Proton transfer unlocks inactivation in cyclic nucleotide-gated A1 channels

Arin Marchesi, Manuel Arcangeletti, Monica Mazzolini and Vincent Torre

Neurobiology Sector, International School for Advanced Studies (SISSA), Trieste, Italy

Key points

- Desensitization and inactivation provide a form of short-term memory controlling the firing patterns of excitable cells and adaptation in sensory systems.
- Unlike many of their cousin K⁺ channels, cyclic nucleotide-gated (CNG) channels are thought not to desensitize or inactivate.
- Here we report that CNG channels do inactivate and that inactivation is controlled by extracellular protons.
- Titration of a glutamate residue within the selectivity filter destabilizes the pore architecture, which collapses towards a non-conductive, inactivated state in a process reminiscent of the usual C-type inactivation observed in many K⁺ channels.
- These results indicate that inactivation in CNG channels represents a regulatory mechanism that has been neglected thus far, with possible implications in several physiological processes ranging from signal transduction to growth cone navigation.

Abstract

Ion channels control ionic fluxes across biological membranes by residing in any of three functionally distinct states: deactivated (closed), activated (open) or inactivated (closed). Unlike many of their cousin K⁺ channels, cyclic nucleotide-gated (CNG) channels do not desensitize or inactivate. Using patch recording techniques, we show that when extracellular pH (pHₒ) is decreased from 7.4 to 6 or lower, wild-type CNGA1 channels inactivate in a voltage-dependent manner. pHₒ titration experiments show that at pHₒ < 7 the I–V relationships are outwardly rectifying and that inactivation is coupled to current rectification. Single-channel recordings indicate that a fast mechanism of proton blockage underlines current rectification while inactivation arises from conformational changes downstream from protonation. Furthermore, mutagenesis and ionic substitution experiments highlight the role of the selectivity filter in current decline, suggesting analogies with the C-type inactivation observed in K⁺ channels. Analysis with Markovian models indicates that the non-independent binding of two protons within the transmembrane electrical field explains both the voltage-dependent blockage and the inactivation. Low pH, by inhibiting the CNGA1 channels in a state-dependent manner, may represent an unrecognized endogenous signal regulating CNG physiological functions in diverse tissues.

(Received 10 September 2014; accepted after revision 28 November 2014; first published online 4 December 2014)

Corresponding author V. Torre: Neurobiology Sector, International School for Advanced Studies (SISSA), Via Bonomea no. 265-34136, Trieste, Italy; Email: torre@sissa.it

Abbreviations BS, binding site; CN, cyclic nucleotide; CNG, cyclic nucleotide-gated; VGIC, voltage-gated ion channel.

Introduction

Cyclic nucleotide-gated (CNG) channels conduct a Ca²⁺-permeable non-selective cation current mediating signal transduction in photoreceptor and olfactory sensory neurons (Hille, 1992; Kaupp & Seifert, 2002; Craven & Zagotta, 2006). Although CNG channels belong to the superfamily of voltage-gated ion channels (VGICs), they open in response to binding of cyclic nucleotides

A. Marchesi and M. Arcangeletti contributed equally to this work.
(CNs) and are only barely modulated by membrane voltage (Hille, 1992; Kaupp & Seifert, 2002; Yu et al. 2005; Craven & Zagotta, 2006; Mazzolini et al. 2010). CNG channel activity is also regulated by a variety of molecules and ions, including protons. Changes in extracellular hydrogen ion concentration ([H⁺]) occur in a variety of physiological and pathophysiological conditions, such as neuronal activity, ischemia and inflammation (Kellum et al. 2004; Isaev et al. 2008; Magnotta et al. 2012). In the retina, pH₄ follows a circadian rhythm (Dmitriev & Mangel, 2001), and pH changes could play a role in adaptation of the retinal response to different light intensities. Moreover, acidosis has been associated with changes in cell excitability in vascular tissues as well as in the CNS (Tolner et al. 2011; Pavlov et al. 2013), in which CNG channels are widely expressed (Zufall et al. 1997; Kaupp & Seifert, 2002; Leung et al. 2010; Lopez-Jimenez et al. 2012). Therefore, determining the effect of pH₄ on CNG channel gating has great clinical and physiological significance.

It is well established that extracellular protons inhibit CNG currents, although conflicting mechanisms have been proposed (Root & MacKinnon, 1994; Rho & Park, 2013; Morrill & MacKinnon, 1999; Martínez-François et al. 2010). For instance, the outward rectification observed at low pH₄ has been attributed to a voltage-dependent proton blockage (Rho & Park, 2013) and more recently to an enhancement of the wild-type (WT) channel inherent mild voltage-dependent gating (Martínez-François et al. 2010). Here we show that when pH₄ is decreased from 7.4 to 6 or lower, WT CNGA1 channels inactivate or desensitize. The term ‘desensitization’ usually refers to a mechanism of activity attenuation following sustained exposure to a chemical messenger, such as a neurotransmitter in ligand-gated channels, whereas inactivation is commonly used for VGICs. In the present investigation, mutagenesis and ionic substitution experiments indicated that the current decline is reminiscent of the usual C-type inactivation observed in Kᵣ channels and consequently we refer to the observed loss of conduction as inactivation. A simple kinetic model in which two protons bind within the selectivity filter provides a conceptual framework to rationalize both voltage-dependent proton block and inactivation. Inactivation thus represents a novel regulatory mechanism of CNG channels that has been neglected.

**Methods**

**Ethical approval**

All studies were approved by the SISSA’s Ethics Committee according to the Italian and European guidelines for animal care (d.l. 116/92; 86/609/C.E.). Oocytes were harvested from female *Xenopus laevis* frogs using an aseptic technique. All *X. laevis* surgeries were performed under general anaesthesia, obtained by immersion in a 0.2% solution of tricaine methane sulfonate (MS-222) adjusted to pH 7.4 for 15–20 min. Depth of anaesthesia was assessed by loss of the righting reflex and loss of withdrawal reflex to a toe pinch. After surgery animals were singly housed for 48 h. Frogs were monitored daily for 1 week post-operatively to ensure the absence of any surgery-related stress. Post-operative analgesics were not routinely used. Considering the simplicity of the procedure, the lack of complications, the effectiveness of anaesthetic regimens and reductions in the number of animals likely to occur compared to the number that would be required if only one surgery were permitted, multiple surgeries on a single animal were performed. Individual donors were used up to five times, conditional upon the relative health of an individual animal. Recovery time between oocyte collection from the same animal was maximized by rotation of the frogs being used. A minimum recovery period of 1 month was ensured between ovarian lobe resection from the same animal to avoid distress. Evidence of surgery-related stress resulted in an extended rest period based on recommendations from the veterinary staff. After the fifth terminal surgery frogs were humanely killed through anaesthesia overdose via 2 h of immersion in a 5 g l⁻¹ MS-222 solution adjusted to pH 7.4.

**Molecular biology**

The CNGA1 channel from bovine rods consisting of 690 amino acids was used (Kaupp et al. 1989). Selected residues were replaced as described (Becchetti et al. 1999) using the Quick Change Site-Directed Mutagenesis kit (Stratagene, La Jolla, CA, USA). Point mutations were confirmed by sequencing, using a LI-COR sequencer (4000 i; LI-COR Biosciences, Lincoln, NE, USA). cDNAs were linearized and were transcribed to cRNA *in vitro* using the mMessage mMachine kit (Ambion, Austin, TX, USA).

**Oocyte preparation and chemicals**

Mutant channel cRNAs were injected into *X. laevis* oocytes (‘*Xenopus express* Ancienne Ecole de Vernassal, Le Bourg 43270, Vernassal, Haute-Loire, France). Oocytes were prepared as described (Nizzari et al. 1993). Injected eggs were maintained at 18°C in a Barth solution supplemented with 50 μg ml⁻¹ gentamycin sulfate and containing (in mM): 88 NaCl, 1 KCl, 0.82 MgSO₄, 0.33 Ca(NO₃)₂, 0.41 CaCl₂, 2.4 NaHCO₃ and 5 Tris-HCl, pH 7.4 (buffered with NaOH). During the experiments, oocytes were kept in a Ringer solution containing (in mM): 110 NaCl, 2.5 KCl, 1 CaCl₂, 1.6 MgCl₂ and 10 Hepes, pH 7.4 (buffered with...
Proton transfer unlocks inactivation in CNGA1 channels. Inactivation was reduced was determined from the ratios τ. CHES was used. Solutions I–V indicating the (is the = n I–V is the voltage across the membrane, = 0.2% (relationships were clearly 9. If not otherwise indicated, data et al. 6a and 28 V 25.5 mV at 23 °C was used as the charge = B = was used and P HEPES was used and = P and 4 g at the indicated E is = G 1981) were recorded with a patch-clamp amplifier (Digidata 1440A; Axon Instruments), using a 10 kHz. Current signals were sampled with a 16-bit A/D converter (Digidata 1440A; Axon Instruments), using a sampling rate of 50 kHz. All other macroscopic current recordings were low-pass filtered at 1 kHz and sampled at 2.5 kHz if not otherwise indicated. Single-channel recordings were low-pass filtered at 2 kHz and sampled at 5 kHz.

Data analysis

Inactivation time constants (τ) were obtained from fitting cGMP activated currents with one-component exponential functions. Single-channel currents (i) were estimated from patches containing only one CNGA1 channel and fitting normalized all-point histograms with two-component Gaussian functions as previously described (Bucossi et al. 1997). The dependency of the fraction of unblocked channels (P_unblocked/P_max, Fig. 2I) on voltage at different pH_o was determined from the ratios of the normalized conductance G/G_o at the indicated pH and at pH_o = 9. If not otherwise indicated, data are presented as mean ± SEM, with n indicating the number of patches. Statistical significance for parametric analysis was determined using unpaired two-tailed t-test or single-variable ANOVA, as indicated. For pairwise comparisons, a Holm–Sidak test was used as post hoc test. P < 0.01 or P < 0.05 were considered significant, as indicated. Data analysis and figures were made with Clampfit version 10.1 (Molecular Devices, Sunnyvale, CA, USA) and Systat Software, Chicago, IL, USA). Kinetic models were generated and evaluated using MatLab 7.9.0 (MathWorks, Natick, MA, USA). The voltage-dependent association constant for protons shown in Fig. 6 was assumed to be of the form:

\[ X(V) = X(0)e^{\delta VF/RT} \]

where X(0) is the association constant at 0 mV, \( \delta \) is the valence (+1), V is the voltage across the membrane, \( \delta \) is the fraction of the membrane voltage at the binding site (the electrical distance) and \( F/RT = 25.5 \text{ mV at } 23^\circ \text{C} \).
The effect of extracellular acidification is best seen in outside-out membrane patches where different proton concentrations could be assessed during the same experiment (Fig. 1F). Mg\(^{2+}\) at 20 mM was used (indicated by solid red bars in Fig. 1F) to block CNG channels – which are continuously exposed to 1 mM cGMP present in the patch pipette – and estimate the leak current (Liu & Siegelbaum, 2000). Upon pH\(_o\) switch to 5, a current corresponding to approximately 20% of the current at pH\(_o\) 7.4 is observed (Fig. 1F). Notably, this current is not stable and exhibits a slow run-down as observed in inside-out patches at the same pH\(_o\) (Fig. 1E). The current decline is fully reversible when pH\(_o\) was reverted from 5 to 7.4 (Fig. 1F, right panel).

As inactivation appears to be stronger at negative potentials (Fig. 1E) and both inactivation and rectification depend on pH\(_o\) (Fig. 1B–E and G), the observed outward rectification (Fig. 2A) could arise from the voltage and pH dependency of the current decline. However, several observations suggest that this is not the case. First, while pH\(_o\) 5...
Proton transfer unlocks inactivation in CNGA1 channels (black circles) and macroscopic $i^+\rightarrow V$ and $i^-\rightarrow V$ equal to 7.3 and 0.41, respectively. $I^+\rightarrow V$ $k_0$ is shown at the right $V=−cGMP$ at $o\rightarrow V$. $I^+\rightarrow V$ $k_0$ indicates the unitary current. $I^+\rightarrow V$ $k_0$ flowing at $−200$ mV) at pH $δ_10$ and $9, 8, 7, 6, 5$ and $4$, respectively. The red lines represent simultaneous fits of all data. The current decline onset (Fig. 1G, green circles). Second, inactivation and rectification appear to have a different pH

Figure 2. Proton block and current rectification in WT CNGA1 channel. A and B, macroscopic current recordings in the inactivated state obtained 2 min after the addition of 1 mm cGMP at 0 mV elicited by voltage steps from $−200$ to $+200$ mV ($ΔV = 20$ mV) at pH 5 (A) and 7.4 (B). Leak and capacitative components were removed by subtracting from the cGMP-activated current those records obtained in response to the same voltage protocol but without cGMP. C, single-channel $i^\rightarrow V$ (black circles) and macroscopic $I^\rightarrow V$ relationships (red circles) obtained from noise analysis (D) and from the recording in A, respectively. The $I^\rightarrow V$ relationship was scaled to the $i$ flowing at $+180$ mV. D, stationary fluctuation analysis at the indicated positive (upper panels) and negative (lower panels) voltages. A linear regression fit throughout each dataset provided the indicated unitary current $i$. Currents were evoked by 30 ms prepulse held at $−200$ mV followed by test potentials ranging from $−180$ to $+200$ mV. E, single-channel current recordings at $+100$ mV at the indicated pHs. Amplitude histogram from recordings at pH 5 is shown at the right (grey area). Dashed brown and blue lines represent a two-component Gaussian fit to histograms obtained at pH 5 and 7.4, respectively. Black dashed lines indicate the 0 current level. C, Oo and O refer to the closed, blocked and open states respectively; $i$ indicates the unitary current. F, as in E but at $−100$ mV. G, macroscopic current recording in the inactivated state elicited by a voltage prepulse held at $+200$ mV followed by test potentials ranging from $−200$ to $+200$ mV ($ΔV = 20$ mV). No tail currents were observed at negative membrane potentials, suggesting a fast mechanism of proton block. Black dashed line indicates the 0 current level. H, a sketch of the energy landscape for a symmetrical two-barrier Woodhull model of block. $U_{ij}$, $k_{ij}$ and $δ$ indicate the heights of energy barriers, rate constants at 0 mV and the electrical distance, respectively. $i$, dependence of the fraction of unblocked channels (see Methods) on $V$ at different pHs (filled circles, open circles, filled triangles, open triangles, open squares and filled squares refer to pHs 9, 8, 7, 6, 5 and 4, respectively). The red lines represent simultaneous fits of all the data with the model shown in F with $k_{01}$, $k_{0-1}$, $k_{21}$, $k_{2-1}$ and $δ$ equal to $7.3 \times 10^7$ m$^{-1}$ s$^{-1}$, 40.8 s$^{-1}$, $8.9$ s$^{-1}$, $1.6 \times 10^7$ m$^{-1}$ s$^{-1}$ and 0.41, respectively. © 2014 The Authors. The Journal of Physiology published by John Wiley & Sons Ltd on behalf of The Physiological Society.
dependency: while a significant increase in rectification was already observed when pH_i is lowered from 8 to 7 (Fig. 1B, C, G), an appreciable inactivation was not detected until pH_i was lowered to 6 (Fig. 1B–D). Lastly, the voltage dependency of inactivation adds to the instantaneous rectification resulting in an enhancement of the steady-state current rectification (Fig. 1H). Thus, the voltage- and time-dependent loss of conductance is not at the origin of the outward rectification observed in the I–V relationship, and the two processes of rectification and inactivation could have different underlying molecular mechanisms.

**Outward rectification arises from voltage dependency of proton blockage**

The almost instantaneous rectification at pH_о 5 (Fig. 2A) was substantially relieved at the usual pH_о 7.4 (Fig. 2B) and could reflect obstruction of the permeation pathway by protons within the transmembrane electrical field. Indeed, at hyperpolarized membrane potentials, protons are expected to be driven into the channel pore, enhancing the block and resulting in an outward rectification. To confirm this mechanism, we estimated the single-channel current i by noise analysis of macroscopic currents at different voltages (Fig. 2D). Estimates of the corresponding i thus obtained are plotted as a function of voltage in Fig. 2C (black dots). As with the macroscopic I–V relationship (Fig. 2C, red dots), the i–V plot is outwardly rectifying and the two curves are almost overlapping. Indeed, the ratio of the macroscopic and unitary currents flowing at +180 and −180 mV (I_180/I_−180 and i_180/i_−180) were 4.5 and 3, respectively, suggesting that the voltage dependency of proton blockage is responsible for a large fraction of the observed rectification.

Estimates of single-channel currents based on stationary fluctuation analysis could be biased by filtering settings (Hille, 1992), and therefore we also attempted to measure single-channel events from membrane patches containing possibly only one channel. Such electrical recordings are shown in Fig. 2E and F and have been obtained at +100 (Fig. 2E) and −100 mV (Fig. 2F) at the usual pH_о 7.4 (right traces) and when pH_о was lowered to 5 (left traces). Proton elevation resulted in a reduction of the single-channel amplitude that was more prominent at negative voltages (Fig. 2F). At −100 mV, amplitude histograms indicate that the i reduced from...
−2.3 to −0.58 pA when pH₀ was lowered from 7.4 to 5 (Fig. 2F), and a substantial relief of block was observed at +100 mV (Fig. 2E, left trace). Indeed, the ratio of the single-channel conductance at +100 and −100 mV (γ⁺/γ⁻) is equal to 3.6 at pH₀ 5. These data suggest that a fast mechanism of proton blockage is the leading cause of the current rectification, a notion that is further substantiated by the lack of tail currents when classical

![Figure 4. Properties of inactivation in pore mutant channels](image)

**A** and **B**, current recordings during a voltage step at −60 mV at pH₀ 5 and 7.4, in symmetrical 110 mM Na⁺ for the WT (A) and for the E363A mutant (B) channels. Currents were evoked by 1 mM cGMP and scaled for comparison. **C** and **D**, inactivation time constant τ (C) and residual fractional current Iₛ/Iₚₑᵃᵏ (D) for the E363A mutant channel at pH₀ 5 and 7.4. An unpaired two-tailed t-test was performed to compare the two conditions. No statistically significant differences were found. **E**, current recording during a voltage step at −50 mV at pH₀ 7.4 in symmetrical Na⁺ for the E363A mutant channel. Currents were evoked by the application of 1 mM cGMP as indicated by solid black bars. The inset shows the fractional recovery Iₜ/Iₘₐₓ for the E363A mutant channel at pH₀ 7.4 (black circles) and WT channel at pH₀ 5 (green circles) plotted as a function of the cumulative interpulse interval. Data were fitted to a single exponential function providing the indicated time constants τ. **F**, current recordings during a voltage step at −60 mV in symmetrical 110 mM Na⁺ for E363A (black trace), WT (green trace), T360A (blue trace) and T359A (red trace) channels, at pH₀ 5. Currents were evoked by 1 mM cGMP and scaled for comparison. **G** and **H**, inactivation time constant τ (G) and residual fractional current Iₛ/Iₚₑᵃᵏ (H) for WT, E363A, T359A and T360A channels. ANOVA and Holm–Sidak post hoc tests were performed to compare each ion against each other (**P < 0.05; ***P < 0.01). **I**, macroscopic current recording in symmetrical 110 mM Na⁺, evoked by 1 mM cGMP and elicited by voltage steps from −200 to +200 mV (∆V = 20 mV) at pH₀ 5 for the T359A mutant channel. **J**, dependence of G/Gₚₑᵃᵏ on V for WT (black dots) and T359A mutant (red squares) channels at pH₀ 5. While abolishing desensitization (F), Thr359 replacement with an alanine does not affect voltage dependency of proton blockage (J). Dashed black lines in A, B, E and F indicate the 0 current level.
Like divalent cations, protons appear to reduce unitary currents by acting inside the conduction pathway of the channel (Dzeja et al. 1999; Seifert et al. 1999), and therefore we examined whether the Woodhull model (Woodhull, 1973) describes proton blockage (Fig. 2H). We assumed that H\(^+\) binds to a binding site (BS) located at a certain electrical distance \(\delta\) from the extracellular side of the membrane (Fig. 2H). Figure 2I illustrates the normalized \(G/G_{\text{max}}\) relationship recorded at different pH\(_{\text{m}}\), the shape of which could be described by the Woodhull model, yielding a \(\delta\) of \(-0.41\). In this framework, the slope of the conductance observed at negative voltages (Fig. 2I) reflects the punch-through of H\(^+\) from the pore. The model, however, does not satisfactorily fit the data at a pH\(_{\text{m}}\) equal to or lower than 5 (filled and open squares, respectively, in Fig. 2I), in which the predicted proton blockage is consistently larger than effectively measured. A very similar model has been described to explain extracellular divalent cation blockage and the blocking site is located at almost the same electrical distance \(\delta\) of 0.45 (Seifert et al. 1999). It is thus conceivable that protons and divalent cations compete for the same BS within the selectivity filter of CNG channels.

**The inactivation gate is located at the selectivity filter**

In K\(^+\) channels, the intimate relationship between the so-called C-type inactivation and the selectivity filter is illustrated by the substantial effects of the permeant ions on inactivation gating (López-Barneo et al. 1993; Starkus et al. 1997). Therefore, we examined whether ionic substitution could also affect the extent and the rate of inactivation in CNGA1 channels. Current decline was observed in the presence of Na\(^+\), NH\(_4\)\(^+\)

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**Figure 5. Voltage dependency of inactivation in WT CNGA1 channel**

*1. Current recordings at pH\(_{\text{m}}\) 5 from the same inside-out patch at the indicated membrane potentials in symmetrical Na\(^+\). Currents were evoked by 1 mM cGMP. Dashed line indicates the 0 current level. B. Time course of inactivation \(I_{\text{f}}/I_{\text{max}}\) at pH\(_{\text{m}}\) 5 for the indicated membrane potentials from experiments as in A where sampling points were averaged every 500 \(\mu\)s. Each current decay was obtained from at least three independent experiments. \(I_{\text{max}}\) was measured at the peak of the cGMP current. C. Dependence of fractional inactivation \((1 - I_{\text{f}}/I_{\text{max}})\) at +60 mV (white dots) and −60 mV (black dots) on pH\(_{\text{m}}\). \(I_{\text{peak}}\) and \(I_{\text{ss}}\) were measured at the beginning and termination of the cGMP pulse, respectively. Blue, green and red lines refer to simultaneous fits of all the data with schemes i, ii, and iii shown in Fig. 6A, respectively. For scheme i association constant \(B(0)\) was \(1.59 \times 10^{4} \text{ m}^{-1}\) (pKa 4.2). The electrical distance \(\delta_{B}\) was 0.37. The forward (i\(_{\text{f}}\)) and backward (i\(_{\text{b}}\)) inactivation rate constants were 0.41 and 0.05 s\(^{-1}\), respectively. For scheme ii association constants \(B(01)\) and \(B(02)\) were \(2.8 \times 10^{4} \text{ m}^{-1}\) (pKa 5.44) and \(2.5 \times 10^{4} \text{ m}^{-1}\) (pKa 4.4), respectively. The electrical distance \(\delta_{B}\) was 0.22. The forward (i\(_{\text{f}}\)) and backward (i\(_{\text{b}}\)) inactivation rate constants were 0.51 and 0.035 s\(^{-1}\), respectively. For scheme iii association constants \(B(0), B(01)\), and \(B(02)\) were \(1.59 \times 10^{4} \text{ m}^{-1}\) (pKa 4.2), \(1.25 \times 10^{5} \text{ m}^{-1}\) (pKa 6.1) and \(5 \times 10^{4} \text{ m}^{-1}\) (pKa 4.7), respectively. The electrical distances \(\delta_{B}\) and \(\delta_{B}\) were 0.37 and 0.45, respectively. The forward (i\(_{\text{f}}\)) and backward (i\(_{\text{b}}\)) inactivation rate constants were 0.41 and 0.05 s\(^{-1}\), respectively.*
and methylammonium (MA⁺) (Fig. 3A), although inactivation was appreciably slower in the presence of NH₄⁺ (τ = 10.5 ± 1.7 s, n = 5) compared to Na⁺ (τ = 7.5 ± 1.2 s, n = 5) and MA⁺ (τ = 7.2 ± 1.2 s, n = 3) conditions (Fig. 3B). Moreover, in the presence of either NH₄⁺ or MA⁺ ions, current decline was significantly less complete (Fig. 3C). When Na⁺ ions were replaced with Rb⁺ and Cs⁺, inactivation properties did not differ among the different conditions (Fig. 3A–C). Thus, the pore occupancy of specific ions appears to be one of the microscopic factors controlling inactivation. Recently we showed that the presence of large cations such as Rb⁺ and Cs⁺ in the bathing medium inhibits the Na⁺ inward current by interacting with Thr359 and Thr360 at the intracellular mouth of the selectivity filter. Indeed, the ring of these threonines contributes to an ion binding site at the intracellular entrance of the selectivity filter, and cations present in the intracellular medium could potentially interact with this site and affect channel gating and permeation (Marchesi et al. 2012). Therefore, we investigated whether occupancy of this binding site by certain ions compared to others could affect inactivation properties. When NH₄⁺ ions in the bathing medium were replaced by an equimolar amount of Na⁺, the NH₄⁺ current not only appeared to be inhibited by the presence of intracellular Na⁺ as the NH₄⁺ peak current reduced by 32% in the presence of intracellular Na⁺ (I(NH₄⁺)/I(Na⁺) = 0.68 ± 0.06; n = 4), but inactivation also proceeded faster (τ = 5.38 ± 0.98 s, n = 4) and was more complete (Fig. 3D). It is therefore conceivable that Na⁺ occupancy of this intracellular site, signalled by the NH₄⁺ current blockage, prompted channel collapse towards the non-conductive, inactivated state.

Previous studies identified the Glu363 residue at the outer mouth of the selectivity filter as the major proton and divalent cation target (Root & MacKinnon, 1993, 1994; Eismann et al. 1994; Rho & Park, 2013; Morrill & MacKinnon, 1999). Therefore, we evaluated whether besides current rectification, proton binding to Glu363 also controls the time- and voltage-dependent loss of conductance discussed earlier. Similar to WT channels when pHₒ was reduced to 5, mutant E363A channels inactivated at the usual pHₒ of 7.4 (Fig. 4A, B, E), as previously described (Mazzolini et al. 2009). Moreover, the current decline observed at pHₒ 7.4 and pHₒ 5 is almost identical in this mutant (Fig. 4B); the inactivation time constant τ is equal to 3.1 ± 0.1 and 2.9 ± 0.2 s, respectively, and the residual steady-state current to 0.04 ± 0.01 and 0.05 ± 0.01 (Fig. 4C, D; n ≥ 4). Thus, neutralization of Glu363 abolishes the pH dependency of inactivation, indicating that proton binding to Glu363 is also responsible for the slow current decline. Remarkably, compared to the WT channel, the onset rate of current decline was faster (Fig. 4F, G) and inactivation was more pronounced in the E363A mutant channel (Fig. 4F, H), while the time course of recovery did not significantly differ in the two channels (Fig. 4E, inset). Inactivation kinetics is from two to three orders of magnitude slower than the gating transitions: indeed CNGA1 channels open in a few milliseconds in response to cGMP concentration jumps (Nache et al. 2006). Relying on this separation of timescales, the onset rate of inactivation depends on both the forward and the backward inactivation rate constants, while the recovery process depends solely on the backward rate constant. As the current decline onset in the E363A mutant is faster but recovery from this effect is not different from the WT channel, the mutation appears to affect only the forward inactivation rate constant, i.e. mutation of Glu363 destabilizes the open conformation of the pore.

To further substantiate the notion that the occupancy of an ion binding site in close proximity to the central cavity controls inactivation, we studied the current decline in Thr359 and Thr360 mutant channels (Marchesi et al. 2012). Inactivation kinetics for WT, T359A and T360A at pHₒ 5 are shown in Fig. 4F. Current decline developed significantly more slowly in the T360A mutant channel (τ = 19.0 ± 1.9 s, n = 3, Fig. 4G) compared to the WT channel (τ = 7.5 ± 1.2 s, n = 5, Fig. 4G). When Thr359 at the intracellular entrance of the selectivity filter was mutated to an alanine, a significant inactivation was not observed within 20 s (Fig. 4F, H). The lack of current decline observed in T359A channels is unlikely to arise from a decreased affinity to protons, as the outward rectification at pHₒ 5 is very similar in WT and T359A channels (Fig. 4I, J).

Voltage dependency of inactivation is expected from proton binding within the electrical transmembrane field

We studied the effect of membrane potential on inactivation kinetics by eliciting currents in response to 1 mM cGMP at different voltages (Fig. 5A). Two key differences among these records are the amplitude of steady-state currents and the rate of inactivation: the former increases as the membrane is made more depolarized, while the latter decreases. This finding is best illustrated in Fig. 5B, in which current decays obtained at different voltages were normalized to the current measured and compared immediately after addition of cGMP. Fitting the time course of inactivation at +70 and −70 mV with a single exponential yields a τ of inactivation equal to 16.3 and 6.7 s, respectively, and a residual steady-state current (I/Iₘₐₓ) equal to 0.63 and 0.27. The I–V relationships measured immediately after exposure to cGMP (filled circles) and after development of current decline (open circles) are shown in Fig. 5C and had a different degree of rectification. Indeed, at steady state,
the outward rectification increased due to the additional contribution of the inactivation (Fig. 5C). Finally, the pH dependency of inactivation at +60 mV (open circles) and −60 mV (solid circles) is shown in Fig. 5D. The fraction of channels residing in the inactivated state appears to be larger at negative membrane potentials and low pH (Figs 1H and 5D).

The slow time course of current decline described here might reflect a reduced accessibility of Glu363 carboxylate to extracellular protons. Indeed, previous structure–function studies suggested that this residue is not directly accessible to extracellular solvent (Sun et al. 1996; Mazzolini et al. 2009; Martínez-François et al. 2009). Yet, this simple interpretation is hard to reconcile with several experimental observations. First, according to this idea, the disruption of the proton binding site should have resulted in a non-inactivating phenotype at pH 5. This is in contrast to the finding that neutralization of Glu363 results in inactivating currents at the usual pHo of 7.4 (Fig. 4E). Indeed, the inactivation onset and recovery in the E363A mutant are still slow and comparable to those observed in the WT channel (Fig. 4E, F). Second, recovery from inactivation is fast – less than 1 s as no obvious kinetics could be resolved – when pHo is reverted from 5 to 7.4 in outside-out patches (Fig. 1F, right panel). If the slow current run-down reflected the low accessibility of Glu363 to protons, then one would have expected a similar slow recovery upon pHo switch to 7.4. Therefore, we suggest that current decline reflects proton-induced conformational changes, of which the kinetics are limited by a transition downstream from protonation. In this view, protons first bind

![Diagram](image_url)

**Figure 6. Proposed mechanism for proton action in CNGA1 channels**
Proton transfer unlocks inactivation in CNGA1 channels

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Desensitization and inactivation are widespread phenomena in ion channels, in which they modulate the amplitude, duration and frequency of electrical signalling within and between cells. Unlike many of their cousin K⁺ channels and other membrane receptors, CNG channels do not desensitize or inactivate at physiological pH when a network of chemical interactions between residues in the pore region is maintained (Mazzolini et al. 2009). We have shown that in response to changes of extracellular pH, CNG channels also inactivate, representing a possible novel regulatory mechanism that has so far been neglected.

Previous studies suggested the presence of multiple H⁺-binding sites in the CNG channel pore. Single-channel current analysis in olfactory CNG channels indicated that the binding of two protons is required to model the proton monovalent current inhibition (Root & MacKinnon, 1994). Moreover, Seifert et al. (1999, fig. 5B, C) showed that a Hill coefficient of ~2 describes the pH dependency of divalent cation blockage in CNGA1 channels, suggesting the binding of at least two protons. In agreement with these earliest observations we were unable to describe the experimental data when the binding of only one proton was considered (Fig. 5). Similarly, a linear scheme assuming two equivalent proton binding sites (Fig. 6Aii) failed to satisfactorily reproduce the experimental data as it overestimates the current inhibition observed at low pHo (compare Figs 1F and 6B, green trace). Therefore, we evaluated a more general kinetic scheme in which two non-equivalent and non-independent proton binding reactions were modelled (Fig. 6Aiii). This model appears to adequately describe key features of the reported current decline, such as the time course of inactivation (Fig. 5B, red lines), its pH dependency (Fig. 5D, red lines), the instantaneous and steady-state rectification (Fig. 5C, red lines) and the fractional current blockage observed at pHo 5 (Fig. 6B, red trace). As we were unable to resolve the kinetics of H⁺ unbinding in outside-out patches (Fig. 1F) as well as the kinetics of proton blockage from voltage jump experiments (Fig. 2G), the proton binding reactions are not likely to contribute to the observed slow current relaxation kinetics and the two binding reactions are therefore assumed to be diffusion limited in this model. Once protons have bound, the slow current decline reflects the very slow inactivation rate constants, which are the rate-limiting reactions in the scheme. Indeed, the time courses of the occupancies for the conductive states (O + 1½O, red lines) and the inactivated states (1H1 + 1½H1, brown lines) are symmetrical (Fig. 6C), yielding at −50 mV the same time constant τ of 6.5 s. Thus, we propose that the non-equivalent binding of two protons within the transmembrane electrical field underlies voltage-dependent blockage and inactivation.

It has also been suggested that voltage-dependent gating contributes to the rectification of the macroscopic currents at very low pHo (pHo = 4) as a consequence of Glu363 side chain titration (Martinez-Franco et al. 2010). Based on noise analysis (Fig. 2C, D), single-channel recordings (Fig. 2E, F) and voltage-jump experiments (Fig. 2G), this mechanism does not appear to significantly contribute to the outward rectification observed at higher pHo (pHo ≥ 5) and has therefore not been included (Fig. 6A).

How could the proposed kinetic mechanism be interpreted in the light of the available structural and functional data? The crystal structure of a recently solved bacterial channel mimicking the CNG channel pore (Derebe et al. 2011) suggests that Ca²⁺ ions are chelated by backbone carbonyl oxygens and not by the Glu363 side chain, which is engaged in hydrogen bonding interactions with its neighbouring residues and buried underneath the external surface of the protein. The authors of this study suggested that the Glu363 carboxylate negative charge could perturb the electron shell distribution along the backbone of the filter residues, making certain carbonyl oxygen atoms, such as those of Gly65, more electronegative and suited to Glu363 within the electrical transmembrane field, quickly blocking ion conduction and destabilizing the pore, which subsequently undergoes a slow structural collapse leading to a virtually closed channel. This molecular picture is captured by the simple linear scheme shown in Fig. 6Ai, involving transitions among an open, blocked and inactivated state, which could successfully describe the pH and voltage dependency of inactivation, yielding a pKₐ of 4.2 (Fig. 5B, D blue lines). Although we could not rule out the possibility that membrane voltage might affect the conformation of charged residues and/or structures located within the electrical field as suggested for bacterial K⁺ channel KcsA (Cordero-Morales et al. 2006), the properties of the observed current decline are well accounted by the voltage-dependent concentration of H⁺ within the electrical field alone, without the necessity of taking explicitly into account a voltage-dependent conformational change. The fraction of the membrane voltage at the binding site thus obtained (δ ~0.37) is consistent with the expected location of Glu363 within the electrical field (Contreras et al. 2010). This scheme, however, appears to be critically deficient in reproducing the instantaneous and steady-state I–V relationship (Figs 5C and 6B, blue lines). The basis of this shortcoming is the different pH dependency of proton blockage and inactivation. Indeed, proton blockage has been associated with an apparent acidic dissociation constant (pKₐ) between 6 and 7 in several reports (Root & MacKinnon, 1994; Seifert et al. 1999). It is therefore possible that the binding of at least two protons underlines proton blockage and inactivation in CNGA1 channels.

Discussion

Desensitization and inactivation are widespread phenomena in ion channels, in which they modulate the amplitude, duration and frequency of electrical signalling within and between cells. Unlike many of their cousin K⁺ channels and other membrane receptors, CNG channels do not desensitize or inactivate at physiological pH when a network of chemical interactions between residues in the pore region is maintained (Mazzolini et al. 2009). We have shown that in response to changes of extracellular pH, CNG channels also inactivate, representing a possible novel regulatory mechanism that has so far been neglected.

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for Ca\(^{2+}\) binding (Derebe et al. 2011). These observations are consistent with several recent structure–function studies, suggesting that interactions between the Glu363 side chain and residues in the P-helix are necessary for normal gating in CNG channels (Mazzolini et al. 2009; Martínez-Francos et al. 2009, 2010). In this view proton blockage could arise from the high-affinity binding to the main chain carbonyl oxygens within the selectivity filter (Fig. 6D, pK\(_a\) = 6.1). A second, low-affinity binding site for protons could be constituted by the Glu363 side chain (pK\(_a\) = 4.2), titration of which results in the neutralization of the negatively charged carboxylate and in the weakening of key interactions anchoring the selectivity filter to the surrounding channel moiety (Fig. 6D) (Mazzolini et al. 2009; Martínez-Francois et al. 2009, 2010). After Glu363 protonation, glycine carbonyl oxygen specificity for protons and divalent cations is lost – the pK\(_a\) of the proton high-affinity binding site reduces from 6.1 to 4.7 – and thus a substantial relief from blockage is observed at low extracellular pH\(_o\) (Fig. 2I). It then follows a slow pore collapse toward a non-conductive, inactivated state. These observations suggest that the two binding sites are non-independent, and explain why neutralization of the same residue (Glu363) affects both proton blockage (Root & MacKinnon, 1994) and the pH dependency of inactivation (Fig. 4).

One of the hallmarks of the C-type inactivation is its intimate dependence on the permeant ion. Indeed, it has long been known that the rate and extent of C-type inactivation is governed by the occupancy of an ion binding site near the external mouth of the pore via a foot-in-the-door mechanism (Yellen, 1998; Kurata & Fedida, 2006). Our results indicate that such a relationship between the permeant ion and inactivation also exists in CNG channels (Fig. 3), although the underlying mechanism appears to be somewhat different. In fact, inactivation appears to be primarily modulated by the nature of the intracellular cation, being faster in the presence of intracellular Na\(^+\) (Fig. 3D). Furthermore, inactivation is abolished when the intracellular ion binding site is disrupted (Fig. 4F), thus suggesting that the occupancy of an ion binding site possibly equivalent to the fourth ion binding site within the selectivity filter (S4 site) of the K\(^+\) channel might control the rate and extent of inactivation in CNG channels. Therefore, it is conceivable that the pore collapse triggered by the protonation of Glu363 implies fine conformational changes also of the intracellular vestibule involving the two threonines Thr359 and Thr360. Although the molecular basis underpinning the C-type inactivation is still of debate (Devaraneni et al. 2013; Armstrong & Hoshi, 2014), according to the predominant view inactivation is associated with a conformational change close to the external mouth of the K\(^+\) channel pore (Yellen, 1998; Kurata & Fedida, 2006; Hoshi & Armstrong, 2013). Nonetheless, rearrangements near the intracellular face of the selectivity filter have been implied as well. Indeed, Posson et al. (2013) demonstrated that the binding affinity of certain intracellular blockers depends on the conformational state of the selectivity filter (open-conductive versus closed-inactivated). The authors of this study argued that the observed state-dependent affinities reflect a conformational change at the intracellular face of the selectivity filter upon inactivation. These rearrangements close to the intracellular end of the selectivity filter could be more pronounced during the CNG channel inactivation process as they may reflect the allosteric connection between the CN-binding domain and the selectivity filter gate. In this view, residues Thr359 and Thr360 might play an important role in the observed current decline. Overall, inactivation appears to be associated with a local remodelling confined to the selectivity filter and neighbouring regions in CNG as well as K\(^+\) channels (Roncaglia & Becchetti, 2001; Kurata & Fedida, 2006; Mazzolini et al. 2009).

In the present study, we demonstrate that the outward rectification observed at pH\(_o\) ≥ 5 at stationary conditions (i.e. after a steady exposure to cGMP for 1–2 min) is mainly caused by asymmetries in the unitary current (Fig. 2). This finding is consistent with several earlier studies suggesting that proton elevation is associated with the stabilization of low conductance states primarily at negative voltages (Root & MacKinnon, 1994; Rho & Park, 2013; Morrill & MacKinnon, 1999). We propose that a Woodhull model of block, alongside divalent cation blockage (Seifert et al. 1999), underlies proton block in CNG channels. This is directly opposite to the conclusions reached by a recent report suggesting that the outward rectification is due to an inherent channel voltage-dependent gating (Martínez-Francois et al. 2010). The reasons for these discrepancies are not clear. Several of the Glu363 mutants display voltage-dependent gating in addition to inactivation (Bucossi et al. 1996; Martínez-Francos et al. 2009). It is thus likely that voltage gating additively contributes to the observed rectification when extracellular pH is further lowered to 4 and slow activation kinetics are observed (Martínez-Francois et al. 2010). Determining the extent that such a mechanism contributes to the outward rectification at very low pH\(_o\) will require further experimentation. Nonetheless, demonstration of common gating features (inactivation and voltage gating) as the result of pore mutation and protonation further illustrates the gating role of the outer pore in CNG channels.

Changes in pH\(_o\) can arise in a variety of physiological and pathophysiological conditions, such as neuronal activity, ischaemia and inflammation (Kellum et al. 2004; Isaev et al. 2008; Magnotta et al. 2012). Low pH
acts as a negative feedback mechanism that inhibits the CNGA1 channel in a state-dependent manner and may represent an unrecognized endogenous signal regulating CNG physiological functions in diverse tissues.

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**Additional Information**

**Competing interests**

The authors declare that there are no competing interests.

**Author contributions**

M.A. and A.M. performed experiments on oocytes. M.M. performed mutagenesis. All authors participated in designing experiments, and analysing and interpreting the data. A.M. wrote the paper, which was revised by all authors in collaboration. A.M. and V.T. supervised the project. All authors approved the final version of the manuscript.

**Funding**

We acknowledge the financial support of the following projects within the Seventh Framework Programme for Research of the European Commission: the SI-CODE project for Future and Emerging Technologies (FET) no. FP7 - 284553, the FOCUS Project no. FP7-ICT-270483 and the NEUROSCAFFOLDS Project no. 604263.