

Family of prokaryote cyclic nucleotide-modulated ion channels

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Cyclic nucleotide-modulated ion channels are molecular pores that mediate the passage of ions across the cell membrane in response to cAMP or GMP. Structural insight into this class of ion channels currently comes from a related homolog, MloK1, that contains six transmembrane domains and a cytoplasmic cyclic nucleotide binding domain. However, unlike eukaryote hyperpolarization-activated cyclic nucleotide-modulated (HCN) and cyclic nucleotide-gated (CNG) channels, MloK1 lacks a C-linker region, which critically contributes to the molecular coupling between ligand binding and channel opening. In this study, we report the identification and characterization of five previously unidentified prokaryote homologs with high sequence similarity (24-32%) to eukaryote HCN and CNG channels and that contain a C-linker region. Biochemical characterization shows that two homologs, termed AmaK and SthK, can be expressed and purified as detergent-solubilized protein from Escherichia coli membranes. Expression of SthK channels in Xenopus laevis oocytes and functional characterization using the patch-clamp technique revealed that the channels are gated by cAMP, but not cGMP, are highly selective for K⁺ ions over Na⁺ ions, generate a large unitary conductance, and are only weakly voltage dependent. These properties resemble essential properties of various eukaryote HCN or CNG channels. Our results contribute to an understanding of the evolutionary origin of cyclic nucleotide-modulated ion channels and pave the way for future structural and functional studies.

cyclic AMP | cyclic GMP | electrophysiology | protein purification

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Insight into the structure of HCN channels has been gained only from crystal structures of the isolated intracellular C-linker and CNBD of mammalian HCN1, HCN2, HCN4, and invertebrate spHCN1. These parts of the channel assemble into tetramers (6–9). Further structural information comes from prokaryote ion channels that are homologous to HCN and CNG channels, such as the bacterial cyclic nucleotide-regulated K⁺ channel MloK1 (10–13). MloK1 lacks a C-linker region, but has a CNBD with an overall structure that is remarkably similar to the CNBD of eukaryote HCN channels (10). Based on the dimer assembly of the MloK1 CNBD in the crystal structure, a gating mechanism has been proposed in which the pore opening in the tetrameric channel arises from the action of the four CNBDs as a dimer of dimers (10). The crystal structure of the MloK1 transmembrane domain (11) reveals a domain topology that resembles that of the voltage-gated K⁺ channel Kv1.2 (14), but with important differences. The MloK1 structure suggests that the S1–S4 domain and its associated linker in MloK1 can serve as a clamp to constrain the gate and possibly function in concert with the CNBD to regulate channel opening (11). Additionally, crystal structures have also been determined for the C-linker and cyclic nucleotide binding homology domain (CNBHD) of related ion channels, including the zebrafish EAG-like (ELK) K⁺ channel (15), the mosquito ERG K⁺ channel (16), and the mouse EAG1 K⁺ channel (17). Structural insight into the mechanism of ion permeation has been derived from a prokaryote ion channel NaK (18), which was mutated to mimic the CNG channel pore region (19). Collectively, these structural data have brought valuable information about the determinants of ion permeation, domain assembly, ligand recognition, channel gating and regulation, as well as effects of disease-causing mutations (20).

Despite this tremendous progress, crystal structures for wholeeukaryote HCN and CNG channels are still not available at present, and structural insight into fundamental aspects of ion channel function is still lacking, such as the inverse voltage sensitivity in HCN channels and the coupling between cyclic nucleotide binding and channel opening by the C-linker domain, which is, as mentioned, absent in the MloK1 channel (10). In contrast, a putative voltage-gated K⁺ channel containing a C-linker region and CNBD similar to eukaryote channels was identified in the genome of the cyanobacterium *Trichodesmium*

Significance

We describe a previously unidentified family of prokaryote cyclic nucleotide-modulated ion channels. In eukaryotes, CNG channels play important roles in signal transduction as they mediate the passage of ions across the cell membrane in response to cAMP or cGMP. In this study, we demonstrate that two prokaryote homologs, AmaK and SthK, can be expressed and purified from *Escherichia coli* membranes. We reveal that SthK has functional properties that closely resemble eukaryote HCN or CNG channels. SthK is gated by cyclic AMP, but not cyclic GMP, and selects K⁺ over Na⁺ ions in a weakly voltage-dependent manner. Our results contribute to understanding the evolutionary origin of cyclic nucleotide-modulated ion channels and pave the way for future structural and functional studies.

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erythraeum (21), here termed TerK, and it was suggested to possibly represent an ancestral HCN or CNG channel (21). However, neither structure nor function of this prokaryote homolog is known. In this study, we report the characterization of Terk and four additional prokaryote ion channels, which all contain six putative transmembrane domains, a C-linker region, and a CNBD, and apparently form a family of prokaryote ion channels with close similarity to eukarvote HCN and CNG channels. We describe the expression in Escherichia coli, detergent screening and biochemical purification of these different homologs. Moreover, we identified two homologs, SthK (from Spirochaeta thermophila) and AmaK (from Arthrospira maxima), which can be stably extracted with detergents and purified in sufficiently high amounts for biochemical and structural studies. Using confocal fluorescence microscopy and electrophysiological recordings, we describe essential functional properties of one homolog, SthK. We find that SthK has electrophysiological properties that closely resemble those of eukaryote CNG channels as it is gated by intracellular cAMP and produces large unitary currents, whereas its activity is relatively insensitive to voltage. However, unlike CNG channels, SthK contains the selectivity filter sequence -TIGYGD-, which is more similar to HCN channels and other K⁺ selective channels. We could experimentally demonstrate that SthK channels are highly selective for K⁺ over Na⁺ ions. Importantly, SthK has several sequence features that closely resemble eukaryote cyclic nucleotidemodulated channels, including a C-linker region, which is missing in previously studied prokaryote homologs, such as MloK1 (10, 12, 13) and MmaK (22). Together, these data make the SthK channel a promising candidate for future structural analysis to learn more about how mammalian CNG and HCN channels work.

Results

Bioinformatics and Sequence Analysis. To identify suitable candidates for biochemical and structural studies on HCN and CNG channels, we screened the bacterial genome database of the National Center of Biotechnology Information (http://blast.ncbi. nlm.nih.gov/) using a standard BLASTP search (23) and the human HCN2 channel sequence as a template. Several ORFs with significant sequence identity to HCN2 (more than 24%) were identified in several species, including Trichodesmium erythraeum IMS101 (TerK), which was previously identified in another study (21), as well as Arthrospira maxima CS-328 (AmaK), Lyngbya sp. PCC 8106 (LpcK), Spirochaeta thermophila DSM 6192 (SthK), and Leptospira biflexa serovar Patoc strain (LbiK). Among these species, Trichodesmium, Arthrospira, and Lyngbya belong to the genus of cyanobacteria or blue-green algae, whereas Spirochaeta thermophila is an extremely thermophilic bacterium (24) and Leptospira biflexa is a saprophytic pathogen (25). Alignment of these sequences using ClustalW (26) and further inspection revealed structural motifs that are common among members of the superfamily of voltage-gated K⁺ channels (Fig. 1A). All identified bacterial sequences contain six putative transmembrane domains (S1-S6), the K⁺-selectivity signature sequence T-V/I-GYG in the selectivity filter (SF) region, and a putative voltage sensor (S4) characterized by the sequence R/Kxx-Rxx-Rxx-R/Kxx-R, where every third residue is positively charged (arginine or lysine) and x is in most cases a hydrophobic residue (a full sequence alignment is shown in Fig. S1). Not all arginine or lysine residues are conserved in each homolog as they range from two positive charges in SthK to five positive charges in LbiK. Similar to eukaryote HCN and CNG channels, the carboxyl-terminal region of the bacterial sequences contain a CNBD connected to the transmembrane region via a C-linker. In contrast to the eukaryote channels, the bacterial channels contain only a very short amino-terminal sequence. The evolutionary relationship of these bacterial homologs is further illustrated in a phylogenetic tree (Fig. 1C) and a pairwise sequence identity diagram (Fig. 1D). The sequence identity among the different bacterial homologs varies between



Fig. 1. Sequence analysis of prokaryote cyclic nucleotide-modulated ion channels. (*A*) Sequence alignment of five prokaryote homologs with spHCN1, human HCN2, human CNGA1, human CNGB1, and Mlok1. The cartoon represents the transmembrane topology of a single-channel subunit. In the S4 segment, positively charged residues are colored in red. Residues are colored in shades of blue by using an identity threshold of 50%. Sequence features are annotated according to the secondary structures features described in the X-ray crystal structure of the HCN2 C-linker region (A'-F') and cyclic nucleotide binding domain (A-C) (6). (*B*) Cartoon representation of the HCN2 C-linker region and cyclic nucleotide binding domain (6). The left panel is a single subunit, and the right panel is a tetrameric assembly. cAMP molecules are shown as spheres (white, carbon; red, oxygen; blue, nitrogen; orange, phosphor). (*C*) Phylogenetic relationship of the homologs described in *A*. (*D*) Pairwise sequence identities of the homologs described in *A*.

32% (AmaK compared with LbiK) and 67% (AmaK compared with LpcK). Compared with human HCN2, the sequence identity varies between 24% for SthK and 32% for AmaK. Compared with human CNGA1, the sequence identity varies between 25% for LbiK and 32% for LpcK. These results indicate that the newly identified homologs could be more suitable to serve as a model for HCN and CNG channels than the previously identified MloK1 channel (10–13), which has only 22% sequence identity with HCN2 (15% with CNGA1) and essentially lacks the C-linker region. Together, these data demonstrate the existence of putative cyclic nucleotide-modulated ion channels in certain bacteria, with marked similarity to HCN and CNG channels as they contain six transmembrane domains connected to a CNBD via a C-linker region.

Expression and Detergent Screening in *E. coli.* Next, we investigated whether the five newly identified HCN/CNG channel homologs could be expressed in *E. coli.* To facilitate this study, we

expressed these homologs from synthetic genes cloned into the vector pWaldo-GFPe (27), which covalently couples GFP-Hisx8 to the carboxyl-terminal end of the target protein (Fig. 24). We transformed each homolog into BL21(DE3), C41(DE3), C43 (DE3), and their pLysS variant strains, and measured whole-cell fluorescence as an initial marker for protein expression. As a positive control, we used the prokaryote K⁺ channel construct KcsA-GFP, which is robustly expressed in *E. coli*. We observed that AmaK-GFP could be expressed at fluorescence levels of 21% in C43(DE3) and 44% in C41(DE3) compared with KcsA-GFP in BL21(DE3) (Fig. 2*B*). No fluorescence significantly higher than background could be observed for the other homologs, except for SthK-GFP, which expressed at ~20% in C41 (DE3) compared with KcsA-GFP.

Next, we conducted a detergent screen to identify detergents that extract AmaK-GFP and SthK-GFP from the membrane in an oligomeric and monodisperse state. We used fluorescence size exclusion chromatography (FSEC) (28) on crude membranes solubilized with a range of different detergents, namely CYMAL-6, lauryldimethylamine *N*-oxide (LDAO), octyl glucoside (OG), nonylglucoside (NG), octylmaltoside (OM), Fos-choline-12 (Fos-12), decylmaltoside (DM), undecylmaltoside (UDM), dodecylmaltoside (DDM) and tridecylmaltoside (TDM), lauryl maltose neopentyl glycol (LMNG), C12E9, CHAPS, and HEGA-10. The



Fig. 2. Biochemical characterization of prokaryote cyclic nucleotide-modulated ion channels. (A) Cartoon presentation of the C-terminal GFP-fusion constructs used for detergent screening. (B) Whole-cell fluorescence obtained from E. coli cultures expressing AmaK-GFP in different strains. Fluorescence intensity was normalized relative to KcsA-GFP (29). (C) FSEC profile (28) of KcsA-GFP after solubilization with 2% (wt/vol) undecylmaltoside and gel filtration in running buffer containing 0.15% undecylmaltoside. (D) FSEC profile of AmaK-GFP after solubilization under the same conditions as in C. The gel scan shows in-gel green fluorescence of His-trap purified protein (lane b) loaded on a 4-15% Miniprotean TGX gel. Lane a contains Precision Plus All Blue standard from Bio-Rad. (E) FSEC profile of SthK-GFP after solubilization with 2% dodecylmaltoside and gel filtration in running buffer containing 0.05% dodecylmaltoside. The gel scan shows in-gel green fluorescence under the same conditions as in D except that an additional ladder (Fluorescent WesternC standard from Bio-Rad) was loaded in lane b. Lane c was loaded with His-trap purified protein.

FSEC method relies on the identification of a symmetric peak running between the void volume of the size exclusion column (~7 mL) and the elution volume of free GFP (~18 mL). For example, the positive control KcsA-GFP behaves as a stable tetramer in a range of different detergents (29) and elutes as an intense symmetric fluorescent peak at 14.5 mL after solubilization with UDM (Fig. 2C). For AmaK-GFP, we also identified UDM as a suitable detergent because the FSEC profile shows a symmetric peak with a retention volume of 12.7 mL (Fig. 2D). Furthermore, we used in-gel fluorescence of nickel affinitypurified AmaK-GFP and observed a fluorescent band near the 50-kDa marker, which likely corresponds to monomeric AmaK-GFP (81 kDa). This phenomenon of gel shifting is commonly observed for membrane proteins and can be explained by the higher number of SDS molecules absorbed compared with water-soluble globular proteins (30). Additionally, we also observed a weaker fluorescent band between the 100 and 150 kDa marker, indicating that an oligometric assembly of AmaK-GFP, possibly dimers, is partially stable in SDS. For SthK, we identified DDM as a suitable detergent and observed an FSEC profile similar to AmaK-GFP. SthK-GFP elutes as a symmetric peak with a retention volume of 13.4 mL (Fig. 2E). The in-gel fluorescence of nickel affinity-purified SthK-GFP shows a single band near the 50-kDa marker, which is similar to AmaK-GFP. In addition, for AmaK and SthK several other hits were obtained from our detergent screen, including Fos-choline-12, HEGA-10, and LMNG (Fig. S2). Together, these results demonstrate that two prokaryote HCN/CNG channel homologs, namely AmaK and SthK, can be expressed in E. coli and extracted from membranes in a monodisperse state. These results pave the way for large-scale production, biophysical characterization, and crystallization screening.

Membrane Localization and Functional Characterization. To test whether SthK and AmaK can be studied in a heterologous expression system suitable for electrophysiological experiments, we tried to functionally express SthK and AmaK GFP-fusion constructs in *Xenopus laevis* oocytes.

First, we used confocal fluorescence microscopy to whole intact oocvtes and investigated whether or not SthK-GFP localizes at the cell membrane, i.e., in the periphery of the oocyte. To visualize the boundaries of the cell membrane, we incubated oocytes in a bath solution containing 5 µM Alexa Fluor 633-labeled wheat germ agglutinin (WGA-Alexa 633), a compound that labels the extracellular face of the plasma membrane. Exposure to 633-nm laser light resulted in bright red fluorescence in the periphery of both SthK-GFP-injected and control oocytes (Fig. S3). In parallel, the WGA-labeled oocytes were exposed to 488-nm laser light, resulting in a bright green fluorescence signal only in SthK-GFP-expressing oocytes (Fig. S3 A and B) but not in control oocytes (Fig. S3 C and D). Additionally, the emission spectrum of SthK-GFP obtained by spectral imaging (SI Experimental Procedures) is clearly different from oocyte autofluorescence and perfectly superimposes with the emission spectrum obtained from GFP expressed in HEK cells (SI Experimental Procedures; Fig. S4). Together, these results indicate that SthK-GFP is expressed in or near the membrane of the oocytes.

To gain further arguments that SthK-GFP is expressed in the plasma membrane and not only in submembranal compartments, we quantified the degree of overlap of the SthK-GFP fluorescence with WGA-Alexa 633 fluorescence at higher magnification. It was observed previously that the construct CNGB1b-TFP:CNGA2:CNGA4, a heterotetrameric eukaryotic CNG channel that produces reliable trafficking to the membrane as confirmed by current recordings, yields a degree of overlap of ~54% between the fluorescence profiles perpendicular to the plasma membrane yielded a degree of overlap of $65.3 \pm 4.43\%$ (n = 3) between GFP and WGA-Alexa 633 fluorescence (Fig. S3B). This result even exceeds the degree of overlap observed

in the experiment with CNGB1b-TFP:CNGA2:CNGA4 and therefore suggests significant membrane localization of SthK-GFP. As previously noted by Nache et al. (31), these results should be interpreted with caution because staining the membrane of control oocytes produced a profile with a half-maximum width of 2.0 ± 0.15 $\mu m (n = 3)$ (Fig. S3D), which by far exceeds the microscope resolution (0.3 mm) and typical membrane thickness of ~3-4 nm. The result can be explained by the extensive folding of the oocvte membrane (31). Despite this optical limitation, our confocal fluorescence microscopy experiments indicate that SthK-GFP produces significant green fluorescence at the oocyte membrane and suggests that SthK-GFP localizes to the cell membrane. Performing the same experiments with AmaK-GFP did not result in GFP-related fluorescence signals, indicating that there is no expression of this construct in the oocyte membrane. Spectral images comparing AmaK-GFP with SthK-GFP injected oocytes are shown in Fig. S5.

Channel Gating. We first tried to record whole-cell currents by using the two-electrode voltage-clamp technique. The result was negative, suggesting that SthK channels are not functional or have a gating mechanism that is different to eukaryote HCN channels. Next, we used the patch-clamp technique with patches in the inside-out configuration, which allows the investigator to apply molecules to the internal face of the membrane. Because SthK carries the -TIGYGD- selectivity filter sequence in the pore loop, suggesting a channel that selects K⁺ over Na⁺ ions, the following experiments were performed in symmetrical solutions containing 150 mM K⁺. In patches containing presumably many SthK channels (macropatches), the application of cAMP (500 μ M) to the bath solution produced robust current responses at positive and negative voltages when applying a voltage ramp from -130 to +130 mV (Fig. 3). Such a response was not observed in the absence of cAMP (Fig. 3A) or in patches from uninjected control oocytes. This result suggests a gating mechanism for SthK similar to eukaryote CNG channels, which can be activated by cyclic nucleotides and which are relatively insensitive to voltage. The cAMPevoked current through SthK channels was reversible when removing the cAMP and the amplitude was stable after an initial run-up of tens of seconds following the cAMP application. There was no indication for any inactivation or desensitization in the presence of cAMP.

Next, we studied the cAMP sensitivity of SthK channels in greater detail. Fig. 3B shows the concentration-activation relationship recorded at 150 mM symmetrical K⁺ solution. Fit of the data points with Eq. S1 yielded the cAMP concentration of half-maximum activation, EC₅₀, of $3.68 \pm 0.55 \mu$ M and the Hill coefficient, H, of 1.33 ± 0.08 (n = 10). The Hill coefficient larger than unity indicates that the subunits cooperate upon channel activation.

Notably, we did not observe any current response when applying cGMP instead of cAMP (Fig. 3 *A* and *C*). To test whether cGMP is not able to bind to the SthK binding domain or acts as a competitive antagonist, we applied 3 μ M cAMP, a concentration near the EC₅₀ value, together with 5 mM cGMP (Fig. 3*C*). Under this condition, cAMP is not able to activate the channel, indicating a competitive of cGMP and cAMP at the binding site. Thus, cGMP acts as a competitive antagonist, which is remarkable, because both cAMP and cGMP are agonists in most of the cyclic nucleotide-modulated ion channels described so far.

Ion Selectivity. To confirm the predicted selectivity of K^+ over Na⁺, we applied voltages from -100 to +100 mV in 20-mV steps at 500 μ M cAMP with either 150 mM KCl or NaCl in the bath and the pipette solution, respectively. The resulting current-voltage relations and representative current traces for each case are shown in Fig. 44. The results demonstrate that SthK channels allow K⁺ but not Na⁺ ions to permeate, similar to the prokaryotic cyclic nucleotide-gated K⁺ channel MmaK (22) and MloK1 (13). At symmetrical potassium, there is only weak



Fig. 3. Ligand selectivity in inside-out macropatches. (A) A voltage ramp ranging from -130 to +130 mV in 4.5 s was applied to an inside-out macropatch excised from a SthK-GFP-expressing Xenopus laevis oocyte. The black trace represents the current response after applying 500 μ M cAMP at symmetrical potassium. There was no current response after removal of cAMP or applying 5 mM cGMP to the same patch, shown as gray and red traces, respectively. (B) The concentration-activation relationship for cAMP was obtained at symmetrical potassium. Means were calculated from 3 to 14 recordings. Error bars indicate SEM. The black curve is the best fit of the mean values using Eq. S1. When fitting single recordings, we obtained the following: $EC_{50} = 3.68 \pm 0.55 \mu M$ (n = 10), H = 1.33 ± 0.08 (n = 10). (C) A voltage ramp under similar conditions as described in A was applied to an inside-out macropatch. The representative traces show current responses after applying 3 µM cAMP (black traces), 3 µM cAMP plus 5 mM cGMP (orange trace), or 5 mM cGMP (red trace). The ligands were washed in the same order as shown. There was no current response as long as cGMP was applied.

outward rectification. This resembles the voltage dependence of eukaryotic CNG channels under symmetrical conditions (32, 33).

Single-Channel Properties. We also studied the activity of single SthK channels. Fig. 4*B* shows a representative example of current traces and the corresponding amplitude histogram. The amplitude of the single-channel current at +100 mV was 7.1 \pm 0.4 pA (n = 4), resulting in a single-channel conductance of 71 pS. This value is in the upper range of reported values for eukaryote CNG channels (20–80 pS) (reviewed in ref. 4), but is much higher than that in eukaryotic HCN channels, which have an extremely low single-channel conductance of only 1–2 pS (34–36). The probability for one channel to be open, P_{o} , at saturating ligand concentration was $13.8 \pm 1.72\%$ (n = 3), which is low compared with the respective P_o values in eukaryotic CNGA1 (94.0%) (37), CNGA2 (99.0%) (38), and HCN2 (>99%) (36) channels.

Discussion

Only a few prokaryotic ion channels have been successfully expressed in *Xenopus laevis* oocytes as heterologous expression system, including the mechanosensitive channel MscS (39), a proton-gated ion channel from the nicotinic acetylcholine receptor family (40), or an ATP-sensitive K⁺ channel (41). Bacterial CNG channels could only be successfully studied as reconstituted proteins in liposomes (13, 42) or in giant *E. coli* spheroplasts (22). In the present study, fluorescence imaging and electrophysiological patch-clamp recording demonstrated that SthK-GFP can be expressed as a functional ion channel protein



Fig. 4. Functional characteristics of SthK in inside-out patches. (A) (Left) Current-voltage relationship as measured under four different jonic conditions. Bath solution and pipette solution contained either K⁺ or Na⁺ ions, respectively. The bath solution additionally contained a saturating cAMP concentration of 500 µM. Positive values indicate outward currents, and negative values indicate inward currents. Current amplitudes are normalized to the maximum value obtained at +100 mV at K_{int}/K_{ex} and K_{int}/Na_{ex} , respectively. Means were calculated from three to nine patches. Error bars indicate SEM. (Right) Representative currents (black traces: at 500 µM cAMP; gray traces: no cAMP) for each of the four ionic conditions. (B) Singlechannel events were recorded at a saturating concentration of 500 μ M cAMP at symmetrical potassium and +100 mV. (Left) Representative currentamplitude histogram for a three-channel patch. The main histogram was fitted by the sum of three Gaussian functions (for one closed and two open levels) (black line). The amplitude of the single-channel current was determined to be 8.1 pA (with a SD obtained from the Gaussian fit of 0.02 pA). The Inset shows the same histogram at higher magnification to visualize the third open level peak. This histogram was fitted by the sum of two Gaussian functions (black line). (Right) Shown are representative traces from the same recording as shown in the left panel. Closed and open levels are indicated as dashed (c) and dotted lines (o).

in the plasma membrane of *Xenopus* oocytes. We investigated functional parameters of SthK channel gating in excised insideout macropatches to learn about its relationship to other members of the family of cyclic nucleotide-modulated ion channels.

Most of the prokaryotic and eukaryotic cyclic nucleotide-gated channels are sensitive to both cAMP and cGMP, very often preferring one ligand over the other (4, 43). It has been shown that, regardless of ligand preference, a highly conserved arginine (Arg618 in human HCN2 or Arg561 in human CNGA1) in the CNBDs of eukaryotic (6) and prokaryotic cyclic nucleotidemodulated channels (13) as well as other cyclic nucleotidedependent proteins (44) is essential for cNMP binding. Mutating this arginine to glutamate decreased the affinity for cNMP dramatically in both eukaryotic CNG and HCN channels (45, 46). A similar effect was observed when neutralizing this residue in the prokaryotic MloK1 (42). Interestingly, this conserved arginine is also present in the sequences of all prokaryotic channels described in our study (Fig. 1), indicating that all these homologs are able to bind cyclic nucleotides.

Channels that bind both cAMP and cGMP have an amino acid residue next to this arginine, which provides a hydroxyl group: a threonine in eukaryotic channels (6) and a serine in MloK1 (13). In SthK, there is a threonine at that very position, arguing for a binding of both cyclic nucleotides. Another arginine residue, which has been described to be important for both cGMP and cAMP binding resides in the C-helix of the binding pocket

(Arg348 in MloK1) (47). This arginine was also found in the SthK sequence. Herein, we showed that, as expected from the sequence, indeed both cAMP and cGMP do bind to the SthK CNBD, however, with opposite effects on channel gating: While binding of cAMP activated SthK channels, cGMP binding did not. This is remarkable because to our knowledge it is the first time that an antagonistic effect of a cNMP to a CNG/HCN channel homolog has been demonstrated. However, the idea that cGMP can act differently than as a pure agonist has been proposed earlier for catfish CNGA2. Young and co-workers described a so-called bimodal agonism, meaning that cGMP exhibits two opposite modes of action: "pro action" that enhances receptor activation and "con action" that suppresses activation (48–50). It would be interesting to investigate whether there is any analogy in the fCNGA2 bimodal agonism and the cGMP antagonism found herein.

The sequence of the SthK pore region contains a TIGYGD motif in the selectivity filter, arguing for a K⁺-selective channel. Our experimental results confirmed this prediction: SthK channels allow K⁺ but not Na⁺ to permeate in both inward and outward direction (Fig. 4*A*). A similar ion selectivity was described for other bacterial CNG channels, such as MloK1 (13) and MmaK (22).

The activation mechanism of the different types of cyclic nucleotide-modulated channels is different. Eukaryote CNG channels strictly depend on the binding of cyclic nucleotides and are only weakly voltage dependent. In contrast, eukaryote HCN channels require hyperpolarizing voltages for activation and are only modulated by cyclic nucleotides. In our study, we show at symmetrical K^+ that SthK is only weakly voltage dependent, which resembles the behavior of eukaryotic CNG channels (32, 33). One possible explanation for the lack of voltage dependence in SthK could be the relatively low number of three positively charged residues in the intracellular half of the S4 domain. In comparison, voltage-dependent eukaryote HCN channels have seven positively charged residues within S4. In addition to S4, the secondary structure of the S4-S5 linker was also shown to play an important role for voltage sensing (51, 52). In voltage-dependent K^+ channels, this linker is a relatively rigid α -helix, transferring the voltage-induced conformational changes of the S4 to the channel gate (51). For eukaryotic CNG channels, it was suggested that the S4-S5 linker is unable to transfer this conformational change as it is a short and flexible loop, making the channel independent of voltage (52). Similar to CNGA1-4 isoforms, the specific amino acid composition of the S4-S5 linker in SthK predicts a flexible loop rather than a rigid α -helix (see full sequence in alignment in Fig. S1). Additionally, the S4–S5 linker in SthK contains the structure-breaking amino acid proline at position 132, which is conserved in almost all eukaryotic CNGA1-4 subunits of different species and which is suggested to significantly participate in making the S4-S5 linker unable to transfer S4 movements to the pore (52).

In summary, several sequence motifs of SthK channels resemble those in eukaryote CNG channels and therefore suggest similarity in function. Certain key amino acids in the cyclic nucleotide binding domain as well as the S4-S5 linker are conserved between SthK and eukaryote CNG channels. In agreement with this conservation, our electrophysiological experiments demonstrate that SthK is gated by the intracellular application of cAMP. However, unlike eukaryote CNG channels, SthK channels contain a sequence in the selectivity filter that resembles HCN channels and other K⁺-selective channels but not CNG channels. Our functional characterization indeed confirms that SthK is a K⁺-selective channel. Importantly, SthK and all of the other homologs described in our study contain a C-linker region, which is absent in previously studied prokaryote homologs, including MloK1 and MmaK. This observation might contribute to a better understanding of the evolutionary origin of CNG channels and suggests that an ancestral CNG-HCN gene may have arisen already in the bacterial stage of evolution. Finally, our biochemical data demonstrate that SthK and AmaK can be expressed and purified in quantities sufficient for future biochemical and structural studies. If available, X-ray crystal structures of SthK and AmaK, most desirably in the absence and presence of cyclic nucleotides, may bring essential progress for our understanding of the channel gating, because these bacterial channels do contain the C-linker region.

Experimental Procedures

LpcK, TerK, SthK, LbiK, and AmaK were expressed as C-terminal GFP fusions using the vector pWaldo-GFPe or pGEM-HE for expression in *E. coli* or *Xenopus* oocytes, respectively. Confocal fluorescence microscopy was used

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to determine protein localization in the membrane. Patch-clamp recordings were carried out in the inside-out configuration. Full methods are described in *SI Experimental Procedures*.

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