Crystall structures of a pentameric ion channel gated by alkaline pH show a widely open pore and identify a cavity for modulation

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Pentameric ligand-gated ion channels (pLGICs) constitute a widespread class of ion channels, present in archaea, bacteria, and eukaryotes. Upon binding of their agonists in the extracellular domain, the transmembrane pore opens, allowing ions to go through, via a gating mechanism that can be modulated by a number of drugs. Even though high-resolution structural information on pLGICs has increased in a spectacular way in recent years, both in bacterial and in eukaryotic systems, the structure of the open channel conformation of some intensively studied receptors whose structures are known in a nonactive (closed) form, such as \textit{Erwinia chrysanthemi} pLGIC (ELIC), is still lacking. Here we describe a gammamaproteobacterial pLGIC from an endo-symbiont of \textit{Tevnia jerichonana} (sTeLIC), whose sequence is closely related to the pLGIC from ELIC with 28\% identity. We provide an X-ray crystallographic structure at 2.3 Å in an active conformation, where the pore is found to be more open than any current conformation found for pLGICs. In addition, two charged restriction rings are present in the vestibule. Functional characterization shows sTeLIC to be a cationic channel activated at alkaline pH. It is inhibited by cations, but not by quaternary ammonium ions, such as tetramethylammonium. Additionally, we found that sTeLIC is allosterically potentiated by aromatic amino acids Phe and Trp, as well as their derivatives, such as 4-bromo-cinnamate, whose cocrystal structure reveals a vestibular binding site equivalent to, but more deeply buried than, the one already described for benzodiazepines in ELIC.

Significance

Pentameric ligand-gated ion channels (pLGICs) mediate fast signal transduction in animal nerve cells through neurotransmitters. Mutation of some of these receptors in the brain causes severe nervous system diseases. The high sequence diversity of prokaryotic receptors makes them unique model systems to understand evolutionary conservation in gating and sensitivity to allosteric modulators. We present the 2.3 Å X-ray structure of a pLGIC (sTeLIC) from a gammamaproteobacteria that is activated at alkaline pH. The structure at pH 8.0 displays an unusually open pore. It is unchanged, but less flexible, in the presence of a positive allosteric modulator that binds in a cavity where benzodiazepines are found in \textit{Erwinia chrysanthemi} pLGIC. This cavity is also present (and druggable) in the SHT\(_2\)-receptor.

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species since 2009, even though they certainly display a wide range of pharmacologies and allosteric conformations. Thus, the wealth of information contained in the diversity of bacterial pLGICs has not been fully explored. Additionally, to understand their evolutionary history, it is important to understand all of the differences between eukaryotic and bacterial pLGICs at the molecular level (22).

GLIC was the first pLGIC to be solved both in an apparently open form, at a resolution of 2.4 Å (23), and in the resting form. The latter form reached only a resolution of 4.35 Å but contained four pentamers in the asymmetric unit, which allowed the unambiguous tracing of all of the chains after noncrystallographic symmetry averaging (24). In molecular dynamics simulations, the structure of GLIC obtained at low pH relaxes to a slightly more open form where some permeation events could be identified (23). Other crystal forms, called the locally closed forms (LC forms), were identified through mutations introducing disulfide bridges at key positions, or even single-point mutations in either the M2 α-helix TMD region or the M2–M3 loop (25–27). It is possible that these LC forms represent a preactivation state on the reaction pathway between the closed to open forms, as suggested by experimental (28) and computational studies (29). An LC form was also found in one crystal structure as coexisting with the open form at pH 4 in the same asymmetric unit (24). Recently, another crystal structure formation was found that was suggested to represent a desensitized form (30). Finally, GLIC presents an excellent model system to study allosteric modulation at the structural level by general anesthetics (6, 7, 11, 31), alcohols (7), xenon (32), and barbiturates (9). Structural studies and determination of the binding sites of allosteric modulators correlate well with chemical affinity-labeling studies in solution (33). The precise mechanism by which protons activate the channel remains unknown, but a systematic mutational analysis identified a key cluster of residues, mainly located at the ECD–TMD interface, as critical for GLIC gating (34).

The other bacterial receptor (ELIC) for which structural information is available, also has many entries (17) in the Protein Data Bank (PDB), involving several mutants and some very informative complexes with divalent cations (35), benzodiazepines (8), and general anesthetics (36). Electronic-spin resonance studies have also been conducted on this channel (37). However, its crystal structure has only been obtained with the ion channel pore in the closed form. Indeed, all ELIC crystal structures are found in a closed–pore conformation, even in the presence of a bound agonist and for mutants favoring the active form (38); therefore, the functional state represented by the crystal structure is still not completely understood. It was even suggested that this closed form is actually an uncoupled form, reminiscent of the uncoupled form of the nAChR (39, 40). It was suggested that ELIC could be a representative member of a new subfamily of pLGICs due to its peculiar behavior in the presence of quaternary ammonium ions that are usually channel blockers, and that its gating properties should be extrapolated only with great care to other pLGICs (41). Analysis of the aromatic residues at the interface of M1/M3 and M4 helices also suggested that ELIC’s TMD is dissimilar to GLIC’s (40).

For these reasons, we decided to get more structural information from bacterial pLGICs closely related to ELIC, especially in the gammaproteobacteria genus, in the hope of crystallizing it in an open form or both its closed and open forms. In the following we describe our results for a gammaproteobacteria pLGIC (referred to as sTeLIC in the rest of the paper), having 28% sequence identity with ELIC, found in the endosymbiont of *Tevnia jerichonana*. This symbiont bacteria is found inside the giant tube-worm *T. jerichonana*, which lives close to deep-sea hydrothermal vents (42).

**Results**

**sTeLIC Is Activated at Alkaline pH.** Initial electrophysiological characterization of sTeLIC expressed in *Xenopus* oocytes showed it is potentiated by MES buffer and inhibited by divalent cations. Therefore, Hepes and Tris buffers were used for all subsequent evaluations, after no effect was found for either buffer. Calcium was removed from the recording solution, whereas to maintain the stability of the membrane during recordings Mg²⁺, the weakest inhibitor, was maintained as a divalent cation. For experiments where long washes were needed, 1 mM CaCl₂ was present in the wash solution only. Under these conditions, sTeLIC was found to be open at basic/natural pHs and closed at acidic pHs (43). Using a perforation buffer at pH 5, which keeps sTeLIC closed, pH-jumps to more basic conditions elicit robust currents characterized by a fast activation, followed by a rapid desensitization. Small currents start to be seen at pH 7.5 and maximal currents occur at pH 9.5. Fitting all pH activation data according to the Hill model yielded a pH₀ of 8.6 ± 0.4 (Fig. 1B). Long duration applications at high pH resulted in a strong current decay (Fig. 1C), which was reversible upon return to pH 5, a property characteristic of desensitization.

**Crystal Structure at pH 8.0 Displays a Widely Open Pore.** Initial crystallization screens yielded crystals of sTeLIC diffracting only up to 6 Å. Extensive optimization, including seeding and dehydration protocols, gave crystals diffracting to 3.5 Å. After further optimization with various additives, we found that adding 6.5 mM monethyl-β-D-glucose (NDG) gave the best crystals, at pH 8.0, diffracting up to 2.3 Å resolution. The structure of sTeLIC was solved by molecular replacement using the open structure of GLIC (PDB ID code 4HFI) (23) as the search model. The final model of sTeLIC, which leaves only six unresolved residues at the N terminus and four residues at the C terminus, has excellent refinement statistics (SI Appendix, Table S1) and electron-density maps of high quality (SI Appendix, Fig. S1). The crystal-packing molecular arrangement shows both parallel and antiparallel, head-to-tail, pentamer interactions, which consist essentially of ECD–ECD or TMD–TMD contacts (SI Appendix, Fig. S2), a phenomenon already seen in the GLIC (15) (PDB ID code 3EAM) and β3-GABAA-R (18) (PDB ID code 4COF) structures.

The overall architecture of the sTeLIC is the same as for other receptors from the pLGIC family. The five monomers are arranged along a fivefold symmetry axis perpendicular to the cell membrane with small deviations from C5 exact symmetry evaluated for pairs (1,2), (1,3), (1,4), and (1,5), respectively, as 0.17, 0.09, 0.14, and 0.14 Å. The receptor is 66 Å in the largest diameter and 114 Å tall (Fig. 1D). The TMD of one monomer is composed of a bundle of four transmembrane α-helices (M1–M4), with M2 facing the ion channel pore (Fig. 1E), flanked by M1 and M3 helices of the same subunit and the M1 α-helix from a neighboring (complementary) subunit. The ECD of the monomer contains 10 β-strands (β1–β10), which form a compact curled β-sandwich, where β4, β7, β9, and β10 compose the inner β-sandwich [in the principal (+) interface], and the rest form the outer β-sandwich [in the complementary (−) interface]. An amphipathic α-helix α1 is inserted between the β3 and β4 strands, at the top ECD vestibule, with its hydrophobic part interacting with the top section of the β-sandwich and its hydrophilic part facing the vestibule (Fig. 1E). Other important regions, such as the β1–β2 loop, the M2–M3 loop, and the Pro-loop at the TMD–ECD interface are also indicated in Fig. 1E, as well as loops known to be important to bind the orthosteric-site agonist, such as loop C (β9–β10 loop) and loop B (β7–β8 loop). In Fig. IF we have highlighted all strictly conserved residues resulting from a multiple alignment of 11 bacterial pLGICs, including sTeLIC, ELIC, and GLIC. In addition, we define here loop Ω (β4–β5 loop), which forms an almost continuous ring in the lumen because of C5 symmetry (Fig. 1E and F).

The columnar-shaped ion channel pore is constituted by the five M2 α-helices, the pore-facing residues of which shape the ion translocation pathway (Fig. 2A and B). The diameter of the pore ranges from 11 to 15 Å (Fig. 2C), and the residues bordering the ion channel pore from the beginning to the end of M2 α-helix, are D4′, D2′, G6′, L9′, I12′, A13′, F16′, T17′, and S20′. The distribution of the pore-lining residues is in accordance with those of other
Overall characterization of sTeL

SD. (\(\alpha\beta\)D\(\beta\)\(\pm\)SI Appendix – 0.4 with a Hill-slope of 2)

E3961 (instead of S or T), D2 via an alanine and D2 or Br (Fig. 2) anomalous peak (generally polar but not shown in Figure 3).

functionally important loops (in brown), and the strictly conserved residues (red ovals) among bacterial receptors (whose multialignment is shown in SI Appendix, Fig. S13).

members of the Cys-loop receptor family (Fig. 2E), with the exception of G6′ (instead of S or T), D2′ (generally polar but not charged), and K-1′, whose charge is counter-balanced by a salt bridge with D2′ of the neighboring subunit (see below). The negatively charged residues, D-4′, whose side chains point to the center of the pore in its cytoplasmic end, create a strong binding site for a Na\(^+\)/K\(^+\) cation (Fig. 3B) and strongly suggest that stTeLIC is a cationic channel. Compared with other members of the pLGIC family, the ion channel pore of stTeLIC has the widest diameter, with a minimum value of 11 Å at the level of D2′ (Fig. 2C).

The closest comparable pore diameter is that found in the open-state structure of the \(\alpha\)-Gly-R, whose structure was determined by cryo-EM at 3.9 Å resolution (20). Superimposition of stTeLIC with the open state of \(\alpha\)-Gly-R reveals that in stTeLIC the top of the M3 \(\alpha\)-helix and the M2–M3 loop move further outward and the pre-M1 region moves closer to the M2 and M3 \(\alpha\)-helices to form a more compact helical bundle (SI Appendix, Fig. S3L).

The widely open conformation of the pore is stabilized by two salt bridges, as illustrated in Fig. 2D, one between D293 in M4 and R225 in M2, and one between K224 (−1′) and D227 (2′). The profile of the pore is wider than other resolved pore profiles that have been assigned to the open-state (Fig. 2C). We therefore assign this conformation to an open/active state, as the transmembrane pore diameter is compatible with permeation of sodium and potassium (or chloride) ions even with their hydration shells (43). In the electron-density map of the pore itself, we found five strong individual peaks (Fig. 3A). The Fo–Fc map on the surface of the ion channel pore during the model building, which were modeled by five NDG detergent molecules. NDG was used as an additive to get the high-resolution diffracting crystals; its curled aliphatic tail appears to be inserted into the crevice formed by two adjacent subunits, while its hydrophilic sugar head faces the channel pore (SI Appendix, Fig. S4–D). A structure obtained from crystals grown without NDG shows that the conformation of the ion channel pore is unchanged (SI Appendix, Fig. S4E), thereby excluding the influence of the NDG upon the widely open transmembrane pore conformation.

Monovalent Ion Binding Sites and Permeation Pathway. To experimentally investigate the interaction of stTeLIC with monovalent cations or anions, we performed crystal-soaking experiments with Cs\(^+\) or Br\(^-\) that have both a substantial anomalous signal at wavelengths attainable in synchrotron beamlines. Soaking the crystals with 150 mM KBr gave no signal in the anomalous map, indicating no strong binding site for bromide anions. On the other hand, crystals soaked at a final concentration of 150 mM CsCl gave strong residual densities in the Fo–Fc map on the surface of the ion channel pore (SI Appendix, Fig. S5A). A structure obtained from crystals soaked with Cs\(^+\) and one between K224 (−1′) and D227 (2′). These residues were mutated into an alanine and the D2′A (D227A) mutant was found to be nonfunctional, while D-4′A (D221A) has a marked loss of function phenotype (Expression S4–D). These residues were mutated into an alanine and the D2′A (D227A) mutant was found to be nonfunctional, while D-4′A (D221A) has a marked loss of function phenotype (Expression S4–D). These residues were mutated into an alanine and the D2′A (D227A) mutant was found to be nonfunctional, while D-4′A (D221A) has a marked loss of function phenotype (Expression S4–D).

Based on the structure, the electrostatic potential was calculated and mapped onto the protein surface (Fig. 3B). Clearly there is an intermediate channel in the lumen of the ECD with a strong negative potential, mainly from residues E69, E70, E106, D88, and E27, explaining the binding sites of the bound cations in the vestibule (Fig. 3C). Taken together, these structural data strongly suggest that stTeLIC is a cationic ion channel. This prediction is consistent with the conclusion of the electrophysiological analysis presented below. Surprisingly, the surface representation reveals that there are two
constriction rings in the ECD, one at the level of K66 in the α1 amphipathic helix, and another one at the level of D88–R86 in loop Ω (β4–β5 loop) (Fig. 3A and B). The carboxylate group of D88 interacts with atom Nε of R86, which indicates a mesomeric form of the guanidinium group where the positive charge is carried by this atom. The distance of terminal Nε amino groups of symmetry-related mates is only 2.8–3.2 Å, showing potential hydrogen bonds between them. If these side chains were frozen in this position in solution, they would prevent ion flow through the vestibule.

The functional significance of the constriction rings was tested by site-directed mutagenesis and electrophysiological experiments in oocytes, which showed that suppression of the charged side chains by mutation to alanine of R86 or K66, or both, produced functional channels and did not significantly affect the pH50 (Fig. 3C). R86 is not conserved in either GLIC or ELIC, whereas D88 is conserved except in GLIC. Negatively charged residues facing the pore at the beginning of M2 are in red, hydrophobic ones in yellow, and polar ones in green.

The crystal structure of the R86A mutant obtained in the same conditions as the WT-sTeLIC shows no change in the pore architecture and simply removes the constriction point seen in the WT structure’s vestibule (SI Appendix, Fig. S7).

**sTeLIC is a Cationic Channel.** Ion charge selectivity of sTeLIC was examined in oocytes using a change in pH from 5 to 8. Reduction of the extracellular Na⁺ and Cl⁻ concentrations (kept equal and referred to as the NaCl concentration) produced a left shift of the reversal potential (Fig. 4A), and the shift varied with the ratio of extracellular NaCl concentrations almost as predicted by the Goldman, Hodgkin, and Katz (GHK) voltage equation assuming PNa/PCl = 0 (Fig. 4B). This analysis, which only supposes constant intracellular ion concentrations, leads us to conclude that PNa/PCl is low for sTeLIC. Subsequent patch-clamp recordings from baby hamster kidney (BHK) cells provided recordings of current elicited by a potentiator at pH 7.5 and measurements of reversal potentials (Erev) at various extracellular NaCl concentrations, using a NaCl-based intracellular solution (Fig. 4C). A graphic plot of the Erev values measured at various extracellular NaCl concentrations showed PNa/PCl < 0.1 (Fig. 4D). From individual Erev values obtained at 17 mM NaCl, calculation of the permeability ratio using the GHK equation indicates a PNa/PCl value of 0.05 ± 0.02 (n = 5). This charge selectivity analysis, based on a PNa/PCl evaluation, leads to the conclusion that sTeLIC is a cationic channel.

**Characterization of Allosteric Potentiators of sTeLIC.** To identify small molecules acting on sTeLIC, known pLGIC agonists (nicotine, aceetylcholine, 5-HT, GABA, glycine, glutamate, and histamine) and

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**Fig. 2.** sTeLIC has a widely open ion-channel pore. (A) Top-down view of sTeLIC M2 α-helices. Amino acids facing the ion-channel pore are shown as sticks with the helical path colored in green (polar), yellow (hydrophobic), and red (charged) according to the property of the corresponding side chain. The distance of negatively charged amino acids with their counterparts from the adjacent and symmetry-related subunits are shown in a dashed line. (B) Side view of M2 α-helices. The pore is colored in blue. Only two helices are shown for clarity, using the same color code as in A. Both absolute (left) and relative (right) numbering of amino acids facing the pore are shown. (C) Pore radius profile of sTeLIC (blue), Gly-R86open 3JAE (pink), GLICopen 4HFI (red), GluClclosed 2NL0 (cyan), and ELICclosed 2YN6 (light green) along their fivefold symmetry axis. (D) Two different views of the bottom part of TMD, highlighting two different salt bridges with a dashed line (D293-R225 and D227-K224). (E) Sequence alignment of the M2 α-helix and part of its neighbor M4 α-helix containing the residues involved in salt bridges between sTeLIC and other members of the pLGIC family with a known structure. The pair D227-K224 is clearly nonconserved, while the R225-D293 pair is conserved except in GLIC. Negatively charged residues facing the pore at the beginning of M2 are in red, hydrophobic ones in yellow, and polar ones in green.
other compounds were tested for the potentiation or inhibition of the pH 7.5-elicited currents, with or without potentiator (SI Appendix, Table S2). Most compounds were inactive, but some amino acids were found to have a small potentiating effect (i-Met, i-His, i-Leu), with the aromatic amino acids i-Phe and i-Trp having the most promising potentiating effect (i-Tyr could not be tested beyond 1 mM due to its low solubility). The effect of aromatic amino acids and their derivatives was systematically studied. It was found that they robustly potentiate the pH 7.5-elicited currents, and their dose-response curves at pH 7.5 were established (SI Appendix, Table S3). All compounds, including the nontitratable Nα-acetyl-i-tryptophan ethyl-ester, do not elicit currents at pH 5, suggesting that they cannot activate the receptor by themselves and that they rather act as allosteric potentiators of the pH-elicited currents (SI Appendix, Fig. S8).

From the study presented in SI Appendix, Table S3, it appears that compounds with an aromatic ring are active at lower concentrations, which may indicate a better affinity for the site involved in potentiation. Substitution by bromine at the “para” position enhances the apparent affinity of the compound. The most potent compound found in the series was 4-bromo cinnamate (4-BrC) with an EC50 of 21 ± 8 μM (Fig. 5A and B) when evaluated at pH 7.5. A strong current decay in the presence of 4-BrC at 3.0 Å resolution (Fig. 5A) indicates a 4-BrC-induced desensitization of sTeLIC. Additionally, a clear indication of cooperativity is seen for most of the potentiating compounds with a Hill number greater than 2. Because X-ray crystallography indicates that 4-BrC fully occupies the five binding sites and that this binding lowers the B-factors of the whole receptor, this large Hill number likely arises from 4-BrC forming cooperative behavior as stated in the Monod–Wyman–Changeux theory (45). Subsequently, 4-BrC was used in conjunction with pH activation for many experiments evaluating the channel characteristics of sTeLIC.

Structural Characterization of Allosteric Potentiator Binding. To gain structural information for molecular recognition between sTeLIC and 4-BrC, we solved the cocrystal structural complex of sTeLIC with 4-BrC at 3.0 Å resolution (Fig. 6A). Both the Fourier Fo–Fc map (contoured at 3 σ) and the anomalous map (contoured at 5 σ) using data collected at the absorption edge of bromine allow unambiguous construction of 4-BrC within the density (Fig. 6C). The binding site of 4-BrC is deeply buried in the central vestibular-facing intrasubunit cavity of the β-sandwich, where the hydrophobic head interacts with the hydrophobic core of the β-sandwich via van der Waals interactions (Fig. 6B). Its carboxylate tail forms a salt bridge with R92 from strand β5 and is exposed to the lumen. The overall conformation of this complex is very close to the conformation obtained for sTeLIC in the absence of 4-BrC, with an rmsd of 0.22 Å. However, the distribution of B-factors throughout the structure is strikingly different from the one of the unbound structure.

The binding site of 4-BrC, with interacting residues indicated in Fig. 6C, largely overlaps with the intrasubunit binding site of flurazepam (FZM) in ELIC, in which FZM also acts as a potentiatior at low concentrations (8). Detailed structural comparison of sTeLIC–4-BrC with ELIC-FZM reveals several distinct features. First, 4-BrC inserts more deeply into the center of the β-sandwich, compared with the position of FZM in ELIC (SI Appendix, Fig. S94). Second, five 4-BrC molecules can be assigned into the pentamer of sTeLIC, with one 4-BrC per monomer (Fig. 6A), whereas just one molecule of FZM could be found in one of the intrasubunit sites for the whole pentamer of ELIC (8). The 4-BrC binding site is very close to the Br-acetate binding site in GLIC (46) (SI Appendix, Fig. S9B).
to ECD lumen. We have tentatively assigned this density to a PEG 200 molecule (SI Appendix, Fig. S9 D and E), the precipitant used for crystallization. This density is localized similarly to the homologous Br-acetate site in GLIC and benzodiazepine FZM in ELIC (SI Appendix, Fig. S9 F and G). This extra density may explain why sTeLIC is maintained in the open state at this pH, as PEG 200, which is present at high concentration (0.9 M) in the crystallization drop, was found to potentiate sTeLIC response, especially in conjunction with DMSO at the concentration used for crystallization (SI Appendix, Fig. S10 and Table S2). It must be emphasized that assignment of the extra density to a PEG 200 molecule is speculative at this stage. Attempts to identify the bound molecule by mass spectrometry were unsuccessful. Strikingly, this density goes away when 4-BrC is bound.

Single-Channel Conductance in the Presence of a Potentiator. Single-channel currents flowing through individual channels expressed in BHK cells were identified at an extracellular pH of 7.5 using applications of 4-BrC to outside-out patches held at a constant membrane potential. The 4-BrC (30 μM) triggered a burst of sTeLIC activity, with a rising phase followed by a decay in the presence of the compound, synonymous with the desensitization shown in oocyte recordings (Figs. 5C and 7). All bursts displayed large current events such as those emphasized in Fig. 7. Inset. The single-channel current amplitude varied linearly with voltage, and reversed near 0 mV, as expected in the symmetrical NaCl solutions used. This indicated a single-channel conductance of 46 ± 2 pS (n = 4) between −80 and −20 mV. Current bursts from most patches in these conditions also displayed current steps with smaller amplitudes but which are much less frequent than the major 46-pS events, indicating conductance substates, which are not further analyzed here.

Inhibition by Divalent Cations but Not by Quaternary Ammonium Cations. The inhibitory effect of several divalent cations was evaluated with pH 8 activation, and the most potent one was found to be Zn²⁺ with an IC₅₀ of 3 ± 2 μM (Fig. 8A). Calcium and barium ions have similar potencies at 180 ± 60 μM and 170 ± 90 μM, respectively. A current trace showing the effect of increasing concentrations of Ca²⁺ ions is shown in Fig. 8B and the corresponding dose–response curves for all ions in Fig. 8C. All of the divalent cations tested showed completely reversible inhibition, as can be seen for Ca²⁺ in Fig. 8B, when long-enough wash times were used. To maintain membrane stability, 1 mM Mg²⁺ was present in all tests. From the few (n = 4) well-behaving oocytes tested without

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Fig. 4. Cationic selectivity of sTeLIC. Ion charge selectivity was evaluated from measurements of the reversal potential of the sTeLIC current using voltage ramps, with both an oocyte two-electrode voltage-clamp set-up (A and B) and a BHK path-clamp set-up (C and D). Simultaneously and equally diminishing the extracellular concentrations of Na⁺ and Cl⁻ (denoted [NaCl]) was counterbalanced with sorbitol. (A) Current-to-voltage plot of typical pH 8, from pH 5, elicited current, in 10 mM Hepes/NMDG, with 1 mM MgCl₂ solution. For each [NaCl] value, currents recorded with the solution at pH 5 were subtracted from currents recorded at pH 8 on the same oocyte. (B) Plot of the shift in reversal potential (Erev - Erev_ref), measured in experiments as shown in A, as a function of the ratio [NaCl]/[NaCl]ref. Erev_ref denotes the reversal potential measured with the reference concentration, 100 mM NaCl ([NaCl]ref). Data, given as mean ± SD, is smaller from three oocytes. The theoretical values of the shift in Erev for anionic and cationic channels, assuming perfect selectivity for Cl⁻ over Na⁺ (PNa/PP = 0), or for Na⁺ over Cl⁻ (PNa/PP = 0) are also shown. (C) Current-to-voltage plot of sTeLIC current recorded from a BHK cell, with 17, 34, or 170 mM [NaCl] (306, 272, or 0 mM sorbitol) outside, and 170 mM [NaCl] inside, in addition to 10 mM MgCl₂ and 10 mM Hepes/Na, at pH 7.5 in the presence of 30 μM 4-BrC. Average of three traces collected after desensitization was subtracted from the average of two traces near the maximal current, in each condition. (D) Plot of the reversal potential (Erev) measured in ion conditions as in C, as a function of [NaCl] outside. Erev values expected for various PNa/PP ratios are also plotted.

It should be mentioned that the difference maps of sTeLIC crystals that were devoid of 4-BrC also revealed a large blob of unexplained electron density in both 2Fo-DFc and mFo-DFc maps at the level of the intrasubunit 4-BrC binding site but closer

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Fig. 5. Modulation of sTeLIC response to pH by 4-BrC. (A) Typical electrophysiological traces in oocytes showing the potentiating effect of increasing concentrations of 4-BrC. Inset shows an enlarged view of 10-μM concentration application which details the sTeLIC response to the protocol used. (B) Corresponding dose–response curve of average ± 5D of normalized values with a nonlinear regression fit for 4-BrC at pH 7.5. The EC₅₀ of 21 ± 8 μM and Hill-slope of 2.5 ± 0.5 were found by fitting individual data and averaging the fit values from six oocytes. Values reported as mean ± SD. (C) Current traces obtained from a single oocyte showing the repeated prolonged application (220 s and 60 s) of 30 μM 4-BrC upon pH 8-elicited currents (60 s and 40 s, Left and Right traces, respectively) from pH 5, with a 5 min stimulation at pH 8 only in between (Center). Double lines (/) represent 2- and 12-min respective recording times. The effects observed in the Center and Right traces indicate that the strong 4-BrC-induced current decay observed in the Left trace is reversible, and corresponds to a desensitization.
divalent ions, the addition of Mg$^{2+}$ was found to inhibit pH 8 currents by $32 \pm 9\%$ and $93 \pm 8\%$ (1 mM and 30 mM, respectively, mean $\pm$ SD), indicating about 100-fold less potency than Ca$^{2+}$.

We used X-ray crystallography to locate binding sites for Ba$^{2+}$ in the sTeLIC structure due to the anomalous signal of Ba$^{2+}$. Two binding sites were found in the anomalous map using data collected at a longer wavelength from crystals soaked in BaCl$_2$. Two of the subunits are highlighted in yellow and cyan. 4-BrC is colored in red and depicted as sticks. The anomalous density map coming from the anomalous signal of a bromine atom is shown in a green mesh contoured at 5 $\sigma$. The Fourier difference 2mFo–DFc map around 4-BrC, contoured at 1 $\sigma$, is shown as a blue mesh.

Fig. 6. Crystal structure of the complex between sTeLIC and 4-BrC (PDB ID 6FLI). (A) Five symmetrical binding sites of 4-BrC in the sTeLIC ECD vestibule. Two of the subunits are highlighted in yellow and cyan. 4-BrC is colored in red and depicted as sticks. (B) Surface representation of the cavity forming the binding site of 4-BrC with the bound allosteric modulator shown in stick and ball representation. (C) Close-up view of the interaction between 4-BrC and sTeLIC. Residues interacting with 4-BrC are depicted as sticks. The anomalous density map coming from the anomalous signal of a bromine atom is shown in a green mesh contoured at 5 $\sigma$ and the Fourier difference 2mFo–DFc map around 4-BrC, contoured at 1 $\sigma$, is shown as a blue mesh.

Fig. 7. Identification of sTeLIC single channel current. Current trace from an outside-out patch hold at $-40$ mV, showing a burst of sTeLIC channels activity triggered by extracellular application of 4-BrC (30 $\mu$M) at pH 7.5. Channel activity rose and then decayed in the presence of the compound, as shown from return to all-channels-closed level (dotted line). An expanded trace of the rising phase is shown as inset, with the corresponding current data points histogram (counts per 0.02-pA interval), showing the current levels corresponding to zero to six channels open simultaneously, as indicated above the histogram.

Structural Analysis of Conserved Residues. As with other prokaryotic members of the pLGIC family, the Pro-loop lacks the disulfide bond conserved in the eukaryotic pLGICs’ Cys-loop. The Pro-loop interacts with the M2–M3 loop through the stacking interactions of F122 and P123 (Y119 and P120 in GLIC, and F119 and P120 in ELIC) and L247 (L246 in GLIC, and L255 in ELIC). The crucial salt bridge between R193 and DI25 is the equivalent of R192–D122 in GLIC (R198–D122 in ELIC), whereas the third partner (D32) of this salt bridge in GLIC is replaced by Q25 (T28 in ELIC) in the $\beta_1$–$\beta_2$ loop (SI Appendix, Fig. S12E). The stacking interaction of W162 on the hydrophobic side of R193 side chain is also conserved (W160 in GLIC and ELIC) (SI Appendix, Fig. S12D), as well as the primordial role of the Y198 side chain (Y197 in GLIC, Y203 in ELIC). Interestingly, E161 of sTeLIC is also making a salt bridge with R193, opposite from D125. E161 is well conserved in bacterial pLGICs but is absent in GLIC, where its role is fulfilled by D32 from $\beta_1$–$\beta_2$ loop.

Strikingly, there is a strong covariation in prokaryotic receptors (10 of 11 cases) of the presence/absence of a negatively charged residue at the positions equivalent to sTeLIC residues 25 and 161 (SI Appendix, Fig. S13). In this respect, GLIC’s covariation results
in a different loop F sequence than any other cationic pLGIC for which a structure is resolved, where the conserved short-sequence motif EW in bacteria (extended to GEW in eukaryotes) is changed to TGW. The W of this XEW motif is otherwise well conserved for cationic ion channels, but is absent in the anionic ones, where it is replaced by a conserved P(Q/S)F motif (SI Appendix, Fig. S12E).

At the ECD–TMD interface, the fundamental role of the conserved P205, at the end of the second turn of the M1 α-helix is confirmed in that it induces a characteristic kink that is further stabilized by an interaction of the carbonyl atom of M200 with the side chain of N240 from (α) subunit (SI Appendix, Fig. S14A). In addition, we noted the presence of salt-bridge R249–D246, that connects two adjacent M2–M3 loops, which is not conserved in other pLGICs, as well as R201–D257, somewhat conserved in prokaryotes, that links the pre-M1 region and the top of M3 from (+) subunit (SI Appendix, Figs. S13–S15).

Discussion

sTeLIC Shows Canonical Loop B, Loop C, and Loop F. The analysis of the known important loops for activation in pLGICs shows that the loop B conformation is “canonical” in sTeLIC, compared with eukaryotic receptors of known structures. With this additional prokaryotic pLGIC structure that conforms to the canonical conformation, the peculiarity of GLIC’s loop B conformations is more evident, marking GLIC’s loop B as an outlier (SI Appendix, Fig. S12B). Loop C in sTeLIC displays no cavity and is in close contact with the complementary subunit, as expected in an active form. It is still unknown what the effect of pH at the level of the agonist site is, although the presence of several histidines around loop C may indicate a possible mechanism for pH-sensing (SI Appendix, Fig. S12A), perhaps through a redistribution of electrostatic energies or dielectric relaxation (47). Loop F maintains the XEW sequence motif of many cationic pLGICs, where the glutamate (E161) has a strong sequence covariation with Q25, the equivalent position of D32 in GLIC, such that the interaction of E161 with the R192-D122 salt bridge is replaced by D32 in GLIC (where XEW is not conserved and replaced by TGW). E161’s role has also been highlighted in ELIC, with the studies of the chlorpromazine binding site (48). In the ELIC structure, loop F contributes to a Ba++ binding site identified in this closed form (35). From the sequence of loop F in sTeLIC, it may be speculated that a homologous binding site involving the side chains of loop F residues E159-E160 may be found in a closed form of sTeLIC, whereas no divalent cation binding site was identified in the present open structure. The assumption that loop F might also play a role in the Ca++ inhibition of sTeLIC by stabilizing a closed form, as shown in ELIC, is consistent with our observation that sTeLIC inhibition by divalent cations does not show the properties typical of an open-channel block (Fig. 8 D–F). If present at the level of loop F, a divalent cation would also block the alternative lateral cation pathway (see below).

sTeLIC Shares Many Features with ELIC. The sequence and structural motifs of sTeLIC, such as an amphipathic α-helix, absent in GLIC, are most similar to ELIC among the pLGICs (SI Appendix, Figs. S3 and S15). We propose that ELIC and sTeLIC are representative members of a subfamily of pLGICs, with specific characteristics of the pore, a possibility already envisaged for ELIC (41).

In addition to the sequence and tertiary structure similarity, sTeLIC belongs to the same gammaproteobacteria family and shares several functional similarities with ELIC. They are both cationic channels, and they are both positively modulated by the same ECD intrasubunit vestibule site (8). Moreover they share additional commonality through their inhibition by divalent ions (35), as well as a lack of inhibition by quaternary ammonium cations, such as TMA, TEA, or TPA (41). However, sTeLIC is activated at alkaline pH, whereas ELIC is activated by amine-containing compounds, such as GABA or cysteamine (49). Here we suggest that sTeLIC is indeed a second member of this subfamily family.
and its high-resolution structure, with a widely open pore, may provide a model for their open-channel conformation.

Comparison with Other Open Conformations. The widely open pore of sTeLIC matches very well the one of the α1-Gly-R (SI Appendix, Fig. S3A), while it is more open than in the GLIC structure at acidic pH. It has indeed been suggested from electron paramagnetic resonance experiments (50) that this GLIC structure may be only semiopen. This interpretation suggests that GLIC’s open-crystal form may represent an intermediate along the reaction path from closed to open forms, trapped because of crystallization conditions, or that the open form of GLIC and other cationic prokaryotic receptors with the same sequence covariations in loop F may have a different conformation. The extreme open form of sTeLIC’s pore may also result from the binding in both structures of either PEG 200 or 4-BrC to the PAM vestibule site.

Overall, the quaternary ECD structures of sTeLIC and the apparently open conformation of GLIC superimpose well after aligning the TMDs, with very little twist (tangential movement) and little bloom (radial movement) (SI Appendix, Fig. S3 B and E). The same comparison of sTeLIC and the open form of α1-Gly-R gives a similar result (SI Appendix, Fig. S3 C and F). However, superimposing sTeLIC and ELIC in the same way reveals both significant twist and bloom movement in the ECD (SI Appendix, Fig. S3 A and D), while the self-consistent factor between the atomic movement present in both of these two structures (but not in GLIC), appears to be shifted by one helical turn along its axis. In conclusion, the quaternary conformation of the ECD of sTeLIC is more similar to the apparently open forms of GLIC and α1-Gly-R than to the closed structures of ELIC, which places at odds the proposal that ELIC’s crystal structure might represent a “decoupled state” with a “relaxed” ECD and a “closed” TMD (40).

Specificities of the Ion Permeation Pathways. The cation pathway in the pore is well illustrated by crystallographic studies in the presence of Cs+ (34). However, this study also suggests that there might be an additional entry pathway for monovalent cations through a lateral door located at the interface between the ECD and the TMD (Fig. 3B), as observed in GABAa-R (18) as well as 5HT3A-R, based on molecular dynamics studies (51). Changing the electrostatics of E160 (loop F) and E28 (loop 2) is enough to reveal a lateral tunnel, as shown in SI Appendix, Fig. S16.

In addition, the presence of two constriction rings in the lumen of the ECD (especially at the level of loop Ω) is intriguing, but these constriction rings in the ECD are not uncommon among other members of the family (GluCra, 5HT3A-R). However, we have shown that the residues causing the constriction rings are not essential to sTeLIC function (Fig. 3C), and that the R86A mutation does not significantly alter the receptor conformation (SI Appendix, Fig. S7). They may play a role as an additional filter, or as a valve, forming a temporarily isolated chamber to regulate the flow of ions inside the lumen of the ECD.

To investigate a possible dynamic role of this region, we calculated low-frequency all-atoms normal modes using a recently developed nonlinear implementation in the framework of the Elastic Network Model (52) and looked for specific modes that could potentially open and close this filter or valve. We found several modes that indeed open the constriction ring in the ECD, either in a symmetrical fashion, mode 10, (Movies S1 and S2) or in a nonsymmetrical fashion (e.g., mode 18) (Movies S3 and S4). Interestingly, mode 10 keeps the pore widely open but displays an up-and-down movement of the TMD, mimicking the functioning of a (linear) “peristaltic pump.” Mode 18, on the other hand, involves the coordinated movement of two subunits, in opposition to the three remaining ones, and it actually closes and opens the pore periodically. The normal mode analysis clearly shows that the two constriction rings at the ECD are dynamic but extensive molecular dynamics simulations in a lipid environment would be needed to further assess a possible mechanism. Single-channel recording experiments of the receptors with mutations of K66 and R86, indicating their influence upon the conductance and ion selectivity of the permeation pathway, as well as the open-channel probability, would be required to show the influence of these constriction rings on permeation and gating.

Allosteric Potentiation of sTeLIC. A single binding site per subunit was present in the cocystal structure for 4-BrC, the most potent positive modulator identified in this study. Therefore, sTeLIC modulation by 4-BrC, which also occurs at a neutral pH and could therefore alternatively be described as a quasi-agonist effect, occurs almost certainly by binding to this vestibular intrasubunit site in the ECD.

If one directly compares the structures with and without 4-BrC, there is almost no difference (rmsd = 0.22 Å). However, comparison of the B-factors of the two structures, crystallized in the same conditions, shows a striking difference, with the 4-BrC bound form more ordered than the unbound form (SI Appendix, Fig. S17); this is not an effect of the resolution because the form with the lower-resolution dataset has the lowest B-factors. Thus, the binding of a 4-BrC apparently reduces the vibrations around the mean conformation (and the entropy), while the total electrostatic energy is also changed due to the extra carboxylate group, resulting in a different free energy of the PAM-bound state. This could result in a better anchoring of the β-sheets to the top of the ECD, a more stable ECD–TMD interface, and an overall stabilization of the open pore by 4-BrC binding. We note that simple elastic (coarse-grained) models of proteins have been shown to be able to reproduce this effect of modulation of the amplitude of thermal fluctuations around a mean structure (53).

A systematic examination of all available structures of pLGICs in the region of the PAM binding site shows that none of the anionic channels have such a vestibular cavity (GluCra, β3-GABAa-R, α1-Gly-R, α3-Gly-R), due to an extension of loop Ω that blocks its entry. In cationic receptors however, we found, in addition to ELIC and sTeLIC, a clear accessible equivalent cavity in the structure of the eukaryotic 5HT3A-R (SI Appendix, Fig. S18). Strikingly, this last receptor also has the same kind of restriction ring in the ECD through a lysine in loop Ω (SI Appendix, Fig. S6E).

Conclusion

In conclusion, sTeLIC is a bacterial pLGIC that is relatively easy to produce and purify for crystallization and whose functional and structural characterization reveals remarkable similarities with ELIC. It thus provides an apparently open-form model of ELIC that has proved up to now resistant to structural elucidation. Further studies will be needed to indicate whether or not sTeLIC forms, together with ELIC, a new subfamily of bacterial pLGICs (as suggested in ref. 41). Crystallizing and solving the structure of sTeLIC’s resting form would considerably help in assigning a functional state for the currently available closed form of ELIC, which is still uncertain. In addition, the identification of its PAM binding site and the finding that a similar cavity exists in the available structure of 5HT3A-R open the way for the rational design of drugs aimed at this human receptor.

Materials and Methods

sTeLIC was expressed in Escherichia coli with its N-terminal fused to MBP. The fusion protein was extract from E. coli membranes by detergent DDM. The recombinant protein was purified using amylase resin and further purified by Superose TM6 Increase 10/300 GL gel-filtration column. MBP was cleaved by thrombin and removed by a second round of gel filtration. Purified protein was pooled and concentrated to 12 mg/mL. Crystals were grown using hanging-drop method. The phase problem was solved by molecular replacement. Two-electrode voltage-clamp electrophysiology experiments were carried out on sTeLIC and its variants expressed in Xenopus laevis oocytes. Patch-clamp currents were recorded from cells of the BHK cell line driven to express sTeLIC, 1–2 d after DNA-calcium phosphate transfection. Full methods are provided in SI Appendix, SI Materials and Methods.
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