Cl⁻-binding site. In the crystals, a Cl⁻ or Br⁻ occupies this site coordinated by main-chain nitrogens and side-chain oxygens of nearby helices. This central binding site is connected through tunnels to the wider, water-filled vestibules, exposed to the intra- and extracellular solutions. Several crystal structures taken together show that the permeant anion likely passes through three binding sites, the central one being the one described above. All three sites have been seen occupied in various crystals with different anions (271). In the first set of crystals, just above the pseudo-twofold axis, Glu148 of helix F projects into the pore, blocking the presumed external Cl⁻-binding site. The crystallized form appears to be that of a “closed” channel. This glutamate residue likely swings out of the way in the open conformation to allow Cl⁻ passage. Mutation analysis indeed shows that the carboxyl group of this glutamate acts as a pseudosubstrate, occupying the external site to block passage (84).
voltage or time dependence, prominent in animal ClCs. Its reversal potential deviates significantly from the one expected from the Nernst's equation of Cl\textsuperscript{−} gradient and from the Goldman-Hodgkin-Katz equation of various Cl\textsuperscript{−} and H\textsuperscript{+} gradients, but is consistent with the expectation of a strictly coupled transport of \(\sim 2\) Cl\textsuperscript{−} for 1 H\textsuperscript{+}. Furthermore, either ion can be made to flow against its own electrochemical gradient. How H\textsuperscript{+} is transported remains unclear. Glu148 is apparently crucial. E148A mutation causes CIC-er1 to allow Cl\textsuperscript{−} to flow down gradient and not to show the H\textsuperscript{+}-dependent transport. The exact mechanisms of ion permeation through ClCs are under investigation. The CIC-ec1 finding echoes a finding in Paramecium, where the XntA gene product behaves as channel in electrophysiology but shows an exchanger-like sequence (see sect.IV). The current understanding challenges the classical distinction made between ion channels and ion exchangers. Current literature now divides the animal ClCs into two classes: those in the plasma membrane (ClC-0, -1, etc.) acting as channels and those in vesicular membranes (ClC-4, -5, etc.), being more homologous to the bacterial ClCs, acting as antiporters. As to the role of bacterial ClCs, evidence supports the view that they are exchangers. Current literature now divides the animal ClCs into two classes: those in the plasma membrane (ClC-0, -1, etc.) acting as channels and those in vesicular membranes (ClC-4, -5, etc.), being more homologous to the bacterial ClCs, acting as antiporters. As to the role of bacterial ClCs, evidence supports the view that they are exchangers.

E. Roles of Ion-Specific Channels in Prokaryotic Physiology

The role of Cl\textsuperscript{−} channel/exchanger in enteric bacteria is described above (145). However, the roles of the cation channels (KcsA, KvAP, MthK, KirBac, NaK, NaChBac, etc.) are unknown. Their roles in the native settings are sometimes difficult to investigate, since some are externophiles, requiring special laboratory conditions for cultivation. Even more frustrating is the fact that we cannot yet fathom the roles in bacteria commonly studied in microbiology, including E. coli, B. subtilis, and S. lividans (132). In the study of the Kch GOF mutants of E. coli, external H\textsuperscript{+} suppresses the ability of K\textsuperscript{+} to block growth of these mutants. About a 10 times increase in [H\textsuperscript{+}] suppresses the toxicity caused by a 10 times increase in [K\textsuperscript{+}] on these mutants. This finding implies that the parameter being regulated has a logarithmic relation to [K\textsuperscript{+}], probably the equilibrium potential of K\textsuperscript{+} (\(E_K\)). E. coli growth depends on the proton-motive force \(\Delta p\). \(\Delta p\) comprises membrane potential (\(V_m\)) and \(\Delta pH\), the pH gradient across the membrane. It is therefore thought likely that the GOF Kch channels open uncontrollably, locking the \(V_m\) near the \(E_K\). When external [K\textsuperscript{+}] is raised, \(E_K\) and therefore \(V_m\) are small and the total \(\Delta p\) is too small to sustain growth. Increasing external [H\textsuperscript{+}] raises the \(\Delta pH\), compensating the loss of \(V_m\) and restoring growth in the GOF mutants. This work shows that Kch is apparently functional in vivo, in the sense that it can pass K\textsuperscript{+} along its electrochemical gradient and that the membrane potential apparently controls it. If this model is correct, many prokaryotic K\textsuperscript{+} channels, like Kch of E. coli, may function like their counterparts in animal cells. While the work of the E. coli GOF mutants is informative, it begs the question of the stimuli (stresses?) that normally opens the wild-type Kch channel. Extensive attempts fail to register the electric current through the membrane of giant E. coli spheroplasts, including those generated from the GOF mutants. Studies of kch\(\Delta\) have failed to discover a clear Kch LOF phenotype in the laboratory (Kuo, Haynes, Saimi, and Kung, unpublished results). On the other hand, the widespread presence of K\textsuperscript{+}-channel genes in prokaryotic genomes clearly indicate the selective advantages they confer. More creative ways to simulate nature in the laboratory are needed to solve this puzzle.

The E. coli GOF mutant studies suggest, though cannot prove, that K\textsuperscript{+} channel regulates membrane potential, like they do in animal cells. In plant roots, there is evidence that certain K\textsuperscript{+} channels are used to take up bulk K\textsuperscript{+}. Interestingly, the MthK-like K\textsuperscript{+} channel of Helicobacter pylori, called HpKchA, apparently serves such a function. hpkchA\(\Delta\) mutant grows poorly at low K\textsuperscript{+} concentration, but well when K\textsuperscript{+} is supplemented. HpKchA is apparently required to colonize murine stomach (359). However, H. pylori, unlike free-living bacteria, are devoid of other K\textsuperscript{+} uptake mechanisms. Bulk K\textsuperscript{+} uptake is unlikely to play a role for K\textsuperscript{+} channels in free-living bacteria, which invariably have several non-channel K\textsuperscript{+} uptake mechanisms (193). In E. coli, three non-channel K\textsuperscript{+} uptakers are well known (Trk, Kdp, Kup), which transport or scavenge K\textsuperscript{+} when needed. kch\(\Delta\) E. coli cells are not K\textsuperscript{+} auxotrophic (193), and prior search for K\textsuperscript{+} auxotrophs did not identify Kch.
IV. ION CHANNELS OF MICROBIAL EUKARYOTES

Research on ion channels focused largely on channels of vertebrates and larger invertebrates. These studies have greatly contributed to our knowledge of the role that ion channels play in the physiology of muscles and the nervous system. Research on microbial channels, on the other hand, offers a unique opportunity for answering questions on the evolution of ion channels as the microbes display large varieties of channels in their cellular membranes. This is best illustrated by the fact that the structures and biophysical properties of microbial channels exhibit variations not seen in plants and animals. The Paramecium genome, for example, reveals a 12-TM subunit of S1 S2 S3 S4 S5 P1 S6 S7 S8 S9 S10 S11 P2 S12 arrangement (190), whereas yeast has an 8-TM, two-pore-unit of S1 S2 S3 S4 S5 P1 S6 S7 S8 S9 S10 S11 P2 S12 arrangement (157, 413).

Ionic currents have been recorded from a number of microbial cells including Dictyostelium (slime mold) (256), Chlamydomonas (a green flagellate) (355, 399) Paramecium and Stylonychia (ciliates) (65, 67, 128, 262, 311), Neurospora (bread mold) (203, 323), Uromyces (a parasitic bean rust fungus) (412), and single-celled fungi Schizosaccharomyces pombe (fission yeast) (410) and Saccharomyces cerevisiae (baker’s yeast) (114, 115). Although the functions of a number of ion channels in ciliates are well understood and interpreted using concepts developed in animal biology, the roles of most ion channels in eukaryotic microbes still remain to be elucidated. Given that fungi and protozoa are highly diverse and occupy widely different ecological niches, there is a great deal to be learned from studies of ion channels in these microorganisms. The yeast K+ channel may illustrate the point. It is gated by the total K+ electrochemical gradient instead of voltage, and this gate has been thoroughly dissected by genetic and biophysical means (218).

The fact that microbes can be genetically modified with relative ease provides significant advantages for functional studies of ion channels. In S. cerevisiae as a key experimental model gain-of-activity mutants have been isolated, the expression of which hampers growth (220). However, the loss of ion channel activities, as in the case of the knock-out mutants, does not necessarily always lead to a discernable phenotype in the laboratory. The inability to discern a laboratory phenotype indicates that these channels serve functions in conditions not yet simulated in the laboratory. In other words, the channel functions cannot easily be converted into differences in growth rates or cell survival, which are commonly monitored as laboratory microbiological phenotypes. Although we may expect that the basic physical and chemical principles underlying channel activities should be universal, the biological roles of ion channels in the physiology of microbial cells will likely be much broader than the roles they play in the physiology of muscles and the nervous system.

A. Ciliates Paramecium and Stylonychia

In ciliates Paramecium and Stylonychia, ion channels are known to generate receptor potentials and action potentials comparable to those found in cell membranes of neurons or sensory organs in metazoa. The perception and transduction of external stimuli in these microorganisms occur at the unicellular level. Consequently, for studies of membrane excitability ciliates may be perceived as “swimming sensory neurons” (68). Such studies in Paramecium and Stylonychia have demonstrated a crucial role that ion channels play in modulating the behavior of these ciliates by generating electrical membrane activity for signal integration and by controlling ion fluxes across their cell membranes.

Paramecium was the first microbial cell penetrated with a microelectrode (150). Being some 100–300 μm in length, Paramecium can easily be investigated by the classical two-microelectrode intracellular recording. Voltage-clamp experiments revealed that membrane depolarization can trigger a Ca2+-dependent action potential leading to an increase in intracilliary calcium concentration, which causes 5,000–6,000 cilia covering the body of a Paramecium cell to beat in a reversed direction (“avoiding reaction”) (146) to propel the cell backward (85, 262). Membrane hyperpolarization can trigger other ion currents causing the cell to spurt forward due to a more rapid ciliary beat in a normal direction (“escape response”) (262, 264). These experiments helped to dissect the K+, Ca2+, and Na+ currents that constitute various action potentials encountered in a Paramecium cell. In addition, as a model organism for genetic studies, Paramecium, and P. tetraurelia in particular, provided means to isolate behavioral mutants, whose avoiding reactions to a range of stimuli were missing or altered, identify the corresponding electrophysiological changes with intracellular microelectrodes (330), and clone the genes of some of the corresponding mutations (331). Since the entire genome of P. tetraurelia has recently been sequenced, many genes corresponding to K+, Ca2+, and Na+ channels as well as nonspecific cationic or anionic channels could be identified (9).

The freshwater hypotrichous ciliate Stylonychia is another ciliate that was extensively examined by the classical two-microelectrode current- and voltage-clamp methods. Because of a variety of outstanding structural and physiological properties characteristic of this unicellular microorganism Stylonychia has served as a valuable model system for cellular physiology (227). For example, simultaneous recording of responses of compound cilia
(membranelles and cirri) in *Stylonychia* using membrane voltage-clamp and high-frequency cinematography showed that membranelles beat at high frequency, whereas the frontal and marginal cirri are quiescent at the membrane resting potential. In contrast, membrane hyperpolarizations and depolarizations specifically activate the cirri, but leave the membranelar frequency unchanged (69). These results indicate that the activity of both frontal and marginal cirri is coupled to the membrane potential similar to the ciliary activity in *Paramecium*, whereas membranelles should have a mechanism of motor control independent of the membrane potential. Although unicellular, *Stylonychia* thus exhibits complexity characteristic of metazoan organisms. This may not be that surprising given that from the view of ion channel evolution, most of the various types of ion channels and their gating mechanisms had already been operating by the time the ciliates emerged some 200 million years ago (183, 186).

1. **Ca**\(^{2+}\)-dependent channels

At least six types of K\(^+\) currents could be identified in *Paramecium* using a combination of genetics and electrophysiology (311). The recently completed *Paramecium* genome-sequencing project (75), however, provided surprising evidence for the existence of 298 K\(^+\) genes, a number three times bigger than the number of K\(^+\) channel genes found in human genome (190); this is a humbling and potentially disturbing fact for an anthropocentrically minded reader. Based on their open reading frames (ORF), 239 genes of the *Paramecium* K\(^+\) channels encode channels similar to the animal channels gated by cyclic nucleotides (CNG-type). The remaining 57 genes encode channels similar to animal ion channels such as inward rectifier, depolarization or hyperpolarization-activated K\(^+\) channels, as well as Ca\(^{2+}\)-dependent BK or SK type K\(^+\) channels. Combined, *Paramecium* also has some 30 Na\(^+\) and Ca\(^{2+}\)-channel genes, which is approximately the number found in the human genome.

Studies of *Paramecium* mutants made it possible to identify a number of Ca\(^{2+}\)-dependent currents controlling its motile behavior. “Panthophobiacs,” a group of behavioral mutants in *Paramecium*, called so because of their overreaction to a variety of stimuli, helped to show that microinjection of the wild-type cytoplasm was able to restore Ca\(^{2+}\)-dependent outward potassium currents lacking in voltage-clamp experiments (329). By fractionating the wild-type cytoplasm, the curing element was later identified as calmodulin, a 17-kDa soluble protein highly conserved in eukaryotes (331). When microinjected to a pantophobiac cell, calmodulin restored the current and cured the misbehavior.

Another group of *Paramecium* mutants called “fast-2” have an opposite phenotype to “panthophobiacs” due to a lack of a Ca\(^{2+}\)-dependent Na\(^+\) inward current. Fast-2 mutants were also identified as having defects in their calmodulin (159). Interestingly, studies of “panthophobiacs” and “fast-2” mutants indicated that Ca\(^{2+}\)-dependent K\(^+\) channels are regulated by the C lobe, whereas Ca\(^{2+}\)-dependent Na\(^+\) channels interact with the N lobe of the calmodulin molecule. This finding was later confirmed in patch-clamp experiments showing that the *Paramecium* Na\(^+\) channel could be directly activated by the Ca\(^{2+}\)-calmodulin complex acting as a detachable subunit (332). Detailed information on regulation of Ca\(^{2+}\)-dependent currents in *Paramecium* by calmodulin can be found in a review by Preston et al. (311). In contrast to the multitude of ion channel genes, there is only a single calmodulin gene in the *Paramecium* genome.

Other Ca\(^{2+}\)-dependent channels found in a *Paramecium* cell include a Ca\(^{2+}\)-dependent Mg\(^{2+}\)-specific current and a hyperpolarization-activated Ca\(^{2+}\)-dependent K\(^+\) channel. Mg\(^{2+}\)-specific current is activated by membrane depolarization as well as by membrane hyperpolarization (310). Divalent ions such as Co\(^{2+}\) and Mn\(^{2+}\) and, to a lesser extent also Sr\(^{2+}\) and Ba\(^{2+}\), can substitute for Mg\(^{2+}\) as charge carriers through the Mg\(^{2+}\) channels. A physiological role of the Mg\(^{2+}\)-specific current consists most likely in maintaining intracellular free Mg\(^{2+}\) at low concentrations (0.1–0.7 mM). This current was found to be affected in some of the calmodulin mutants. Single gene mutants of *P. tetraurelia* with a reduced ability to respond behaviorally to Mg\(^{2+}\) were isolated and called “eccentrics” (*xntA* and *xntB*). Voltage-clamp experiments show a deficit of the Ca\(^{2+}\)-dependent Mg\(^{2+}\) current in these mutants (312, 313). Because the macronucleus of *Paramecium* is large, DNA can be directly injected through a micropipette into a mutant macronucleus to effect phenotypic transformation. It is possible to clone certain genes through such complementation by sorting a library of wild-type genomic DNA fragments to be injected. With the use of this method, the XntA gene was cloned and its predicted product found to be unlike any known ion channels but similar to animal Na\(^+-/Ca\(^{2+}\) exchanger (126). This is the first example of an exchanger-like entity behaving like an ion channel electrophysiologically. This surprising finding is recently echoed in the study of CIC-ec1, the bacterial chloride “channel” behaving like an exchanger (see sect. nD).

A hyperpolarization-activated Ca\(^{2+}\)-dependent K\(^+\) channel with a conductance of ~70 pS was recorded from the *Paramecium* plasma membrane blisters using the patch-clamp technique (333) (Fig. 2). This channel is weakly voltage dependent, requiring ~41 mV-e-fold change in its activity. A treatment of the channel with trypsin, pronase, or thermolysin from the intracellular side in inside-out excised plasma membrane blister patches removed the Ca\(^{2+}\) dependence of the channel activation, which suggested that this K\(^+\) channel has a cytoplasmic inhibitory domain that can be removed by...
proteolysis (180). The physiological activation of this channel by Ca\(^{2+}\) thus could be due to a temporary removal of this inhibition. Hyperpolarization-activated Ca\(^{2+}\)-dependent K\(^{+}\) currents have also been identified and characterized by two-microelectrode voltage-clamp experiments. Since these currents are thought to regulate the resting membrane potential in *Paramecium* (322), the 70-pS hyperpolarization-activated Ca\(^{2+}\)-dependent K\(^{+}\) channel may correspond to one of these currents.

2. Mechanosensitive channels

When observed under a low power microscope, *Paramecium* cells transiently stop or back up briefly by reversing their ciliary beat direction (ciliary reversal). These avoiding reactions (146) occur spontaneously and can also be induced by a variety of stimuli. Mechanical impact at the anterior end of a *Paramecium* cell elicits a Ca\(^{2+}\)-based receptor potential (85). Other divalent cations are also able to carry this membrane-depolarizing current (281), which can, in turn, elicit a Ca\(^{2+}\)-based action potential underlying the ciliary reversal. Mechanical impact at the posterior end of the cell elicits a K\(^{+}\)-based hyperpolarization (85), prompting the cell to spurt fast forward due to an increase in frequency of the ciliary beat in normal direction. Since cilia in *Paramecium* can be removed with ethanol, the experiments on deciliated cells that were mechanically stimulated showed that deciliated cells retained the depolarizing and hyperpolarizing mechanosensitive responses, which indicated that mechanosensitive Ca\(^{2+}\) and K\(^{+}\) channels were localized on the cell soma (228). In addition, ciliated and deciliated cells showed a decrease in depolarizing mechanosensitivity and a subsequent increase in hyperpolarizing mechanosensitivity, when the site of stimulation was shifted from the anterior to the posterior end of the cell. These experiments indicated that the Ca\(^{2+}\) and K\(^{+}\) mechanosensitive channels were distributed over the somatic cell surface in the manner of overlapping gradients (281). Although both mechanosensitive macroscopic currents in *Paramecium* have been well characterized by the voltage-clamp method, the genes and proteins behind these receptor potentials are yet to be identified.

3. Voltage-dependent Ca\(^{2+}\) channels

The composition of ionic currents underlying the Ca\(^{2+}\) action potential and its relation to ciliary reversal in *Paramecium* are well understood and have been extensively described over the years (85, 226, 250, 337). A depolarization-activated current is carried by Ca\(^{2+}\) through voltage-dependent calcium channels, which inactivate in a Ca\(^{2+}\)-dependent manner (39, 280). The channels are located in the ciliary membrane, since deciliation abolishes the depolarization-activated Ca\(^{2+}\) current (228). This current is abolished or reduced in “pawn” mutants that show no backward swimming in response to stimulation (185, 280). Microinjection of cytoplasm from a wild-type cell into the pawn mutants belonging to any of the three complementation groups (pawnA, pawnB, pawnC) restored the ability of the mutants to generate action potentials and hence swam backward (116). In addition, the cytoplasm from a pawn of a different complementation group as the microinjected mutant could also restore the action potential and backward swimming. Consecutive studies demonstrated further that microinjection of the total wild-type genomic DNA into the macronucleus of a *Paramecium* cell could restore the Ca\(^{2+}\) current and excitability in clonal descendants of the microinjected pawnA cell. By sorting a library of total DNA, the factor required for restoring the Ca\(^{2+}\) current was identified as a 2.3-kb DNA fragment corresponding to a novel protein that may be glyco phosphatidylinositol anchored (129). The same method was used to clone the pawnB gene, which corresponds to a peptide with two transmembrane domains but no clear homology to known proteins (127).

Similar to *Paramecium*, mechanical stimulation of the anterior part of *Stylonychia* evokes a depolarizing mechanoreceptor potential, which gives rise to an action potential. The action potential in *Stylonychia* is, however, more complex compared with the *Paramecium* action potential. It has two components, which are represented by a first, large graded peak and a second, small “all-or-none” peak (65). The two components were shown to correspond to two types of Ca\(^{2+}\) currents based on their differential sensitivity to Co\(^{2+}\) and Cd\(^{2+}\) (67). In both types of currents Ca\(^{2+}\) can be replaced by Ba\(^{2+}\) or Sr\(^{2+}\). These Ca\(^{2+}\) currents are separable by their localization in the membrane. The smaller Ca\(^{2+}\) current corresponding to the “all-or-none” peak is restricted to the membranelles (row of compound cilia) since in cells that spontaneously released their membranellar band the small current disappeared and the larger current corresponding to the graded peak remained unaltered. It has been suggested that the two voltage-dependent Ca\(^{2+}\) currents may be functionally related to different beating behavior of membranelles and other types of the compound cilia in *Stylonychia* (67).

B. Slime Mold (*Dictyostelium*)

1. A model system of chemotaxis without ion channel function

*Dictyostelium discoideum* cycles between free-living single amoebae and social multicellular organisms for spore formation. After starvation, the amoeba secrete cAMP, and cAMP-chemoreception drives the amoebae to form a multicellular slug (chemotaxis), which, in turn, moves around to find an appropriate site to form fruiting
body to develop spores. This chemotaxis is the central study of *Dictyostelium*, since it shares molecular mechanisms with the chemotaxies of neutrophils, lymphocytes, and even cancer cells (46, 100, 288, 309, 391). Briefly, upon reception of external cAMP by G protein-coupled receptor proteins (cAR1 and cAR3), a series of reactions including Ga2βγ activation, small GTPase (Ras and Rho family members) activation (45), PI3K (class 1 phosphatidylinositol-3 kinases, IP3K, -2, -3) activation (104, 143, 373), phosphorylation of PI(3,4,5)P3 into PI(3,4)P2 and finally F-actin polymerization resulting in pseudopod development for chemotactic movement. Activation of Rho GTPases or their functional homologs, on the other hand, leads to activation of PTEN (PI 3-phosphatase) at the opposite site of the signal source that causes a reduction of PI(3,4,5)P3 (45). Thus, through changing the local concentrations of PI(3,4,5)P3 at the front and back, the amoebae amplify the external cAMP signal (as little as 2% difference) and convert into internal signal gradient to determine for correct polarity in chemotaxis. Although the main cascade of signal transduction has been so established, the chemotactic orientation of *Dictyostelium* cells is hardly affected after genetic deletions of PI3Ks and reduction in PI(3,4,5)P3 contents (213, 373), implying redundant pathways. Genetic screens have identified a second pathway utilizing PLA2A (Patatin-like phospholipase A2 homolog; Ref. 48). Arachidonic acid has been implicated in the PLA2A pathway (48). There is yet another pathway for chemosensation in *Dictyostelium* that uses phospholipase C (PLC) (156, 174, 381).

Although chemosensation in *Dictyostelium* is often associated with Ca2+ fluxes (196, 225, 268, 338) and PLA2A may be regulated by intracellular Ca2+ (381), there is no evidence that Ca2+ signals are involved in chemotaxis (374), nor is there evidence that membrane potential plays a role in regulation of chemotaxis (380). This lack of ion channel involvement in chemotaxis is in stark contrast to the axonal growth cone that also senses chemical cues and requires TRP channel function (386, 389). The genome of *D. discoideum* has been completely sequenced (88), and one can identify at least two possible TRP channel genes, a Ca2+ channel gene, and several K+ channel genes. Its membrane potential is largely regulated by an electrogenic proton pump like fungi (379). Curiously, there is no recognizable gene for CNG channels in this organism, which is the champion of cAMP chemotaxis. Nuclear pores in nucleus membranes are obviously critical for all eukaryotic cells. Recent efforts on *Dictyostelium* nuclear pores show new features of the pore complex (15). An amoebapore (protozoan pores that affect humans) is also found in the database (405). As to channel activities, three single-channel activities have been reported from the results of patch clamp in the cell-attached mode; DI (11pS), DII (6pS), HI (3 pS) (256, 257). DI and DII are permeable to K+, and HI may be permeable to Ca2+. DII is quite low in density.

2. P2X, ATP-activated channel

P2Xs are purinergic receptor channels found and studied extensively in animals. These are channels that open upon the binding of extracellular ATP. The channels are multisubunits of subunits, each with two transmembrane helices (277). *Dictyostelium* has a gene, which has weak homology to these P2X channels. Fountain et al. (99) expressed this gene (DsP2X) in HEK cells, where it apparently forms trimeric ATP-gated channels of ~8 pS conductance with some Ca2+ selectivity. Permeability to Na+ is lower (PNa/PNa = 1.5) and those for larger organic cations still lower. The DdP2Xs have key residues for ATP binding, such as Lys67 (Lys71 in rat P2X) and Lys289 (Lys308 in rat P2X), which have critical functions in ATP sensitivity. Unlike mammalian P2X, DdP2X is not activated by ATP analogs. Cu2+ blocks DdP2X channels at micromolar concentrations, but not mammalian P2Xs. This gene product in *Dictyostelium*, unlike the HEK cell expression, is localized to the contractile vacuole, the water pumping organelle. Upon osmotic downshock, the DdP2X knockout *Dictyostelium* suffers swelling. Also, HEK cells expressing DdP2X swell upon inhibition by Cu2+. One possible hypothesis is that distension of the contractile vacuole leads to the entry of ATP from the cell cytoplasm, which activates DdP2X receptors, allowing vacuolar Ca2+ to pass back into the cytoplasm and initiate contraction (99). Curiously, *Dictyostelium* contractile vacuole apparently includes a voltage- and K+-dependent K+ channel, which may also function in water regulation (398).

C. Yeasts and Other Fungi

Much of the current knowledge on fungal channels came from the study of the budding yeast *Saccharomyces cerevisiae*. We will describe the various ion channels studied therein and then point out their homologs in other fungi.

1. Mechanosensitive channels

Like bacteria, yeast cells have to face rapid and large changes in osmolarity. Given the knowledge on prokaryotic mechanosensitive channels, it is not surprising that yeasts and other fungi also have such channels. Possible fungal homologs of MscL and MscS are listed in section II.A.

a) Plasma membrane mechanosensitive channel. A patch-clamp survey in 1988 of the yeast plasma membrane revealed an ion-nonspecific stretch-activated channel of 36 pS (115). This conductance has poor selectivity, al-
though preferring cations over anions. The \textit{MID1} product is required for Ca\textsuperscript{2+} influx in yeast (see sect. \textit{S. cerevisiae}). When expressed in mammalian cells, \textit{MID1} was reported to support a pressure-activated, Gd\textsuperscript{3+}-suppressible 32-pS conductance (153, 154). However, the plasma membranes of \textit{mid1Δ} yeast cells retain the 36-pS stretch-activated channels (187). Despite the availability of the complete \textit{S. cerevisiae} genome sequence, the molecular identity of this channel remains unknown to date. At this writing, each ORF that corresponds to recognizable channel sequence in this genome has been individually deleted. None of the deletions remove this 36-pS conductance from the plasma membrane (Zhou, Saimi, and Kung, unpublished results). It, therefore, is likely that it corresponds to a novel channel structure. The surface of protoplast protuberances of the fission yeast \textit{Schizosaccharomyces pombe} has also been examined with the patch clamp (410). There is a 180-pS conductance that prefers K\textsuperscript{+} over Cl\textsuperscript{-}, but discriminates poorly among cations. The spores of the bean-rust fungus \textit{Uromyces appendiculatus} germinate, and the growing germ tubes are apparently guided by the topography of bean-leaf surface to the stomata, where they differentiate into invading structures. A 600-pS mechanosensitive conductance can be registered from the surface of protoplasts derived from the germlings (412). This conductance is cation specific, nearly impermeable to Cl\textsuperscript{-}. It passes Ca\textsuperscript{2+} and is blocked by Gd\textsuperscript{3+}. It seems possible that this channel can transduce the membrane stress induced by the leaf topography into a flux of Ca\textsuperscript{2+} that initiates differentiation.

\textbf{2) TRP CHANNELS (Yvc1).} First identified from a near-blind \textit{Drosophila} with an abbreviated transient receptor potential, TRP channel is now a large superfamily of channels largely studied in animals. Attention is drawn to them in recent years because they apparently underlie sensation of certain tastes, of hot and cold (and the chemical mimics such as pepper and menthol), and of force (osmotic, impact, tension, vibration, etc.). Each TRP channel is believed to be a tetramer of 6-TM subunits akin to \textit{Shaker} K\textsuperscript{+} channel (Fig. 5). The seven subfamilies of animal TRPs, however, are quite diverse, with sequence similarity only at TM5-P-TM6 segment and just beyond it (52). The yeast genome includes a gene containing this sequence, and its product forms a channel in the vacuolar membrane. It is called Yvc1, for yeast vacuolar channel, or TRPY1, for a TRP channel in yeast (286). Yeast cell wall can be removed by enzyme digestion; its plasma membrane can be removed by osmotic manipulation. The exposed vacuole can directly be examined with a patch-clamp pipette. Yvc1 displays an inwardly rectifying conductance of 350 pS, rather large for TRP channels. This conductance was first discovered by reconstituting vacuolar membranes into planar lipid bilayers (388). Like other TRP channels, Yvc1 passes cations but discriminates poorly among them, and its open probability is regulated by H\textsuperscript{+} and Ca\textsuperscript{2+} (21, 286). Using transgenic aequorin as an indicator, Denise and Cyert (72) found that Yvc1 releases Ca\textsuperscript{2+} from the vacuole into the cytoplasm upon osmotic upshock. Correspondingly, Yvc1 was found to respond directly to membrane stretch force under patch clamp (409). Combining forward genetics and patch clamp, Yvc1 is being dissected towards understanding its detailed structure-function relationship and the submolecular bases of regulation by force and by Ca\textsuperscript{2+}. GOF mutations were found to be mostly those on aromatic amino acids. A mutation at the base of TM6 near the presumed gate was found to destabilize both the closed and the open state. Of all the 19 variations at this residue, only the aromatics can apparently anchor the gate properly (408). Another aromatic mutation at the cytoplasmic end of TM5 displays a constitutively open channel kinetics, phenocopying a special allele of the \textit{Drosophila} TRP channel (363). Yvc1 has direct homologs in over 30 species of fungi except \textit{Schizosaccharomyces pombe} and the parasitic \textit{Zygomycetes}. The ones from \textit{Kluyveromyces lactis} and \textit{Candida albicans} have been expressed in the vacuole of \textit{S. cerevisiae} and found to also function as mechanosensitive cation channels (411). Yvc1 (TRPY1) and its direct relative are considered more related to TRPML of animal TRP channels (51). A gene in the fission yeast \textit{S. pombe} is reported to encode a channel more similar to animal TRPP (285), although it appears to have minimal homology.

\textbf{2. K\textsuperscript{+} channels (Tok1)}

The activities of a K\textsuperscript{+} channel were evident in the first patch-clamp examination of the yeast plasma membrane in 1986 (114). The yeast genome was the first eu- karyotic genome to be completely sequenced in 1996. Four groups independently recognized a gene that predicted a K\textsuperscript{+} channel (157, 201, 320, 413). The cloned gene can be expressed and examined in yeast itself or in \textit{Xenopus} oocytes. Unlike the 2-TM KcsA or 6-TM \textit{Shaker} motif, the channel subunit predicts a peptide with 8 TM helices and 2 P domains (Fig. 5D) and is named TOK1, for two-pore domain K\textsuperscript{+} channel (157). It is K\textsuperscript{+}-specific, rectifies outwardly, and has a unitary conductance of \~30 pS (413). Although TOK1 may take up K\textsuperscript{+} in certain mutant context (94), the complete loss of TOK1 function in \textit{tok1Δ} cells has no easily discernible growth phenotype in the laboratory. GOF mutants, however, can be isolated that stop growth when the mutant channel is expressed. Examined in oocytes, such mutant channels are largely unable to enter a closed state distal to the open state. Many mutations are located at the "post-pore" regions, at the cytoplasmic end of the predicted TM6 and TM8 (220). Both are the inner helices comparable to TM2 of KcsA; the "post-pore" region is now shown to be near the K\textsuperscript{+}-channel gate in crystal structure (80). The proximal
Ca\textsuperscript{2+} products, Cch1 and Mid1, are needed for this Ca\textsuperscript{2+} endoplasmic reticulum stress (34). The entry of animals (209) and play roles in the response to static mechanism analogous to the capacitative calcium located in the plasma membrane, where they apparently voltage-gated Ca\textsuperscript{2+} residue glycoprotein, which has no homologs in animals Mid1 (for mating-pheromone-induced death) is a 548-residue protein, which has homologs in animals 

3. Ca\textsuperscript{2+} channels (Cch1)

Animal Ca\textsuperscript{2+} channels are large proteins of over 2,000 molecular weight. Besides the flaming cytoplasmic domains, the transmembrane body has four repeats, each has a 6-TM arrangement similar to the one described for K\textsuperscript{+} channels above. The yeast genome sequence revealed a Ca\textsuperscript{2+} channel, but unlike its animal counterparts, there has been no report on electric current through this channel in situ or in heterologous cells. No Ca\textsuperscript{2+} currents have yet been reported in the plasma or the vacuolar membranes with patch-clamp electrodes at different voltages or stretch force. The studies on this channel are therefore genetic and not electrophysiological.

Mating pheromones induce a Ca\textsuperscript{2+} influx. Two gene products, Cch1 and Mid1, are needed for this Ca\textsuperscript{2+} influx and for the survival through the mating process (95, 284). Mid1 (for mating-pheromone-induced death) is a 548-residue glycoprotein, which has no homologs in animals (140). Cch1 (for calcium channel) is a homolog of animal voltage-gated Ca\textsuperscript{2+} channel (141). Mid1 and Cch1 are both located in the plasma membrane, where they apparently function together and can be communoprecipitated (209). Evidences suggest that they are parts of a homeostatic mechanism analogous to the capacitative calcium entry of animals (209) and play roles in the response to endoplasmic reticulum stress (34). The cch1\textDelta mid1\textDelta double mutant is apparently sensitive to cold and to high iron concentration in the medium (296). The osmotic upshock-induced cytoplasmic Ca\textsuperscript{2+} response is reduced in the cch1\textDelta mid1\textDelta double mutant (241). However, a recent examination of a collection of 4,810 yeast-gene deletants shows that this reduction is due to the depletion of Ca\textsuperscript{2+} in the vacuole and not due to defects in the Ca\textsuperscript{2+}-releasing channel Yvc1 (S.H. Loukin, M.M. Kuo, X.L. Zhou, W.J. Haynes, C. Kung, Y. Saimi, unpublished data). Alkaline shock induces a Ca\textsuperscript{2+} flux as evidenced by Ca\textsuperscript{2+}-aequorin luminescence. This flux is largely missing in cch1\textDelta or mid1\textDelta, suggesting a role of Ca\textsuperscript{2+} channel in defense against alkaline stress (384). Even though Cch1 cannot yet be studied directly with electrodes, insights from the dissection of the yeast Cch1 can be transferred to animal Ca\textsuperscript{2+} channels. For example, LOF phenotypes were found in a cch1 mutant, mutated at a glycine that is completely conserved. The corresponding mutant in an animal Ca\textsuperscript{2+} channel is then found also to be nonfunctional under voltage clamp (142).

4. NH\textsubscript{4}\textsuperscript{+} channels (Mep)

Fungal Meps (methylamine permeases) are homologous to Amts (ammonium transporters) in the bacteria (see sect. iiiC). The first NH\textsubscript{4}\textsuperscript{+} channel was cloned as a transporter that complements the yeast mep1-mutant defect of poor growth with ammonium as the sole nitrogen source (232). There are three such transporters in yeast, Mep1, Mep2, and Mep3, of different affinity and capacity, as measured by \([^{14}C]\)methylammonium accumulation into live yeast cells (231). Biochemical studies showed that the NH\textsubscript{4} terminal of Mep2 faces outward and is N-glycosylated (230), and its function requires the serine/threonine kinase Npr1 (33). Mep2 apparently also functions as a sensor for rapid activation of the protein kinase A pathway (382). Nitrogen starvation of diploid yeast on a fermentable carbon source triggers pseudoephedrine growth, presumably to better explore the environment. Mutant of mep2, but not mep1 or mep3, abolishes pseudoephedrine growth, leading to the suggestion that Mep2 may be the sensor (215). There appears to be no study on Meps in more reduced system such as oocytes or liposomes, and no discussion on interpreting the substrate-accumulation data with the presumed channel mechanism.

5. Cl\textsuperscript{-} channels (Gef1)

Mutations of a gene called GEF1 yield the petite phenotype that can be suppressed by added iron. GEF1 was found to give a product of 779-residue, 13-TM structure, similar to the animal Cl\textsuperscript{-} channel subunits (110) (see sect. iiD). This channel, also called yClC-1 (138), was first found to be located in the Golgi apparatus (343) or pre-vacuolar fraction. It was found to be a necessary component in the homeostasis of Cu\textsuperscript{2+} and other cations in vesicular compartments by regulating their pH. gef1 defects can be complemented with certain animal or plant CICs, indicating functional similarity (105). Gef1 is apparently proteolytically cleaved in the secretory pathway into
a 30- and a 70-kDa fragment, which function together but not individually (387). Electrophysiologically, a 10-pS anion conductance was observed when microsomal fraction of the wild-type, but not the gef1 mutant, was reconstituted into planar lipid bilayers (97). There is a recent report showing that GEF1 can be expressed in oocytes or HEK cells and apparently producing a permanently open channel in the plasma membrane, blockable by known ClC blockers. In that study, electron microscopy showed that GeF1 is located in the plasma membrane as well as internal membranes (214). GEF1 homologs can be found in other yeast species as well as in filamentous fungi. Two are found in the human pathogenic yeast Cryptococcus neoformans, one of which is required for virulence (415). Three such homologs, AnCLCA, -B, and -C, are found in the filamentous Aspergillus nidulans (279).

D. Algae (Chlamydomonas)

Green algae comprise single cells or loose colonies of cells. They are true plants with chloroplasts for photosynthesis. Some of them have flagella or cilia to move about, seeking light (phototaxis) and/or avoiding intense light (photophobic response, also called photoshock response) (310, 349, 392). Chlamydomonas reinhardtii has been exploited as a model system in the studies of photosynthesis and cell motility for a number of years. Recently, its entire genome has been sequenced (245) along with another nonflagellated yet free-living green alga Ostreococcus tauri (74). Sixty-one ion channel genes have been predicted in Chlamydomonas (245). However, patch-clamp analyses of channel activities have proven to be difficult because of low resistance seal with patch pipettes.

In Chlamydomonas, light is detected with the eye spot, which contains rhodopsins. These rhodopsins consist of protein opsins and photopigment retinals, but unlike mammalian rhodopsins (type 2), they mostly consist of all-trans retinals, similar to prokaryotic rhodopsins (type 1). These types of rhodopsins have similar structural designs (7 transmembrane domains) but are dissimilar in terms of amino acid contents. In fact, the opsins of Chlamydomonas are numerous and mixed. At least six opsin genes have been identified in the genome. Light sensation by the eye spot in Chlamydomonas causes membrane depolarization, which in turn activates the Ca^{2+} channels in flagella (14, 121, 135, 136, 349). Ca^{2+} in flagella is known to regulate the flagella motion. At low Ca^{2+}, the flagella movement is in the form of breast stroke (or ciliary type motion), whereas at higher Ca^{2+}, it becomes undulating (or flagellar type). Because the two flagella have different Ca^{2+} sensitivity (152), the net translational movement of the cell can be rotational when the light stimulation is low, resulting in turning the cell body until it orients towards the light source, thereby shielding the eye spot from further excitation. At a high light intensity, both flagella beat in the flagellar type motion, thus moving the cell away from the light source (349).

1. TRP homologs

The general features of TRP channels are given in section vClC (318). They are found among mammals, worms, insects, yeasts, ciliates, but, mysteriously, largely absent in plants. One exception is Chlamydomonas; there are at least seven homologs of TRP-channel genes in its genome. One of them is a homolog of PKD2 or TRPP2, a gene whose mutant forms cause a polycystin disease in kidney (255). In the kidney, the NH_{2} terminus of PKD2 interacts with the COOH terminus of PKD1, a protein with multiple transmembrane domains, to form a functional channel unit, which lets Ca^{2+} into the cell (120). These channels reside in the ciliary membrane and probably are activated by mechanical movement of cilia (120, 267). Interestingly, PKD1’s NH_{2} terminus may be cleaved at GPS domain (G protein-coupled receptor proteolytic site) (315, 403), which may be important for the function of the PKD2-PKD1 complex in the ciliary membrane. Thus PKD2 is the pore-forming subunit, whereas PKD1 may have multiple functions such as chaperone and G protein activation (70). PKD1 does have modest homology to PDK2. Thus PKD1 alone might also form channels (175). Cilia or flagella are assembled and maintained by the intraflagellar transport (IFT), motorized with kinesin-2 and dynein 1b or 2 (326). IFT particles comprise ~17 proteins that are conserved among mammals, fish, flies, and worms (10, 204). In Chlamydomonas, one of the IFT subunits has been identified as a PKD2 homolog (295). Chlamydomonas PKD2, CrPKD2, is a protein of 1,626 amino acids with 6 or even more transmembrane domains. Its NH_{2} terminus extracellular domain is ~1,000 amino acids long (137). The whole cell extracts show 210-, 120-, and 90-kDa proteins, recognizable by antibodies against CrPKD2. The latter two are the in vivo proteolytic products of the 210-kDa protein after the long extracellular domain, and only these two proteins are found in the flagellar fraction, while the 210-kDa protein is not detected in flagella. CrPKD2 can be visualized with a GFP tag to be in puncta that move along the flagella (137). This movement is dependent on IFT as such fluorescence movement is not observed in the fla10 mutant under a restrictive condition. RNAi knockdown of CrPKD2 causes reduced mating reactions, suggesting CrPKD2 may be necessary for Ca^{2+} influx after flagellar adhesion in the cascade of mating reaction in Chlamydomonas (137). Whether CrPKD2 forms the functional channel by itself or with other subunits such as PKD1 in Chlamydomonas has yet to be determined. Also, which product(s) of CrPKD2 (the 210 kDa found in the cell body or 120/90 kDa proteins...
found in the flagella) actually function as ion channels has to be determined in the future. Some of the other TRP channels may work together with CrPKD2 in the function of Ca\textsuperscript{2+} signaling during the mating reaction. It would also be interesting to see whether the *Chlamydomonas* CrPKD2 has the purported mechanosensitivity of PKD2.

2. Channel rhodopsins

In the study of phototaxis and photophobic responses of *Chlamydomonas*, at least seven eye pigment proteins have been identified. *Cop1* and *Cop2* (splicing variants) (66, 102) are similar to the invertebrate opsins and *Cop4* (also called channelrhodopsin) (ChR) 1, 2 (260, 261); CSRA, B (350); *Acop1*, 2 (371) are similar to prokaryotic (type 1) rhodopsins that are H\textsuperscript{+} pumps utilizing light energy. Many of the key residues necessary for prokaryotic rhodopsin’s H\textsuperscript{+} pump activity are preserved in ChRs (350). There are other opsins (Cop5–7) similar to prokaryotic “sensory” rhodopsins (356) in the *Chlamydomonas* genome (155). Bacterial and archaenal rhodopsins are type 1 rhodopsins and can be classified into two categories: transporters which pump H\textsuperscript{+} or Cl\textsuperscript{−} and electrogenic [see Stoekenius (360) for a historic account] and receptors (SRI and SRII) which interact with their cognate transducers HtrI or HtrII to activate the cascade of Che gene reactions in phototaxis (356). Such type 1 rhodopsins are also found in eukaryotes (345). The structures of some of these rhodopsins have been solved at atomic resolution.

A surprising finding on ChR1, 2 in *Chlamydomonas* is that these ChRs behave more like ion channels than pumps. ChRs are photosensitive, having their peak absorption at 510 and 470 nm, respectively. Both ChRs are expressed in *Chlamydomonas* and can be expressed in *Xenopus* oocytes as well as mammalian cultured cells and yeast cells (12, 260, 261, 350). Patch-clamp examinations revealed that ChR1 are highly selective for H\textsuperscript{+} (260). On the other hand, ChR2 is selective for mostly H\textsuperscript{+} but also other cations including Ca\textsuperscript{2+} (261). Unlike prokaryotic pump rhodopsins, both currents passed through both ChRs can reverse directions and are dependent on the electromotive force of H\textsuperscript{+} or other cations (259–261), indicating that they behave more like ion channels. However, the estimated conductance appears very small (<300 fS) (155). The response time of ChR2 takes less than 200 μs (261), while that of ChR1 is slow (260, 261).

3. Ca\textsuperscript{2+} channels

Depolarization of the *Chlamydomonas* membrane activates voltage-dependent ion channels. There are about 10 Ca\textsuperscript{2+} channel genes and more than 5 K\textsuperscript{+} channel genes identifiable in the *Chlamydomonas* genome. Changes in internal Ca\textsuperscript{2+} concentrations have been demonstrated to alter flagellar beat patterns (339). For phototaxis, which results from high intraflagellar Ca\textsuperscript{2+} concentrations, Ca\textsuperscript{2+} channels are suspected to be involved, because Ca\textsuperscript{2+} channel blocker ω-conotoxin blocks phototaxis (131). Such channel activity can be recorded with a macropipette covering part of *Chlamydomonas* cell, where light-induced eye-spot current and flagella-origin current can be measured (14). Partial amputation of flagella causes loss of the current of flagellar origin, which recovers as flagella regenerate to their original length, indicating such channels are distributed along the length of flagella (14). Ca\textsuperscript{2+} channels may also be involved in other aspects of cell biology in *Chlamydomonas*. Deflagellation of *Chlamydomonas* can be induced by Ca\textsuperscript{2+} influx triggered by cytoplasm acidification or Ca\textsuperscript{2+} release from internal storage (317). Such acid-induced Ca\textsuperscript{2+} influx is not sensitive to common Ca\textsuperscript{2+} channel blockers, Cd\textsuperscript{2+}, SKF-96365, or flufenamic acid (316). Most interestingly, isolated flagella do not accumulate Ca\textsuperscript{2+} upon acid stimulation, whereas the cell bodies do, indicating the entity for Ca\textsuperscript{2+} influx is not in the flagellar membrane (316), which is different from the Ca\textsuperscript{2+} channel distribution (see above). Currently, it is not clear whether that entity is an ion channel or transporter.

4. Mechanosensitive channels

In *Chlamydomonas*, there is no reported behavior that is associated with mechanical stimulation or osmotic stress, though in a related green alga *Spermatozopsis*, a touch-induced backward movement has been reported (177). However, there are reports on negative pressure-induced mechanosensitive ion channels in *Chlamydomonas* (399), although low seal resistance of the recording
system makes the channel activities less certain. Given the CrPKD2’s presence in the flagella and its possible mechanical activation (see above), mechanically sensitive ion channels may indeed exist in the flagella and/or the cell body of *Chlamydomonas*. Note that there are other TRP channels identified in the genome, which could also play a role in the potential mechanosensitivity.

Another mechanosensitive ion channel is MSC1 (266) that is a homolog of prokaryotic MscS. (see sect. 1.A4). MSC1 has three putative transmembrane domains, and high homology to bacterial MscS can be found in TM3. When expressed in *E. coli*, it corresponds to a ~400 pS conductance, though its NH₂-terminal signal sequence has to be removed prior to expression. MSC1 is activated by membrane tension but not by membrane potential changes. It requires moderate tension, ~80% of tension that activates MscL of *E. coli*. The channel activity diminishes upon hyperpolarization despite continued membrane tension, whereas the activity is maintained upon depolarization. The channel prefers anions over cations (\(P_{K^+}:P_{Cl^-}:P_{H^+}:P_I = 1.7:9:9\)), whereas *E. coli*’s MscS is less selective (\(P_{Cl^-}:P_{K} = 1.5-3.0:1\)) (238, 354, 364, 368).

The most astonishing feature of MSC1 expressed in *E. coli* is its hysteresis behavior depending on how tension is applied to the membrane. It is not clear why MSC1 exhibits hysteresis. In *Chlamydomonas*, it is localized in chloroplasts. Its homologs of *Arabidopsis* (MSL2, 3) are also known to be localized in chloroplasts and are necessary for maintenance of chloroplast morphology (125). The immunolocalization signals are also found in cytoplasm of *Chlamydomonas*, which appears to be fortuitous cross-reactions (266). RNAi knockdown of MSC1 reduces chlorophyll autofluorescence. In *Chlamydomonas*, there are at least two additional homologs, but these have yet to be investigated. Given the mechanosensitive function of the *Arabidopsis*’ MSL3 in rescuing of the mutant *E. coli* lacking mechanosensitive channels upon osmotic downshock (125), it is likely that these eukaryotic MscS homologs also function as mechanosensors in vivo. Since these channel proteins are functionally expressed in *E. coli*, it raises a possibility of purification and reconstitution in artificial lipid membrane for further biophysical studies. It also makes it possible that large quantities of these proteins can be used for crystallization in the future.

### V. BACTERIAL TOXINS FORMING ION CHANNELS

Bacterial toxins are widely distributed proteins that usually induce an unrestricted flux of ions and molecules across membranes of mammalian cells (106). They are frequently cytotoxic as they perforate the cell membranes by creating unregulated pores across the membrane of susceptible cells, leading to cell death and lysis. Broadly classified into two groups, they are thought to interact with the membrane either through α-helices or β-sheet structures such as β-barrels. The mechanism of pore formation is much better characterized for those toxins, which interact with membranes of target cells through β-sheet structures.

The most widespread bacterial toxin encountered by the human organism is α-hemolysin secreted by *Staphylococcus aureus* (24). The α-hemolysin toxicity is a result of its capacity to form large pores permeant to ions, ATP, and water, leading to irreversible osmotic swelling and cell lysis. α-Hemolysin assembles as a homohexamer from a water-soluble monomer to form a membrane-bound β-barrel on the surface of targeted cells. The mechanism of heptamer assembly occurs in a stepwise manner, resulting in a water-filled transmembrane channel having a pore of ~40 Å in diameter (252). The three-dimensional structure of α-hemolysin has been solved by X-ray crystallography to 1.9 Å resolution (352), showing the mushroom-shaped heptamer consisting of a 14-strand antiparallel β-barrel. Ion conductance and anionic selectivity of α-hemolysin is sensitive to pH (176). Furthermore, in the presence of divalent and trivalent cations, the α-hemolysin channel can be inactivated by voltage in a dose-dependent manner (244). In very good agreement with the experiments, molecular dynamics simulation studies of α-hemolysin helped to visualize permeation of ions, small solutes, and water through this channel at atomic scale (372).

RTX toxins (repeats in the structural toxin) present another type of pore-forming bacterial peptides with a molecular mass of 100–200 kDa. They are produced by a broad range of pathogenic gram-negative bacteria and, in particular, species of the family *Pasteurellaceae*. RTX toxins mostly exhibit a cytotoxic as well as a hemolytic activity. They were named after a repeated structure of iterative glycine-rich nonapeptides binding calcium on the COOH-terminal half of the protein. The toxic activities of RTX toxins in host cells usually lead to necrosis and apoptosis. The molecular mechanism of their toxicity is currently under investigation (101).

Among the bacterial toxins, the pore-forming colicins were first to be discovered in the process of searching for ways to kill bacteria (109). Colicins are produced by colicinogenic strains of nonpathogenic *E. coli* and some related species of *Enterobacteriaceae*. Colicins primarily target membranes of susceptible bacterial strains to inhibit their growth and thus present a major weapon in the on-going bacterial warfare. They form voltage-dependent ion channels by interacting simultaneously with several components of the complex membrane of susceptible cells and transforming themselves into α-helical membrane-spanning channel proteins (198). Although most colicins kill susceptible bacteria by forming ion channels in the plasma membrane of target cells, other types of colicins, such as E3, E4, E6, and cloacin DF13 or E2 and

*Physiol Rev* • VOL 88 • OCTOBER 2008 • www.prv.org
E7-E9, function as specific 16S rRNA endonucleases or nonspecific DNA endonucleases, respectively, while other colicins like colicin M or pesticin I inhibit cell wall synthesis or hydrolyze it (171).

A large family of bacterial toxins is formed by the cholesterol-binding toxins (CBTs). CBTs are also known as thiol-activated toxins and cholesterol-dependent cytolysins (106, 253). Among the best-known CBTs are streptolysin from *Streptococcus pneumoniae*, pneumolysin from *Streptococcus pyogenes*, perfringolysin from *Clostridium perfringens*, and listeriolysin from *Listeria monocytogenes*. They interact with membrane cholesterol at concentrations between 1 and 100 nM. The presence of cholesterol, which has been identified as a CBT receptor in membranes of target cells, is thus a prerequisite for CBTs to form oligomeric pores that can be made up of up to 50 subunits. CBT oligomers exist in two forms, as arc- or ring-shaped oligomers. CBT oligomerization occurs in a concentration-dependent manner (106), such that smaller arc-shaped oligomers form smaller pores compared with the larger pores of ring-formed oligomers. Given that only 10–100 CBT monomers are required for their cytotoxic action, this suggests that arc-formed pores may be more important for the cytotoxic action of CBTs than the larger ring-formed pores.

An interesting case on its own presents VacA, the pore-forming toxin of the stomach ulcer causing bacterium *Helicobacter pylori*. In contrast to most bacterial toxins, which usually form membrane pores without preference for permeating ions, VacA exhibits electrophysiological properties characteristic of the CIC channels of eukaryotic host cells (64). In addition to sharing similar conductance and ionic selectivity with the CIC channels, VacA also shares an open probability dependent on the molar ratio of permeable ions as well as single-channel events resolvable as two independent, voltage-dependent transitions, which are the very characteristic of the CIC channels. The only feature distinguishing VacA from the CIC channels is the membrane potential at which the two channels close. VacA toxin thus largely mimics the electrophysiological behavior of CIC channels of the host cells, suggesting a novel mechanism of toxin action.

### VI. VIRAL ION CHANNELS

Before discovery of the microbial world, the term *virus*, which originates from a Latin word meaning “venom,” had been used to describe any disease-causing agent. As cellular parasites, viruses depend entirely on their host cell machinery for propagation. The genome of a virus consists of either DNA or RNA, but never both. The viral genome is surrounded by the capsid, a protein coat, which together with the genome constitutes the nucleocapsid. The nucleocapsid of complex viruses may be surrounded by an additional protein covering, which in enveloped viruses has also a lipid bilayer covering, which contains membrane glycoproteins functioning as ion channels. The ion channels of enveloped viruses, such as influenza A, influenza B, and influenza C virus as well as the human immunodeficiency virus type 1 (HIV-1) form homo-oligomers of short membrane proteins of ~100 amino acids in length. These channels are called M2 (influenza A), NB (influenza B), CM2 (influenza C), Vpu (HIV-1) (93, 96, 370), and Vpr (HIV-1) (304).

The M2 protein from the influenza A virus is a minimalistic primitive ion channel, which has only one 22-residue-long transmembrane domain. Most likely four of these domains assemble in a tetrameric ion channel selective for protons (305, 306). Protons are conducted through the channel pore in response to applied membrane voltage. Proton selectivity of these channels, which is ensured by the presence of four histidines, one from each of the four helices, is essential to viral function. The channels are blocked by the drug amantadine, the only drug against influenza (251). M2, NB, CM2, and Vpu proteins do not share any sequence homology, although they have in common an amino acid sequence of ~22–26 residues with a high content of hydrophobic residues corresponding to the transmembrane domain. NB, Vpu, and Vpr are selective for cations (93, 304, 370), whereas the ion selectivity of CM2 has still to be determined. The exact role of NB and CM2 in the life cycles of the viruses is not yet identified, although they may also play a role in virus entry into a host cell (96). The ion channel activity of Vpu is thought to be associated with the ability of Vpu to facilitate the budding of new virus particles. Except for a partial high-resolution structure available for Vpu, there is no higher order structure available for any of the viral ion channels (251). Another recently identified viral ion channel is the *Chlorella* virus PBCV-1 Kcv protein. A distinguishing property of the Kcv ion channel is its high selectivity for potassium, which is believed to be critical for virus replication, since blocking the channel by amantadine or barium also inhibits virus plaque formation (308).

### VII. APPLICATIONS OF MICROBIAL ION CHANNELS AND TOXINS IN NANOTECHNOLOGY AND MEDICINE

Membrane proteins and peptides that form pores and channels in bilayers of biological membranes comprise a diverse class of molecules. The diversity of regulation and selectivity of these membrane pores together with recent advances in redesigning these pro-
teins by means of molecular biology and chemical technology provide excellent prospects for building custom molecules with wide-ranging biotechnological applications. Studies on microbial ion channels have been primarily driven by curiosity to understand structure and function, physiological role, as well as evolutionary origins of these fascinating molecules. Inevitably, as a byproduct of the basic research on these molecules, practical applications in nanotechnology and medicine have recently begun to emerge. The following examples of practical applications resulting from the research on bacterial mechanosensitive channels and the α-hemolysin toxin may illustrate this.

The MscL channel of *E. coli* was used as a template to construct a light-driven channel-nanovalve (168). For this purpose, light-sensitive synthetic compounds that undergo light-induced charge separation were attached to the channel molecule allowing the MscL nanovalve to be opened by long-wavelength ultraviolet radiation and closed by visible light. MscL was subsequently also converted into a pH-sensitive nanovalve. The opening and closing of this pH-sensitive channel can be fine-tuned by changing the pKₐ of the chemical compounds modulating the MscL properties (167). Thus the modified MscL channel can be reversibly opened and closed by light instead of by membrane tension. Such a light-driven nanovalve could, for example, find applications in pharmaceutical industry in liposome-based drug delivery systems.

Recently, it has also been recognized that bacterial MscL and MscS channels have a potential as selective targets for novel types of antibiotics (272). Both channels not only can be opened by amphipaths (235, 237) but also by parabens, the alkyl esters of p-hydroxybenzoic acid as well as eriochrome cyanine R, a biological stain commonly used as a spectrophotometric reagent for the detection of aluminium (Nguyen and Martinac, unpublished data). The difference between the action of amphipaths and parabens is that parabens appear to interact directly with the gate of the channels (272) and not indirectly via the lipid bilayer, although a recent study seems to suggest that parabens may also act on the bacterial MS channels through their action on the lipid bilayer (151). One could envisage that the novel antibiotics based on parabens or eriochrome cyanine R as lead compounds, for example, would work as channel openers of MscL and MscS. Opening of the large pores of these MS channels should in turn compromise cellular turgor of bacterial cells leading to inhibition of cell growth and proliferation. Given that, to date, MscL has been identified in over 120 prokaryotic species, including many bacterial pathogens (307), but its homologs have not been found in animal and human cells (117), this channel in particular appears as an excellent target for the novel type of antibacterial agents. An antibiotic targeting MscL could be expected to be broad-spectrum, selective for a range of pathogenic bacteria and most likely having minimal side effects to infected patients.

Recent experimental studies exploring a possibility of using the α-hemolysin protein as a model system to decipher ionic signature patterns in the genetic code of nucleic acids present an example of a practical application of basic research on microbial toxins in nanotechnology (40, 61). This application of α-hemolysin is based on the fact that single-stranded nucleic acid polymers can be transported across an α-hemolysin channel under the action of an applied electric field. Given that the translocation of the nucleic acid polymers causes transient blockades in the ion current through the channel pore and each type of nucleic acid seems to have a characteristic “current signature,” the α-hemolysin channel could thus function as a “sequencing nano-device.” Recently, the physical and chemical details of the interactions between polymer, channel, and ionic solution that lead to the electric current blockade events have been investigated.

Furthermore, the exchange of cationic α-helical peptides between the bulk aqueous phase and the transmembrane β-barrel of the α-hemolysin protein pore were examined at the single-molecule level (254). This exchange was found to be strongly dependent on voltage and peptide length. At high transmembrane potentials, the alamine-based peptides, which include bulky lysine side chains, were found to bind more strongly to the lumen of the α-hemolysin pore than highly flexible polyethylene glycols, for example. In contrast, the peptides bind weakly to the lumen of the pore and the affinity decreases with the peptide length at zero transmembrane potential. Consequently, these results indicate that α-hemolysin may also serve as a versatile nano-device for examining the interactions between positively charged signal peptides and a pore of membrane-spanning β-sheets. In addition to α-hemolysin, numerous other pore-forming peptides are being developed as antimicrobial agents with potential application as antitumorigenic agents as well as biosensors for a range of different analytes (287).

As shown here, besides gaining further knowledge from microbial channels, some of these channels can be used as tools in further research or even in therapy. As another example, transgenic channelrhodopsin of *Chlamydomonas* has become a tool to depolarize neurons with light (291, 341). This then allows us to investigate the roles of the targeted neurons in the network of behaving animals. Other technological opportunities are currently being explored. Besides the existing channel blockers in cardiological use, blockers and activators of other channels hold great promise in medicine, since all cells in the human body, excitable or not, have ion channels in their plasma membrane as well as organelle membranes. Microbial ion channels, with their fundamental similarity to human channels and
their endless variations reflecting microbial diversity, will continue to figure heavily in biomedical research, development, and practice.

VIII. CONCLUSIONS AND OUTLOOK

A. Possible Physiological Roles of Ion Channels in Microbes

The achievement and intellectual significance of the high-resolution structures of microbial channels are indeed astounding. In contrast, current understanding of the physiological roles of these channels in the microbes themselves lags behind. As stated at the outset of this review, ion-channel studies are rooted in animal physiology, especially neurobiology. The study of Paramecium ion channels reinforced the connection between channels and motile behavior, even though a Paramecium is but a single cell (see sect. nA). Recent research presents two egregious disconnects between motile behavior and ion channels. First, two major models of microbial motile behaviors are currently understood mechanistically without the involvement of ion channels. E. coli chemotaxis is understood as a series of phosphate transfers from the receptor to the effector molecules, without an ionic component (89) (although the flagellar motor is constantly driven by $\text{H}^+$ or $\text{Na}^+$). cAMP taxis of Dictyostelium amoebae is said to begin with a G protein-coupled receptor and proceed with a series of phosphorylations, again without an ionic component in signal transduction (see sect. nB1). Second, as evident throughout the entire review above, microbes that are immotile and do not have overt animal-like behavior nonetheless have various types of ion channels found in animals. While we do not yet know the true physiological roles of most of the identified ion channels in microbes, it is clear that they function far beyond regulating motilities.

The role of the mechanosensitive channels in protecting bacteria from hypsomotically induced lysis is now well established (see sect. u). The roles of ion-specific channels in microbes, however, are largely unknown. This is especially frustrating, since they include channels in well-known model microbes, such as Kch of E. coli (see sect. uF) as well as Yvc1 (see sect. nCl1b) and Tok1 (see sect. nC2) of S. cerevisiae. Knocking out kch, yvc1, or tok1 resulted in no discernable growth phenotype in various cultures or under various stresses. These genes, together with some 30% of the genes in the E. coli or yeast genome, are annotated as being “nonessential,” meaning that they are not required for growth in the pampered conditions of the laboratory. The near-universal existence of ion-channel genes in the highly streamlined microbial genomes, however, strongly indicates selective advantages they confer. It therefore seems clear that these channels are employed to handle environmental challenges, which we have yet to simulate in the laboratory. The role of the mechanosensitive channels in hypsomotic stress (see sect. uI) and CIC-ec1 in extreme acid-resistance response are cases in point (see sect. uD). A major task for microbial physiologists now is to find the environmental stresses, in which various microbial channels are called upon to function. It seems likely that these ion channels are components of novel signal transduction pathways.

As reviewed here, microbes harbor a large variety of ion channels in their cell membranes (further reading on bacterial ion channels can be found in Kubalski and Martina, Ref. 179). The cytoplasmic membranes of bacteria, archaea, algae, and fungi play an essential role in energy transduction that requires tight regulation of ion channels to allow cellular functions. A leaky membrane would not allow for energy transfer according to the chemiosmotic theory (249), because membrane potential is closely linked to the proton and sodium electrochemical gradient, which provide an interconvertible energy source that links the chemical energy of the cell to membrane transport and motility (37). Another component contributing to membrane potential in bacterial cells is the $\text{K}^+$ gradient across the cell membrane, since intracellular $\text{K}^+$ in most microbes ranges between 300 and 700 mM. Compared with animal cells, whose resting membrane potential is mainly determined by the $\text{K}^+$ gradient, membrane potential in bacterial cells, for example, is much more negative ranging between $-120$ and $-200$ mV, indicating the importance of the proton pumping and sodium-motive force in determining the resting membrane potential in bacteria. Given that the cytoplasmic pH is relatively constant, varying within a narrow range of pH 6.0 for acidophiles and pH 8.5 for some alkalophiles, the pH gradient in microbes varies significantly with the pH of the environment with the consequence that the membrane potential component of the proton and sodium electrochemical gradient varies with external pH (35, 37).

It is obvious that, with such high values for the membrane potential, strong driving forces exist for ion movements through ion channels in bacterial membranes. Given the very small capacitance of the cytoplasmic membrane of a single bacterial or archaean cell ($\sim 0.1$ pF), only a very small number of ions flowing through an ion channel would suffice to cause a substantial change in the resting membrane potential (35). Consequently, most of the time ion channels must remain tightly closed and open only transiently to ensure undisturbed energy transduction in a bacterial cell. MS channels (238) are an exception to this rule as they have evolved specifically to serve as emergency valves for fast release of osmolytes in osmotically challenged bacterial cells. A specific role for $\text{K}^+$ or any other type of ion channels in most microbes has not yet been established, although $\text{K}^+$ channels may, for
example, serve as efflux systems, when fast potassium efflux or rapid hyperpolarization is required. For such physiological studies, development of methods to monitor the membrane potential in vivo from these microbes seems necessary. Although TPP has been successfully used to monitor slow changes in membrane potential, dye-based fluorescent molecules may be useful for faster reactions. Alternately, GFP conjugated to a voltage sensor may also be used in a less invasive manner.

B. Microbial Channels, A Treasure Trove

Life originated from microbes (Fig. 1). They have dominated this planet in number, in mass, and in variety, and they will continue to dominate in the future. Although we do not have instinctive emotional connections to microbes as we have with dogs, cats, and pandas, their importance cannot be denied. If biology is to be a comprehensive understanding of all life forms, we need to pay more attention to microbes.

The biological world is vast. The six “model organisms” chosen for mechanistic investigations by contemporary biologists (the mouse, the fly, the worm, the cress, budding yeast, and E. coli) do not even begin to capture the tremendous variations in life forms and life-styles. Such variations are especially evident among microbes. They have conquered nearly all imaginable niches of the planet, in deserts, in ice, deep underground, fathoms beneath the oceans, as well as in and on other life forms. Some thrive in extreme pH, salinity, temperature, aridity, or redox environments. Some even derive energy from sources not traceable to the sun. There are dazzling varieties even among the more familiar microbes: “the swimming neuron” Paramecium, the social amoebae Dictyostelium, and the plant-animal chimera Clamydomonas. From phenomena that capture our visual imagination such as bacterial bioluminescence to those that seem beyond our awareness such as metabolism miles under our feet, the microbial world provides an endless fascinating frontier for exploration and investigation. Fascinating puzzles beckon the curious mind. Why should a unicellular Paramecium have more ion-channel genes than humans? How come the chemotactic migration of Dictyostelium amoeboid chemotaxis does not seem to employ ion channels? What are K⁺- or Na⁺-specific channels for in bacteria and archaea? Why would Synechocystis PCC6803 need a glutamate receptor channel? Answers to these questions on “molecular natural history” will certainly enrich our understanding of our living planet.

Even if our biology is ultimately selfish, aiming only at the betterment of human life, understanding microbes is still of utmost importance. History has shown that we learned about intermediary metabolism from studying yeast, learned about photosynthesis from algae, and about DNA, RNA, and their workings from bacteria and their viruses. As to the ion channels reviewed here, we now learned about mechanosensitivity, voltage sensitivity, and ion filtration from microbial ion channels. These momentous advances in our understanding are no accident. On the one hand, evolution dictates that important molecular mechanisms evolve early and therefore are universal, existing in microbes as well as macrobes. On the other hand, microbes offer laboratory advantages for molecular biology, genetics, and crystallography. We can therefore look forward to solving additional molecular mysteries from the study of microbes and their ion channels. Besides those reviewed here, such channels as the sensory channels (e.g., TRPs) and channels gated by various ligand (e.g., neurotransmitter) may also find their roots in the vast and varied microbial world, making their studies easier and deeper.

Beyond understanding, we can now also look forward to the use of microbial ion channels in biomedical practice. As described in section VII, the days may be near when we can sequence DNA through channels, release drugs through engineered MscL, effectively kill pathogens by targeting their channels, or even regulate the behavior of animals and the mental state of humans by opening engineered channels in their brains with a flash of light!

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