

Functional asymmetry of transmembrane segments in nicotinic acetylcholine receptors

Jörg Grandl · Christophe Danelon · Ruud Hovius · Horst Vogel

Received: 13 March 2006 / Revised: 22 May 2006 / Accepted: 8 June 2006 / Published online: 13 July 2006
© EBSA 2006

Abstract Nicotinic acetylcholine receptors are heteropentameric ion channels that open upon activation to a single conducting state. The second transmembrane segments of each subunit were identified as channel-forming elements, but their respective contribution in the gating process remains unclear. Moreover, the detailed impact of variations of the membrane potential, such as occurring during an action potential, on the transmembrane domains, is unknown. Residues at the 12' position, close to the center of each second transmembrane segment, play a key role in channel gating. We examined their functional symmetry by substituting a lysine to that position of each subunit and measuring the electrical activity of single channels. For 12' lysines in the α , γ and δ subunits rapid transitions between an intermediate and large conductance appeared, which are interpreted as single lysine protonation events. From the kinetics of these transitions we calculated the pK_a values of respective lysines and showed that they vary differently with membrane hyperpolarization. Respective mutations in β or ϵ subunits gave receptors with openings of either intermediate or large conductance, suggesting extreme pK_a values in two open state conformations. The results

demonstrate that these parts of the highly homologous transmembrane domains, as probed by the 12' lysines, sense unequal microenvironments and are differently affected by physiologically relevant voltage changes. Moreover, observation of various gating events for mutants of α subunits suggests that the open channel pore exists in multiple conformations, which in turn supports the notion of functional asymmetry of the channel.

Abbreviations

nAChR	Nicotinic acetylcholine receptor
TM1/TM2	First/second transmembrane segment
GFP	Green fluorescent protein

Introduction

The nicotinic acetylcholine receptor (nAChR) is a paradigm of ligand-gated ion channels (Absalom et al. 2004). Muscle-type nAChR are heteropentamers with a subunit stoichiometry $(\alpha 1)_2, \beta 1, \delta, \gamma$ (the γ subunit being replaced during maturation by the ϵ subunit) and two binding sites for the natural agonist acetylcholine, located at the interfaces between the α and their adjacent subunits (α/δ , α/γ or α/ϵ). Upon ligand binding the receptor switches from its closed conformation in apparently one single step to the open state (Maconochie et al. 1995). How the implied structural rearrangements within the asymmetric structure are realized in detail is still unclear (Grutter and Changeux 2001).

J. Grandl · C. Danelon · R. Hovius · H. Vogel (✉)
Laboratory of Physical Chemistry of Polymers and Membranes, Ecole Polytechnique Fédérale de Lausanne, Faculté des Sciences de base, Station 6, Lausanne 1015, Switzerland
e-mail: horst.vogel@epfl.ch

Present Address:

J. Grandl
Department of Cell Biology, The Scripps Research Institute, La Jolla, CA 92037, USA

The conformational changes are not necessarily the same for all subunits (Dahan et al. 2004; Grosman and Auerbach 2000) and structural data from electron microscopy from *Torpedo californica* nAChR propose that rotational movements of the α subunits promote channel gating (Miyazawa et al. 2003; Unwin 2005).

From various experiments it is known, that in nAChR the highly conserved second transmembrane segments (TM2) of each subunit (Fig. 1) form the membrane spanning part of the channel (Akabas et al. 1994; Hucho et al. 1986). Lysine substitution along the δ -TM2 in mouse nAChR has recently been shown to locally probe the microenvironment of the pore (Cymes et al. 2005), and protonation of the ϵ -NH₂ group reduced channel conductance. Within a channel pore the proton exchange reaction is greatly slowed down so that individual events become measurable compared to bulk conditions, where such processes happen on time scales shorter than the typical time-resolution of patch-clamp experiments. The degree of channel blocking reflects, due to repulsion of equal charges, the distance of the lysine to the ion pathway. Gating rate constants directly mirror the protonation kinetics. They can hence be used to calculate local pK_a values of single lysines in the channel, which might differ from the pK_a value of lysine in a bulk environment (~10.2).

Here, we extended this approach to the 12' positions of TM2 in all subunits. Moreover, we studied the effect of engineered lysines on single-channel properties as a function of membrane potential to sense structural and functional differences between individual subunits.

Materials and methods

Site-directed mutagenesis

The single-point mutations α T254K, β T265K, γ T263K δ S268K and ϵ T263K were produced using cDNA coding for subunits of human muscle-type nAChR

using the QuikChange Site-Directed Mutagenesis Kit (Stratagene, La Jolla, CA, USA) and their correctness was confirmed by sequencing (MWG, Germany).

Cell preparation

Human embryonic kidney cells were cultured in D-MEM:F-12 medium (Invitrogen, Basel, Switzerland) containing 2.5% fetal calf serum (Sigma, Buchs, Switzerland) at 37°C in a humidified 5% CO₂ atmosphere. Cells were transiently transfected using the Effectene Transfectent Reagent (Qiagen, Hombrechtlikon, Switzerland). A total of 0.2 μ g cDNA coding for the α , β , δ and ϵ or γ subunits of the human muscle nAChR was mixed with green fluorescent protein (GFP) cDNA in the ratio of 2:1:1:1:1 (α : β : ϵ or γ : δ :GFP) or in equal amounts for experiments combining mutated and native α subunits and applied to a 35 mm culture dish with 2 ml medium. Experiments were performed on the adult-type receptor, for its larger single-channel conductance (59 vs. 39 pS for fetal-type receptors) and therefore its higher signal-to-noise ratio in single-channel measurements (Newland et al. 1995), except for measurements on the γ T12'K mutant. Experiments were carried out 1–3 days after transfection upon transferring cells to a solution containing: NaCl 147 mM, glucose 12 mM, HEPES 10 mM, KCl 2 mM, MgCl₂ 1 mM (all Sigma), pH adjusted to 7.4 with NaOH.

Lysine substitutions in α subunits

To investigate the effects of lysine substitution in each of the two α subunits (α_δ or α_ϵ), three sets of single-channel experiments were performed upon transfection of DNA coding for: (i) the adult wild-type receptor (α , β , ϵ , δ) in the ratio 2:1:1:1, (ii) only the mutated version of the α subunit (α T12'K, β , ϵ , δ) in the ratio 2:1:1:1, yielding receptors containing the mutation in both α subunits (further: 2 \times α T12'K), (iii) both mutated and wild-type nAChR (α T12'K, α , β , ϵ , δ) in

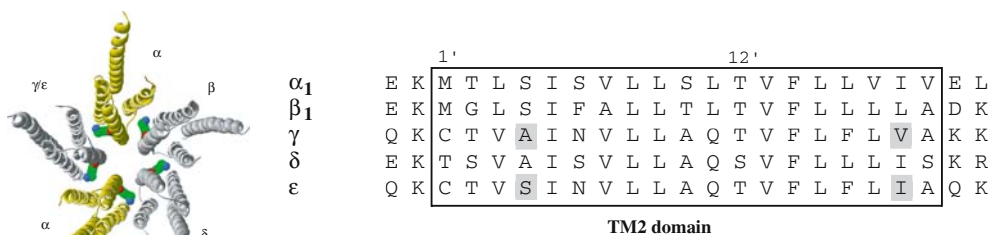
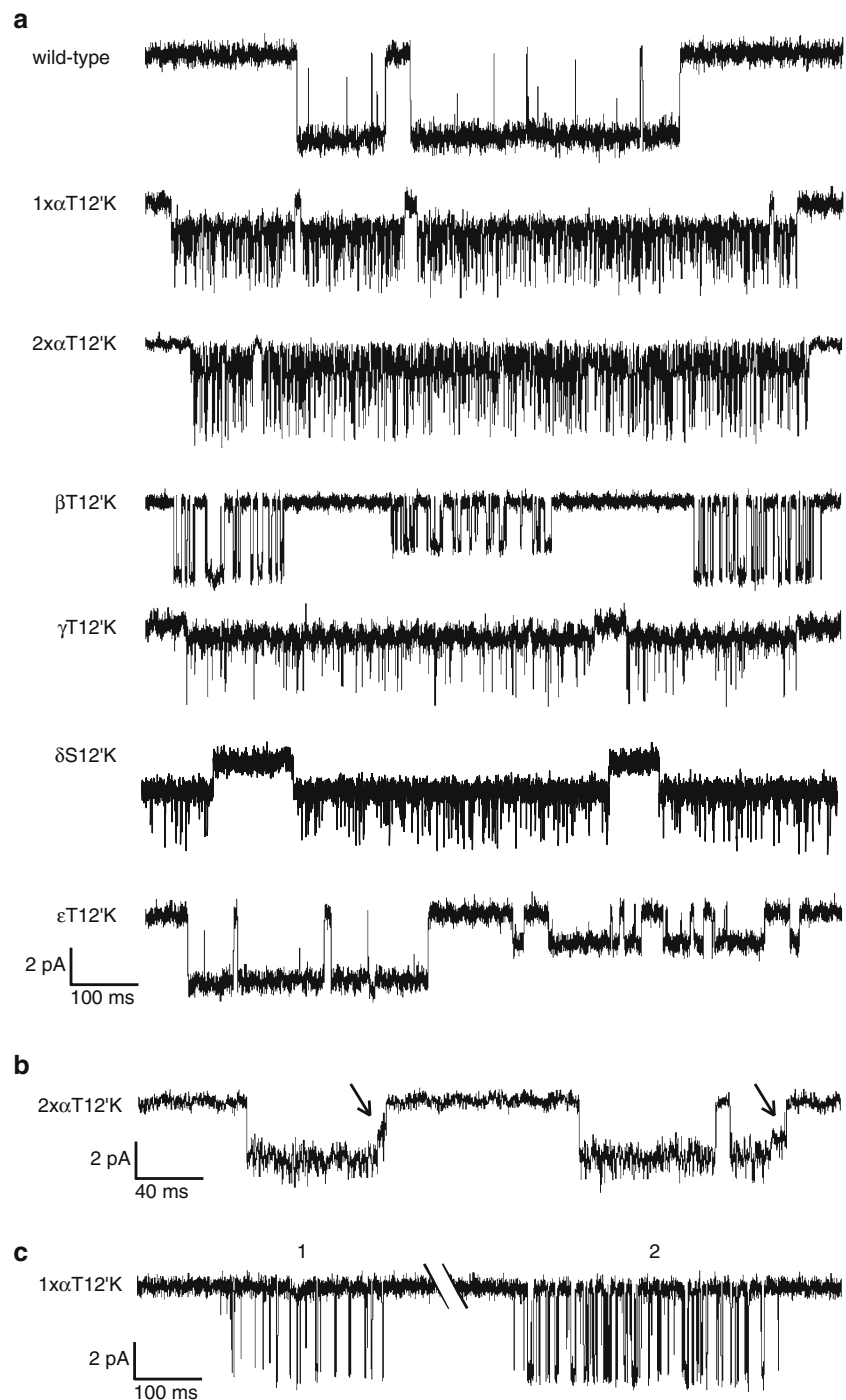


Fig. 1 Channel structure and TM2 alignment of human nAChR. Structure of the nAChR transmembrane segments of *Torpedo californica* nAChR (Unwin 2005), showing engineered lysines at

12' positions in green (left). Alignment of the TM2 domains of human muscle nAChR subunits (right). Differences within the TM2 (box) residue between subunits γ and ϵ are highlighted

Fig. 2 Single-channel current traces of 12'K nAChR mutants. **(a)** Predominant gating behaviour when stimulated by 10 μ M ACh. Downward deflections are channel openings. **(b)** For $2 \times \alpha$ T12'K receptors opening with largely increased open channel noise and stepwise channel closing (marked with *arrows*) was frequent. **(c)** For $1 \times \alpha$ T12'K receptors two different gating behaviours (marked 1 and 2) with distinct lifetimes (~ 1 and ~ 5 ms) that open towards a large conducting state (presumably wild-type conductance) were observed within the same patches



equal amount, resulting in expression of receptors containing either none, one (further $1 \times \alpha$ T12'K) or two ($2 \times \alpha$ T12'K) mutations.

Patch-clamp recordings

Recordings were performed at 18°C in buffer solution (see above) on cells that showed green fluorescence. Borosilicate patch pipettes (GB 150F-8P, Science

Products, Basel, Switzerland) were pulled with a P-87 micropipette puller (Sutter Instruments, Novato, USA) and had a resistance of 5–10 M Ω when filled with a solution containing NaCl 140 mM, EGTA 10 mM and HEPES 10 mM, pH adjusted to 7.4 with NaOH. Currents were measured with an EPC9 patch-clamp amplifier (HEKA, Lambrecht, Germany).

Single-channel measurements were carried out in the cell-attached configuration with 10 μ M ACh

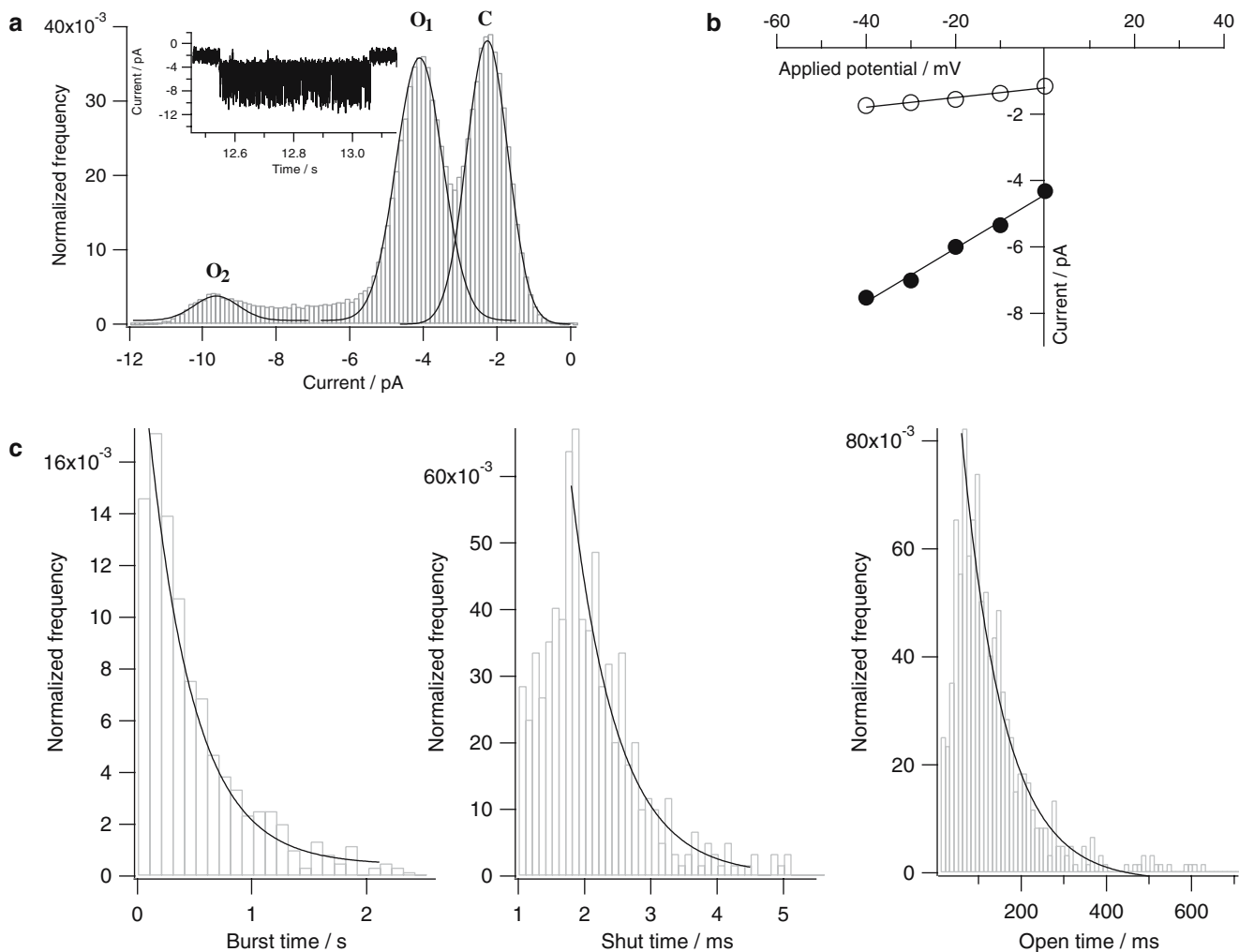


Fig. 3 Analysis of single-channel current traces of the $1 \times \alpha T12'K$ receptor. **(a)** Amplitude histogram of current trace (inset) fitted with three Gaussian distributions corresponding to the three states C, O₁ and O₂. **(b)** Values of current differences between the closed state C and the open states O₁ (open circle) and O₂ (filled circle), deduced from midpoints of Gaussian fits to amplitude histograms as a function of the applied voltage. *Lines*

are linear fits with respective slopes of 14 ± 2 and 76 ± 6 pS. **(c)** Dwell time histograms for burst time τ_b , the small component of the shut time τ_s and open time τ_o , derived from single-channel current traces. *Black lines* are single exponential fits to the data with time constants $\tau_b = 395 \pm 29$, $\tau_s = 0.7 \pm 0.1$ and $\tau_o = 106 \pm 10$ ms

(Sigma, Buchs, Switzerland) added to the pipette solution. The data were filtered with a Bessel filter at 2.9 kHz and sampled at 10 kHz.

Determination of single-channel conductances

Current differences between the open states O₁ and O₂ from the shut state C were obtained from Gaussian fits to current amplitude histograms for each measured voltage. Data were plotted as a function of applied voltage and fitted linearly, where the slope corresponds to the conductance of the respective states and the intersection at zero current to the reversal potential. For each patch, the applied voltage was corrected by

the reversal potential in order to obtain the actual value of the transmembrane voltage.

Dwell-time analysis

Segments of current traces showing single-channel activity were selected and idealized using the SKM algorithm of QuB (Qin et al. 1996). Histograms of open-channel time distributions showed a single component and hence could be well fitted with single exponential curves using Igor Pro and their means were calculated (Fig. 3c). Closed-channel time distributions showed two well-separated components. The smaller component had typically a lifetime of ~ 1 ms and its

upper limit (typically ~ 5 ms, which did not interfere with events of the larger component) was used as a critical time value t_{crit} . In the following, bursts were defined as a series of closely spaced openings into a conducting state that were longer than 20 ms and preceded and followed by closed intervals longer than t_{crit} .

Rate-kinetics of the fast gating transitions (protonation/deprotonation) were fitted (MIL-algorithm) for each patch experiment separately with correction for missed events, imposing a dead-time of 90–100 μs (Qin et al. 1996).

Results and discussion

Transitions between subconducting and fully conducting states have been observed recently in nAChR containing single lysines in the δ -TM2 domain and were interpreted as protonation–deprotonation events of single ϵ -NH₂ groups in the open channel pore (Cymes et al. 2005). Here, we address the question of symmetry between pore lining domains of the individual subunits by comparing experiments on lysine mutations at identical positions in each subunit.

12' Lysine mutation affects gating differently in α , γ and δ than in β and ϵ subunits

Whereas wild-type receptors show openings from a closed state C to a single open state, receptors with the introduced 12' lysine in α , γ or δ subunits feature within a burst a large conducting state O₂ and an open state O₁ of intermediate conductance and rapid transitions between these states, as seen in single-channel patch-clamp recordings at pH 7.4 (Fig. 2a). To the contrary, receptors β T12'K and ϵ T12'K opened either to a state of intermediate O₁ or large O₂ conductance and direct transitions between O₁ and O₂ have not been observed (Fig. 2a).

Different assemblies and gating behaviors for alpha mutated receptors

Co-transfection of both alpha wild-type and mutant α T12'K cDNA yields stochastic assembly of wild-type, $2 \times \alpha$ T12'K and $1 \times \alpha$ T12'K hybrid receptors (see methods), with the latter carrying the mutations in either α_δ or α_ϵ subunit. To identify openings specific to $1 \times \alpha$ T12'K hybrids we removed gating events similar to those observed for measurements with wild-type or $2 \times \alpha$ T12'K receptors. Amongst the remaining openings we observed predominantly openings with intermediate conductance (Fig. 2a) and more rarely openings from

the closed state towards a large conducting state (presumably identical to O₂), with two distinct lifetimes (Fig. 2b). These two gating types might reflect the two possible populations of $1 \times \alpha$ T12'K receptors containing the lysine in either α_δ or α_ϵ subunit.

$2 \times \alpha$ T12'K receptors showed in addition within bursts, next to O₁ \rightleftharpoons O₂ transitions, frequent fast transitions from the subconductance state O₁ to a closed or low conductance state. We assign this transition to the protonation of the second ϵ -NH₂ group, since its voltage dependence is similar to that of the first protonation, as shown in detail below.

In addition to the mentioned activities (Fig. 2a) alternative gating features were occasionally ($\sim 2\%$ of all bursts) observed for the $2 \times \alpha$ T12'K containing receptors (Fig. 2c). $2 \times \alpha$ T12'K receptors open to a large conductance state with largely increased open channel noise and channel closing through intermediate conducting states. These channel activities were observed sequentially (not simultaneously) to the ones shown in Fig. 2a in the same patches, indicating that this gating does not stem from multiple channels with different subunit assemblies, (mis-)foldings or conformations, but rather from one single channel. This demonstrates further complexity of open channel conformations and their gating pathways.

All mutant receptors gate between two open states

The existence of only two open states is most obvious for β T12'K and ϵ T12'K receptors due to their long lifetimes. Since receptors containing lysines engineered in α , γ and δ subunits showed gating with much shorter lifetimes, we analyzed power density spectra of current fluctuations for the exemplary δ S12'K mutant. We obtained single Lorentzian curves (not shown), supporting that the rapid current fluctuations are also well described by a simple two-state Markov process (Sigworth 1985). Therefore, we interpret these transitions as protonation events.

Channel currents are blocked dissimilarly by protonation of different 12' lysines

Conductance values g_1 and g_2 of the states O₁ and O₂, obtained as shown in Fig. 3, are summarized for all receptors in Table 1. We calculated the extent of channel block

$$(g_2 - g_1)/g_2 \quad (1)$$

which is thought to reflect the proximity of the lysine to the ion pathway (Cymes et al. 2005). The value of the δ

subunit is significantly lower ($56 \pm 14\%$) than those of α and γ subunit (82 ± 13 and $82 \pm 17\%$). The conductance value corresponding to the second protonation event is below the resolution limit and indistinguishable from zero and might reflect a nearly completely blocked ion pathway.

Current differences between states O_1 and O_2 are relatively small for receptors β T12'K and ϵ T12'K compared to the other receptors, suggesting little channel blocking (40 ± 11 and $27 \pm 8\%$, respectively). However, as discussed later, we cannot be sure that lysine protonation is the origin. For this reason we limited further analysis concerning these two mutants to the determination of gating rate constants for O_2 at different membrane potentials (Fig. 4).

Analysis of gating kinetics reveals voltage dependence of pK_a values

In the following we focus on receptors $1 \times \alpha$ T12'K, $2 \times \alpha$ T12'K, γ T12'K and δ T12'K, with emphasis on their fast gating transitions (as shown in Fig. 2a), which we associate with 12'K lysine protonation. Shut times τ_s , open times τ_o , $C \rightleftharpoons O_1 + O_2$ and burst times were not voltage dependent within experimental error for receptors $1 \times \alpha$ T12'K, γ T12'K and δ T12'K. We observed that burst times of δ S12'K receptors were about two-fold larger (757 ± 159 ms) than that of $1 \times \alpha$ T12'K (395 ± 29 ms) or γ T12'K (414 ± 58 ms) receptors, whereas respective apparent equilibrium constants did not vary significantly (Table 1). Presently, we cannot interpret this feature in concrete structural or functional terms. However, together with the significantly different extent of channel block it is evident that the lysine engineered in the δ subunit is not located identically to lysines in α and γ subunits with respect to the ion pathway and that the gating transmission of the δ subunit differs to that from other subunits.

Assuming a first-order proton binding reaction to lysine side chains, the protonation rate k_{21} (s^{-1}) is

defined as the closing rate from state O_2 to O_1 , while the deprotonation rate k_{12} (s^{-1}) is defined as the opening rate from O_1 to O_2 . The actual protonation rate constant (in $M^{-1}s^{-1}$) is given by $k_{21}/[H^+]$. Figure 5 shows that k_{12} and k_{21} (and additionally the rate constants of channel gating $C \rightleftharpoons O_1$ for $2 \times \alpha$ T12'K) are voltage dependent. Whereas k_{21} was larger than k_{12} at all examined voltages, the latter increased more strongly with membrane hyperpolarization (Fig. 5). Extrapolated values at zero voltage are reported in Table 2. We calculated pK_a values of individual lysines at pH 7.4 as

$$pK_a = -\log(k'_{12} \times [H^+]/k_{21}) \quad (2)$$

(Fig. 6 and Table 2).

Both, the pK_a value and extent of channel block found for the δ S12'K receptor are in good agreement with similar measurements on the mouse nAChR (Cymes et al. 2005), which shows that the high homology between these receptors on the level of the primary sequence (>98%) seems also to extend to the structural level. Additionally, we found that all pK_a values, except for γ T12'K, decrease with membrane hyperpolarization, which is mainly due to the voltage dependence of the deprotonation rate. This might result from reorientation of these residues towards a more hydrophobic environment with hyperpolarization or the increased electric force on the proton.

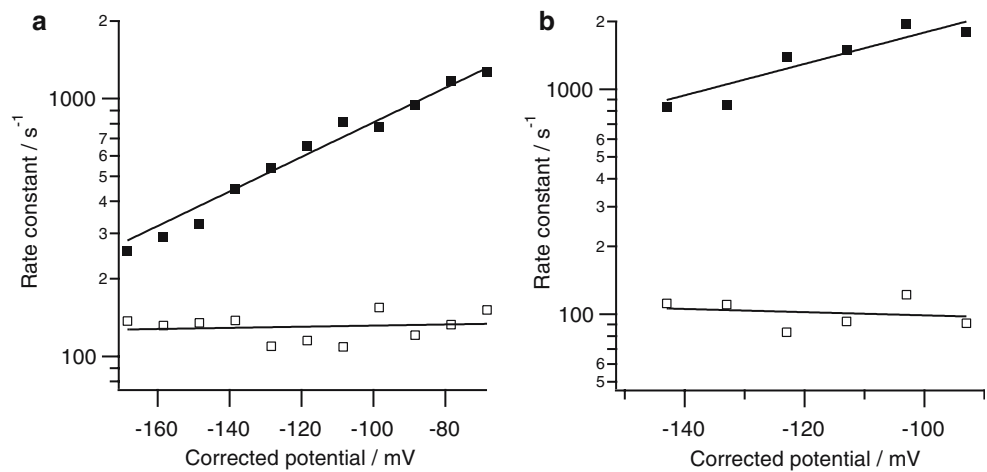
Extrapolation of the data shows, that pK_a values for protonation of one 12' lysine would be similar at zero voltage (~ 8.6). This might reflect that at zero voltage the lysines have virtually identical microenvironments, which diverge with membrane hyperpolarization. We speculate that the observed voltage dependence of pK_a values (which ranks γ T12'K < δ S12'K $\sim 1 \times \alpha$ T12'K) might reflect the degree of displacement of the respective 12' residue with membrane polarization. Protonation of the second lysine in the $2 \times \alpha$ T12'K receptor seems to be less favorable ($\Delta G \approx 3.9$ kJ/mol

Table 1 Single channel conductance and dwell time analysis of 12'K receptors

Receptor	g_1 (pS)	g_2 (pS)	Channel block (%)	Burst length (ms)	Shut time τ_s (ms)	Open time τ_o (ms)	Apparent equilibrium constant Θ	No.
$1 \times \alpha$ T12'K	14 ± 2	76 ± 6	82 ± 17	395 ± 29	0.7 ± 0.1	106 ± 10	151 ± 33	4
$2 \times \alpha$ T12'K	26 ± 2	76 ± 3	66 ± 9	–	–	–	–	2
β T12'K	42 ± 5	70 ± 2	40 ± 11	–	–	–	–	3
γ S12'K	11 ± 2	60 ± 3	82 ± 13	414 ± 58	1.0 ± 0.2	111 ± 25	111 ± 59	3
δ T12'K	25 ± 2	57 ± 4	56 ± 14	757 ± 159	1.1 ± 0.2	205 ± 27	189 ± 69	4
ϵ T12'K	55 ± 2	75 ± 3	27 ± 8	–	–	–	–	3

Single channel conductances g_1 and g_2 , extent of channel block $(g_2 - g_1)/g_2$ in percent, burst length, shut time τ_s , open time τ_o , apparent gating equilibrium constant $\Theta = \tau_o/\tau_s$ and number of patches evaluated for the gating transition between the closed state C and the open states O_1 and O_2

Fig. 4 Rates for channel closing (filled square) and channel opening (open square) for the transition $C \rightleftharpoons O_2$ as a function of voltage for $\beta T12'K$ (a) and $\epsilon T12'K$ (b) receptors. Fitting (lines) the data to $k(V) = k_{V=0} \exp(-z\delta F/RT) V$ gave for $\beta T12'K$: $k_{12,V=0} = 138 \pm 20 \text{ s}^{-1}$, $z\delta = -0.01 \pm 0.03$, $k_{21,V=0} = 3,779 \pm 307 \text{ s}^{-1}$, $z\delta = -0.39 \pm 0.02$ and $\epsilon T12'K$: $k_{12,V=0} = 84 \pm 39 \text{ s}^{-1}$, $z\delta = 0.04 \pm 0.10$, $k_{21,V=0} = 8,913 \pm 3,740 \text{ s}^{-1}$, $z\delta = -0.40 \pm 0.10$



at 0 V) as compared to protonation of the first lysine, as can be seen from the lower $pK_{a,V=0}$ value (~ 7.7). This is consistent, considering that the first protonated lysine hinders the second protonation event due to electrostatic repulsion. The observed voltage dependence demonstrates also that the precision of

pK_a values determined by this approach is at least ten-fold lower as estimated before (Cymes et al. 2005), since it is dominated by the error of the exact membrane potential measurement.

Fig. 5 Rates of the first [k_{12} (open square)/ k_{21} (filled square)] $O_1 \rightleftharpoons O_2$ and the second [k_{12} (open circle)/ k_{21} (filled circle)] $C \rightleftharpoons O_1$ channel blocking (only receptor $2 \times \alpha T12'K$) as a function of the corrected potential for receptors $1 \times \alpha T12'K$ (a), $2 \times \alpha T12'K$ (b), $\gamma T12'K$ (c) and $\delta T12'K$ (d). Parameters of fits (solid lines) are summarized in Table 2

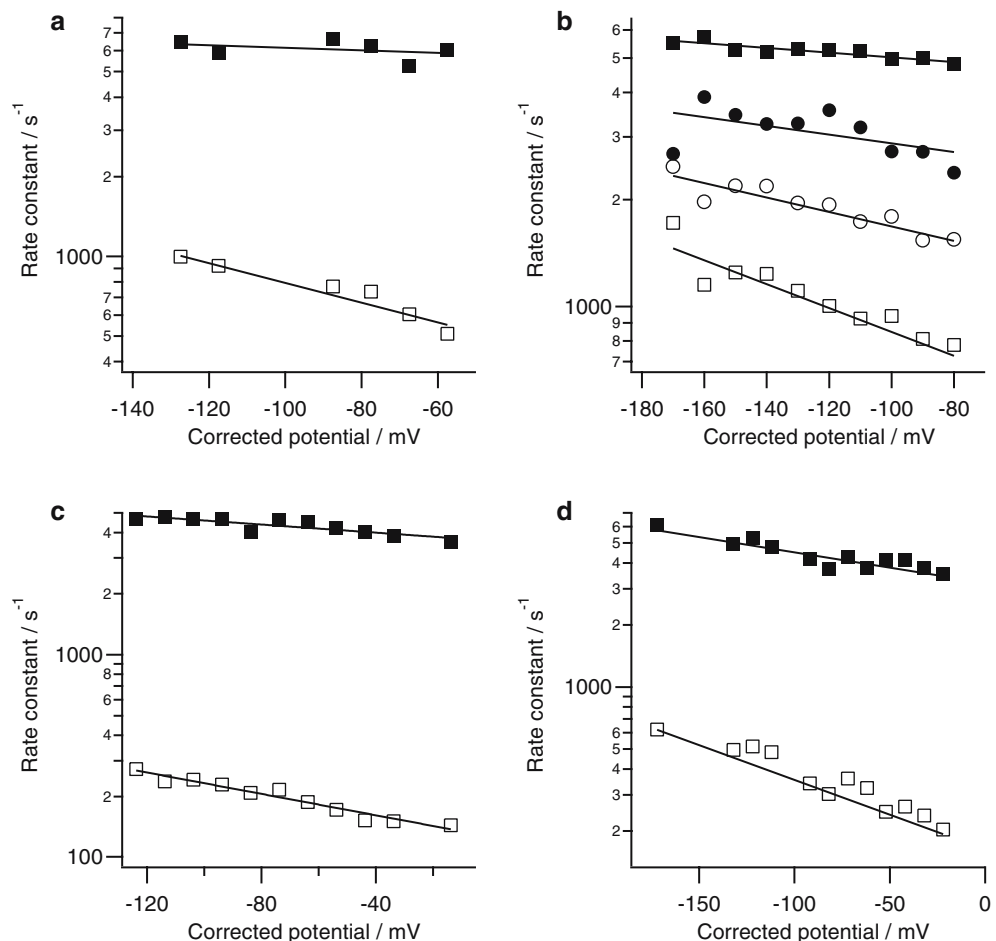


Table 2 Voltage dependence of (de-)protonation rates and pK_a values

Receptor	Deprotonation		Protonation		pK_a	
	$k_{12,V=0} \text{ s}^{-1}$	$z\delta$	$k_{21,V=0} \text{ s}^{-1}$	$z\delta$	$pK_{a,V=0}$	$z'\delta'$
$1 \times \alpha\text{T12}'\text{K}$	256 ± 36	0.33 ± 0.04	$2,311 \pm 381$	0.09 ± 0.05	8.6 ± 0.1	0.08 ± 0.02
$\gamma\text{S12}'\text{K}$	127 ± 5	0.15 ± 0.01	$3,644 \pm 139$	0.06 ± 0.01	8.7 ± 0.1	-0.01 ± 0.01
$\delta\text{T12}'\text{K}$	195 ± 14	0.18 ± 0.02	$3,245 \pm 148$	0.09 ± 0.01	8.7 ± 0.1	0.06 ± 0.01
$2 \times \alpha\text{T12}'\text{K}_{(1)}$	406 ± 71	0.19 ± 0.03	$4,332 \pm 155$	0.04 ± 0.01	8.4 ± 0.1	0.06 ± 0.01
$2 \times \alpha\text{T12}'\text{K}_{(2)}$	$1,076 \pm 109$	0.12 ± 0.02	$2,242 \pm 440$	0.07 ± 0.04	7.7 ± 0.1	0.02 ± 0.02

Fit values to opening and closing rates and pK_a values as a function of voltage V , according to $k(V) = k_{V=0} \exp(-z\delta F/RT) V$ and $pK_a(V) = pK_{a,V=0} + (z'\delta'F/RT) V$ (Auerbach et al. 1996) (Figs. 5, 6), where z is the amount of charge moving across the fraction δ of the electric field, F the Faraday constant, R the gas constant and T the temperature. For the receptor $2 \times \alpha\text{T12}'\text{K}$ the first ($\text{O}_2 \rightleftharpoons \text{O}_1$) and the second ($\text{O}_1 \rightleftharpoons \text{C}$) protonation events are indexed (1) and (2), respectively

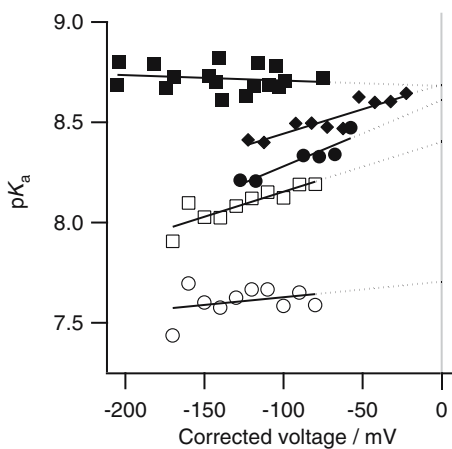


Fig. 6 pK_a values as a function of voltage. Calculated pK_a values for introduced 12' lysines as function of voltage for $2 \times \alpha\text{T12}'\text{K}$ (first blocking: open square, second blocking: open circle), $1 \times \alpha\text{T12}'\text{K}$ (filled circle), $\gamma\text{T12}'\text{K}$ (filled square) and $\delta\text{S12}'\text{K}$ (filled diamond) receptors. Solid lines are linear fits to the data, dashed lines extrapolations thereof, and the grey line indicates zero voltage. Fit parameters are summarized in Table 2

The different gating of receptors with $\beta\text{T12}'\text{K}$ or $\epsilon\text{T12}'\text{K}$ might have another origin

Similar gating behaviour and single-channel conductances to that of the $\beta\text{T12}'\text{K}$ receptor have been already observed after transfection of *Xenopus oocytes* with cDNA for all but β subunits (Camacho et al. 1993). We can therefore not exclude that at least in the case of the $\beta\text{T12}'\text{K}$ mutation efficient assembly with wild-type stoichiometry might be compromised. However, assuming a correctly assembled receptor, the lack of fast channel blocking (~ 1 ms) would indicate that lifetimes of protonated and deprotonated states are much larger than the respective burst lifetimes (~ 100 ms), which could be explained by extreme pK_a values of the respective lysines in at least two largely different environments. From the above values we can estimate (see Eq. 2) that pK_a values of the lysines in

the open states O_1 and O_2 would be superior to ~ 9.4 and inferior to ~ 5.4 , respectively, in order to explain that no fast current fluctuations are observed.

Differences between highly homologous subunits γ and ϵ

The difference in gating behavior of $\gamma\text{T12}'\text{K}$ and $\epsilon\text{T12}'\text{K}$ receptors is unexpected, since their respective TM2 regions differ only in positions 4' and 18' (Fig. 1). Their overall primary sequence homology is $>90\%$, which shows that this approach is not limited to the structural level, but is rather sensitive to the functional contributions of subunits. The structure of *Torpedo californica* nAChR (Fig. 1) suggests that the introduced 12' lysines in the muscle nAChR do not point directly towards the channel, but rather feature close proximity to TM1 (position 12') and TM2 (position 14') of the adjacent subunit. However molecular dynamics simulations show the 12' residue to be much closer to the central pore axis and the conformational flexibility of lysines opens multiple structural possibilities, making it difficult to determine specific interactions (Kim et al. 2004).

Conclusion

Lysines 12' of TM2 of the different subunits are probing different microenvironments as reflected by their different gating behaviours, their respective kinetic parameters and the voltage dependence of their pK_a values. The latter is different for each subunit, and reflects and quantifies the structural distortions such as they might occur during channel gating.

Acknowledgments We thank Karen Martinez for critical manuscript reading and Pierre-Jean Corringer and David Beeson for providing cDNA of wild-type nAChR subunits. This work was supported by the Swiss National Science Foundation

(31–57023.99 and 31–00A0–102062/1) and the European Commission via contract LSHG-CT-2004–504601 (E-MeP).

References

- Absalom NL, Lewis TM, Schofield PR (2004) Mechanisms of channel gating of the ligand-gated ion channel superfamily inferred from protein structure. *Exp Physiol* 89:145–153
- Akabas MH, Kaufmann C, Archdeacon P, Karlin A (1994) Identification of acetylcholine receptor channel-lining residues in the entire M2 segment of the alpha subunit. *Neuron* 13:919–927
- Auerbach A, Sigurdson W, Chen J, Akk G (1996) Voltage dependence of mouse acetylcholine receptor gating: different charge movements in di-, mono- and unliganded receptors. *J Physiol* 494(Pt 1):155–170
- Camacho P, Liu Y, Mandel G, Brehm P (1993) The epsilon subunit confers fast channel gating on multiple classes of acetylcholine receptors. *J Neurosci* 13:605–613
- Cymes GD, Ni Y, Grosman C (2005) Probing ion-channel pores one proton at a time. *Nature* 438:975–980
- Dahan DS, Dibas MI, Petersson EJ, Auyeung VC, Chanda B, Bezanilla F, Dougherty DA, Lester HA (2004) A fluorophore attached to nicotinic acetylcholine receptor beta M2 detects productive binding of agonist to the alpha delta site. *Proc Natl Acad Sci USA* 101:10195–10200
- Grosman C, Auerbach A (2000) Asymmetric and independent contribution of the second transmembrane segment 12' residues to diliganded gating of acetylcholine receptor channels: a single-channel study with choline as the agonist. *J Gen Physiol* 115:637–651
- Grutter T, Changeux JP (2001) Nicotinic receptors in wonderland. *Trends Biochem Sci* 26:459–463
- Hucho F, Oberthur W, Lottspeich F (1986) The ion channel of the nicotinic acetylcholine receptor is formed by the homologous helices M II of the receptor subunits. *FEBS Lett* 205:137–142
- Maconochie DJ, Fletcher GH, Steinbach JH (1995) The conductance of the muscle nicotinic receptor channel changes rapidly upon gating. *Biophys J* 68:483–490
- Miyazawa A, Fujiyoshi Y, Unwin N (2003) Structure and gating mechanism of the acetylcholine receptor pore. *Nature* 423:949–955
- Kim S, Aaron K, Chamberlain, James U. Bowie (2004). A model of the closed form of the nicotinic acetylcholine receptor m2 channel pore. *Biophys J* 87:792–799
- Newland CF, Beeson D, Vincent A, Newsom-Davis J (1995) Functional and non-functional isoforms of the human muscle acetylcholine receptor. *J Physiol* 489(Pt 3):767–778
- Qin F, Auerbach A, Sachs F (1996) Estimating single-channel kinetic parameters from idealized patch-clamp data containing missed events. *Biophys J* 70:264–280
- Sigworth FJ (1985) Open channel noise I. Noise in acetylcholine receptor currents suggests conformational fluctuations. *Biophys J* 47:709–720
- Unwin N (2005) Refined structure of the nicotinic acetylcholine receptor at 4A resolution. *J Mol Biol* 346:967–989