Delimiting the Binding Site for Quaternary Ammonium Lidocaine Derivatives in the Acetylcholine Receptor Channel

JUAN M. PASCUAL*† and ARTHUR KARLIN*‡§

From the *Center for Molecular Recognition, †Department of Neurology, ‡Department of Physiology and Cellular Biophysics, and §Department of Biochemistry and Molecular Biophysics, Columbia University, New York 10032

ABSTRACT The triethylammonium QX-314 and the trimethylammonium QX-222 are lidocaine derivatives that act as open-channel blockers of the acetylcholine (ACh) receptor. When bound, these blockers should occlude some of the residues lining the channel. Eight residues in the second membrane-spanning segment (M2) of the mouse-muscle α subunit were mutated one at a time to cysteine and expressed together with wild-type β, γ, and δ subunits in Xenopus oocytes. The rate constant for the reaction of each substituted cysteine with 2-aminoethyl methanethiosulfonate (MTSEA) was determined from the time course of the irreversible effect of MTSEA on the ACh-induced current. The reactions were carried out in the presence and absence of ACh and in the presence and absence of QX-314 and QX-222. These blockers had no effect on the reactions in the absence of ACh. In the presence of ACh, both blockers retarded the reaction of extracellularly applied MTSEA with cysteine substituted for residues from αVal255, one third of the distance in from the extracellular end of M2, to αGlu241, flanking the intracellular end of M2, but not with cysteine substituted for αLeu258 or αGlu262, at the extracellular end of M2. The reactions of MTSEA with cysteines substituted for αLeu258 and αGlu262 were considerably faster in the presence of ACh than in its absence. That QX-314 and QX-222 did not protect α2.258C and α2.262C against reaction with MTSEA in the presence of ACh implies that protection of the other residues was due to occlusion of the channel and not to the promotion of a less reactive state from a remote site. Given the 12-Å overall length of the blockers and the α-helical conformation of M2 in the open state, the binding site for both blockers extends from αVal255 down to αSer248.

KEY WORDS: cysteine mutagenesis • reaction kinetics • methanethiosulfonate • open-channel block

INTRODUCTION

There is considerable evidence that certain noncompetitive inhibitors (NCIs)† of the acetylcholine receptor act as open-channel blockers, binding within the open channel and occluding it (Adams, 1976; Ruff, 1977; Ascher et al., 1978; Neher and Steinbach, 1978; Colquhoun and Sheridan, 1981; Farley et al., 1981). NCIs, binding either within the channel or at lower affinity sites (Krodel et al., 1979; Heidmann et al., 1983; Herz et al., 1987; Lurtz et al., 1997), can also interact with closed states of the receptor (Adams, 1976; Farley et al., 1981; Neher, 1983) and promote desensitization (Magleby and Pallotta, 1981; Karpen et al., 1982; Maleque et al., 1982; Siné and Taylor, 1982; Neher, 1983; Oswald et al., 1983; Boyd and Cohen, 1984; Clapham and Neher, 1984; Gage and Wachtel, 1984). Based on the likelihood that NCIs bind within the channel, reactive NCI derivatives were used to identify channel-lining residues. Chlorpromazine (Revah et al., 1990), triphenylmethylphosphonium (Huchu et al., 1986), and meproafen mustard (Pedersen et al., 1992) all labeled residues in or flanking the predicted second membrane-spanning segment (M2) (Fig. 1) of one or more Torpedo species acetylcholine (ACh)-receptor subunits (Fig. 2). In particular, the labeling of α2.248 and the aligned residues in β, γ, and δ by chlorpromazine and triphenylmethylphosphonium (Fig. 2) provided evidence that the channel was on the axis of the pentameric complex (αβγδ) surrounded by the five M2 segments.

Other crucial evidence for the topology of M2 and for its central role in the channel was provided by the effects on conductance and selectivity of mutating charged residues flanking M2 in each of the subunits (Imoto et al., 1988). In addition, the effects of mutations of the M2 residues α2.248 and α2.252 and of the aligned residues in the other subunits (Fig. 2) on the kinetics of QX-222 block (Fig. 3) provided further evidence that open-channel blockers bind in the channel in the vicinity of these M2 residues (Charnet et al., 1988; Leonard et al., 1988). Furthermore, the mutation L247T in the homopentameric neuronal ACh receptor formed by the α7 subunit (Anand et al., 1997; Cooper...
et al., 1991) resulted in the loss of sensitivity to block by QX-222 (Revah et al., 1991). (α7)L247 aligns with mouse-muscle αL251 (Fig. 2).

Not all evidence, however, points to the region between αS248 and αS252 as the sole binding site in the channel. Quinacrine azide photolabeled residues at the extracellular end of αM1, specifically in the open state of the channel (Karlin, 1989; DiPaola et al., 1990), and mutations at the extracellular end of αM1 affected quinacrine but not chlorpromazine binding (Tama-mizu et al., 1995). Nevertheless, chlorpromazine and a number of other NCIs retarded the labeling by quinacrine azide (DiPaola et al., 1990) and competed with the binding of quinacrine (Lurtz et al., 1997). In addition, meproadin mustard labeled αE262 at the extracellular end of M2 (Pedersen et al., 1992). Different NCIs may bind to nonidentical, possibly overlapping sites in the channel.

We report here on another approach to the localization of the sites of action of channel-blocking NCIs, using them in conjunction with the substituted-cysteine-accessibility method (SCAM). SCAM was applied previously to the α and β subunits, identifying residues exposed in the channel along the entire length of M2 (Fig. 2), in the M1–M2 loop, and in the extracellular third of M1 (Akabas et al., 1992, 1994; Akabas and Karlin, 1995; Zhang and Karlin, 1997, 1998; Wilson and Karlin, 1998). These residues, when substituted by cysteine, reacted with small, positively charged sulfhydryl-specific methanethiosulfonate reagents such as 2-aminobenzyl methanethiosulfonate (MTSEA). The rate constants for the reactions with the different residues depended on the state of the receptor (Pascual and Karlin, 1998) and on the side of application of the reagent (Wilson and Karlin, 1998). In this paper, we test to what extent the channel blockers QX-222 and QX-314 (Fig. 3; Neher and Steinbach, 1978; Horn et al., 1980; Neher, 1983; Charnet et al., 1990) protect Cys-substituted channel-lining residues in αM2 from reaction with extracellularly applied MTSEA. Previously, Cys-substituted residues in the M2 segment of the GABAA receptor α1 subunit were protected by picrotoxin (Xu et al., 1995), Cys-substituted residues in the Na channel were protected by tetrodotoxin (Kirsch et al., 1994), Cys-substituted residues in Kv2.1 potassium channel were protected by tetethylammonium (Pascual et al., 1995), and Cys-substituted residues in SHaker potassium channel were protected by agitoxin (Gross and MacKinnon, 1996). We find that QX-222 and QX-314, in the presence of ACh,
Cells were kept in culture and used for recording on days 1–10. oocyte bath solution contained (mM): 115 NaCl, 2.5 KCl, 1.8 MgCl$_2$, 10 HEPES, pH 7.2, except where indicated otherwise. Solutions flowed at 7 ml/min first through a stainless steel coil immersed in a thermostat at 18.0°C, and then past the oocyte, which was held in a rectangular chamber with a cross-section normal to the direction of solution flow of 4 mm$^2$. An agar bridge connected a Ag/AgCl reference electrode to the bath and was placed as close as possible to the oocyte. The bath was clamped at ground potential. We used beveled agarose-cushion (Schreibmayer et al., 1994) glass micropipettes filled with 3 M KCl, of resistance $\sim$0.2–0.5 M$\Omega$, for both current-passing and voltage-recording electrodes. A few uninjected oocytes from each batch were tested for the presence of endogenous ACh-induced currents, which were never found. The function of wild-type and mutant receptors was assayed as the ACh-induced current elicited by the application of brief (3–25 s) pulses of ACh, at a concentration 10$^4 \times$ the EC$_{50}$ as determined for each mutant, and at a holding potential of $-50$ mV, except where indicated otherwise. ACh-induced currents ranged from 1 to 25 $\mu$A at $-50$ mV.

**Characterizing Inhibition by QX-314 and QX-222**

The concentrations (IC$_{50}$) of QX-314 and of QX-222 that inhibited the ACh-induced current by 50% were determined at several holding potentials (see Fig. 4). ACh was applied at 10$^4 \times$ its EC$_{50}$ for each mutant. Leak currents were measured at holding potentials of $-50$, $-125$, $-75$, and $-25$ mV, each held for 400 ms. ACh was added and, at the peak of the current, the holding potential was stepped through the same four values. After a 5-min wash, QX-222 or QX-314 was applied continuously for 2 min, during which time leak and ACh-induced currents were determined as before. This protocol was repeated with increasing concentrations of blocker. We calculated the net ACh-induced currents in the presence (I$_{ACh,QX}$) and absence (I$_{ACh}$) of blocker, and we fit the data at each membrane potential to the equation

$$I(t) = I_{ACh}(0 mV) \times \exp(-t/\tau)$$

where $\tau$ is the charge on the blocker, $F$ is the Faraday constant, $\psi$ is the transmembrane potential difference (inside-outside), and $\delta$ is the apparent electrical distance from the extracellular medium to the blocker binding site.

**Protection**

The protection of substituted Cys against reaction with MTSEA afforded by bound blocker was estimated from the second-order rate constants of the reaction in the presence and absence of blocker. The second-order rate constants ($k$) were determined as previously described (Pascual and Karlin, 1998). Oocytes were superfused for 20 s with ACh (at 10$^4 \times$ the EC$_{50}$ as determined for each mutant), for 3 min with bath solution, for 2–25 s with MTSEA in the presence of ACh, or for 1–4 min with MTSEA in the absence of ACh, for 3 min with bath solution, for 20 s with ACh, and for 3 min with bath solution; this sequence was repeated several times. MTSEA was applied at concentrations ranging from 5 $\mu$M for the rapidly reacting residues up to 10 mM for the slowly reacting residues. MTSEA was applied also in the presence of QX-314 and QX-222, both in the presence and absence of ACh. The peaks of the ACh-induced currents obtained before and after the exposure to MTSEA were fit to $I = I_0 + (I_{0} - I_{eq}) \exp(-t/\tau)$, where $I_{eq}$ is the initial ACh-induced peak current, $I_{0}$ is the ACh-induced current after a cumulative time, $t$, of application of MTSEA at concentration, $x$, and $L$ is the ACh-induced current after complete reaction of the Cys. Note that the fraction of unmodified receptors $= (I_{0} - L_{eq})/(I_{0} - L_{eq})$ (Pascual and Karlin, 1998).

![Figure 3. Structures of QX-314 and QX-222.](image-url)
In the presence of ACh, the receptors are distributed among a number of states: closed, open, and desensitized. During the 2–25 s that MTSEA and ACh (at 10× EC_{50}) were applied, the receptor was predominantly in the open state or the rapid-onset desensitized state. There was little slow desensitization during this time, and in any case the rate constants for the reactions of two mutants in the slow-onset desensitized state were small (Pascual and Karlin, 1998; Wilson and Karlin, 1998). The rate constant for the reaction in the mixed open state and rapid-onset desensitized state we call $k_{\text{open}}$. When blocker was also added, it could have bound to the open state and to the rapid-onset desensitized state (Neher, 1983). To compare the effects of QX-314 and QX-222, at different concentrations, on the rate constants for the reactions of MTSEA with the different mutants, we estimated the rate constants for the reactions of MTSEA with the blocked state(s), $k_{\text{blocked}}$, from the observed rate constants in the presence of blocker, $k_{\text{obs}}$, and from the IC_{50} for the particular blocker and mutant. We assumed that the observed rate constant, $k_{\text{obs}}$, is approximated by $k_{\text{obs}} = k_{\text{open}}(1 - y) + k_{\text{blocked}}y$, where $y$, the fraction of channels that was blocked, is given by $y = 1/(1 + IC_{50}/[QX])$.

With each oocyte, we first applied MTSEA (in the presence of ACh) several times in the absence of blocker to determine $k_{\text{open}}$, and then applied MTSEA several times in the presence of blocker to determine $k_{\text{obs}}$. The concentrations of blocker used were in the range of 2–5× IC_{50} for the given blocker and mutant, so that $y$ ranged from 2/3 to 5/6. If we take the extent of protection against reaction to be $1 - (k_{\text{blocked}}/k_{\text{open}})$, then $1 - (k_{\text{blocked}}/k_{\text{open}}) = (1/y)[1 - (k_{\text{obs}}/k_{\text{open}})]$; i.e., the (corrected) extent of protection was 1.2–1.5× the observed extent of protection.

**Results**

Eight residues in αM2 were substituted, one at a time, by Cys and expressed in *Xenopus oocytes* together with wild-type β, γ, and δ subunits. In each case, the current elicited by ACh was inhibited by QX-314 and QX-222, and the inhibition increased with increasing concentration of blocker. This inhibition is illustrated for αL251C and QX-314 in Fig. 4. We derived the IC_{50}s for the inhibition by the blockers of the ACh-induced currents at each holding potential from such experiments. The IC_{50} for inhibition of the mutants by QX-314 ranged from 0.4× (αT244C) to 4.6× (αV255C) the IC_{50} for wild type (Table I). The range for QX-222 was from 0.05× (αT244C) to 1× (αS252C) the IC_{50} for wild type. Typically, when QX-314 and ACh were washed out, there was an immediate increase in the amplitude of the current before it returned to the baseline (Fig. 4, B–D). This increased current resulted either from the blocker dissociating from its site of inhibition before ACh dissociated from its binding sites or from the faster dilution below effective concentration of the blocker (initially at 2–5× IC_{50}) than of ACh (initially at 10× EC_{50}).

For both blockers and for all mutants, the more negative the holding potential, the greater the inhibition of current (Fig. 4). The voltage dependence of the inhibition was consistent with the site of inhibition being in the open channel. The voltage dependence of the IC_{50} was fit by the Woodhull (1973) equation to yield the blocker charge times the apparent electrical distance, $\delta$. For these quaternary ammonium blockers, $\delta = 1$. For QX-314, $\delta$ was 0.35 for wild type and ranged from 0.28 for αV255C to 0.57 for αL251C (Table I). For

**Figure 4.** Inhibition of ACh-induced current in αL251C by QX-314 at different holding potentials. The top trace shows the holding potential. Both before and 3 s after the application of 25 μM ACh, the holding potential was stepped from −50 mV to −125, −75, −25, and back to −50 mV. The bottom trace shows the current. (Inset) The peak ACh-induced current during the voltage steps minus the current during the voltage steps before ACh was added, with an expanded time axis. (A) No QX-314, (B) 0.1 mM QX-314, (C) 0.3 mM QX-314, (D) 1 mM QX-314. The four sections are part of a single experiment. Between the sections shown, the oocyte was washed and QX-314 at the indicated concentration was applied for 45 s before, and in addition to, the time indicated in the traces.

614  Open-Channel Blocking Site in the Acetylcholine Receptor
Woodhull equation to the IC50 as a function of holding potential.

The interpretation of the apparent electrical distances, \( \delta \), estimated from the fit of the Woodhull equation to the IC50 as a function of holding potential.

Table I: The Characteristics of QX-314 and QX-222 Inhibition of the ACh-induced Currents Mediated by Wild-Type and Mutant ACh Receptors

| Mutant   | IC50 \( \mu M \) | \( \delta \) | IC50 \( \mu M \) | \( \delta \) |
|----------|------------------|-------------|------------------|-------------|
| Wild type| 78 ± 12 (4)      | 0.35 ± 0.02 | 2780 ± 230 (7)   | 0.80 ± 0.01 |
| \( \alpha E262C \) | 92 ± 11 (4)      | 0.39 ± 0.03 | 765 ± 80 (4)     | 0.55 ± 0.05 |
| \( \alpha L258C \) | 284 ± 21 (4)     | 0.29 ± 0.03 | 558 ± 112 (4)    | 0.34 ± 0.02 |
| \( \alpha V255C \) | 563 ± 51 (3)     | 0.28 ± 0.01 | 426 ± 93 (4)     | 0.47 ± 0.07 |
| \( \alpha S252C \) | 175 ± 15 (3)     | 0.45 ± 0.03 | 2900 ± 294 (4)   | 0.66 ± 0.08 |
| \( \alpha L251C \) | 297 ± 48 (4)     | 0.57 ± 0.02 | 530 ± 9 (5)      | 0.53 ± 0.03 |
| \( \alpha S248C \) | 84 ± 8 (3)       | 0.38 ± 0.03 | 1339 ± 131 (4)   | 0.68 ± 0.04 |
| \( \alpha T244C \) | 30 ± 2 (4)       | 0.51 ± 0.02 | 140 ± 14 (4)     | 0.68 ± 0.03 |
| \( \alpha E241C \) | 256 ± 12 (4)     | 0.37 ± 0.01 | 1862 ± 299 (4)   | 0.42 ± 0.04 |

The IC50s were determined at holding potentials of \(-25, -50, -75, \) and \(-125\) mV (see Fig. 4 and METHODS). The means and SEMs of the IC50s at \(-50\) mV are presented. The voltage dependence of the block is characterized by the apparent electrical distance, \( \delta \), estimated from the fit of the Woodhull equation to the IC50 as a function of holding potential.

QX-222, \( \delta \) was 0.80 for wild type and ranged from 0.34 for \( \alpha L258C \) to 0.68 for \( \alpha T244C \) and \( \alpha S248C \). The effects of the mutations on the apparent electrical distances for QX-314 and for QX-222 were not correlated. The interpretation of the apparent electrical distances is difficult because the IC50s are dependent not only on the equilibrium dissociation constants of the blockers, but also on the kinetics of the transitions between functional states.

The results were obtained with \( \alpha L251C \) and \( \alpha S248C \) at concentrations several times their IC50s. By contrast, the reactions of MTSEA in the presence of ACh with Cys-substituted residues deeper in the channel were markedly slowed by the addition of these blockers. For example, the irreversible reaction of MTSEA in the presence of ACh with \( \alpha S248C \) was nearly stopped by the addition of QX-314 (Fig. 5, C and D).

QX-314 retarded the reactions of MTSEA only in the presence of ACh at the two residues on which this was tested, \( \alpha T244C \) at the intracellular end of the channel and \( \alpha L251C \) in the middle of the channel. The time course of the reaction of \( \alpha T244C \) with MTSEA in the presence of ACh, first with, and then without, QX-314 is shown in Fig. 6A. The rate of reaction was faster after QX-314 was removed. On the other hand, the rate of reaction of MTSEA in the absence of ACh was not changed by the addition of QX-314 (Fig. 6B). Similar results were obtained with \( \alpha L251C \).

The extent of the retardation of the MTSEA reaction depended on the concentration of blocker. This is illustrated by the rate constants of the reactions of MTSEA in the presence of ACh with \( \alpha S248C \) and \( \alpha V255C \), in the additional presence of different concentrations of QX-314 (Fig. 7). To compare the effects of QX-314 and QX-222 on the rates of reaction of the different mutants, we estimated the rate constants for the reactions

Figure 5. The effect of QX-314 on the irreversible inhibition of the ACh-induced current in two mutants. (A and B) \( \alpha L258C \). (C and D) \( \alpha S248C \). A and C show the test responses to ACh between applications of MTSEA in the presence of ACh and in the presence or absence of QX-314. B and D show the test currents as a function of cumulative exposure time to MTSEA. (■) Initial responses and, in C and D, also extra final responses without any further application of MTSEA. (△) MTSEA in the presence of ACh. (○) MTSEA in the presence of both ACh and QX-314. ACh was applied at 25 \( \mu M \) to \( \alpha L258C \) and at 100 \( \mu M \) to \( \alpha S248C \). 1 mM MTSEA was applied in 3 s pulses to \( \alpha L258C \) and 1.5 mM MTSEA was applied in 15 s pulses to \( \alpha S248C \). QX-314 was applied at 1 mM to \( \alpha L258C \) and at 0.2 mM to \( \alpha S248C \). The calibration bars are 3 s and 5 \( \mu A \).
of MTSEA with the blocked state from the observed rate constants in the presence and absence of blocker and from the IC_{50} for the particular blocker and mutant (see Methods). The IC_{50} for a particular blocker and mutant is a function not only of the equilibrium dissociation constant for the blocker binding to the open channel, but also of kinetic constants characterizing the isomerizations of the receptor among closed, open, and desensitized states. Although the IC_{50}s (Fig. 7, arrows) were two to four times smaller than the concentrations of blocker giving half of the maximum retardation of rate, two to four times smaller than the concentrations of blocker giving half of the maximum retardation of rate, the use of the IC_{50} to estimate the fraction of a given mutant blocked at different concentrations of blocker yielded consistent rate constants for the blocked state.

We take as the extent of protection by blocker, 1 - (k_{blocked}/k_{open}). At the extracellular end of M2, αL258C and αE262C were not protected by QX-314 or QX-222 (Fig. 8). At the intracellular end of M2, αE241C, αT244C, and αS248C were nearly completely protected. Between the fully protected and the unprotected residues, αL251C was protected ~70% and αV258C ~60% by both blockers. αS252C was protected 58% by QX-222 and not at all by QX-314.

The reaction of αT244C with an uncharged reagent, MTSEH, was also retarded by QX-314 in the presence of ACh: the protection was 88% ± 2% (n = 5). QX-314

2If MTSEA itself first bound reversibly at its site of reaction, and then reacted, the discrepancy between the IC_{50} for QX-314 and the QX-314 concentration that gave half-maximal retardation of the reaction of MTSEA could be explained by the competition of QX-314 and MTSEA for reversible binding in the channel. However, even at 10 mM MTSEA, the highest concentration tested, MTSEA had no effect on the ACh-induced current mediated by wild-type receptor. In addition, there was no rapidly reversible component of the inhibition by MTSEA of the ACh-induced current mediated by any of the mutants. Hence, MTSEA does not reversibly bind in the channel to an appreciable extent, and thus, before its covalent reaction in the channel, MTSEA is unlikely to compete with the binding of QX-314.

*ND.
had no effect on the rate of the reaction of MTSEH in the absence of ACh.

**Discussion**

We found that QX-314 and QX-222, two quaternary ammonium lidocaine derivatives that act as open-channel blockers, protected channel-lining residues substituted by Cys from reaction with positively charged MTSEA. The protection was partial from αV255C to αL251C and was nearly complete from αS248C to αE241C (Fig. 8). There are a number of possible mechanisms for this protection, including steric hindrance, charge repulsion, and allosteric stabilization of unreactive receptor states.

The last mechanism is not a likely explanation for the observed protection. The rate constants for the reactions of some of the mutants are quite different in different functional states of the receptor, so that the blockers could retard the reactions of these mutants by promoting the less reactive state. Such a mechanism, however, would affect all of the state-dependent and none of the state-independent reaction rates, contrary to our results. The rate constants for the reactions of αE241C, αT244C, αL251C, and αV255C with extracellularly applied MTSEA were much larger in the presence of ACh (in the open and rapid-onset desensitized states) than in the absence of ACh (in the predominantly closed state) (Pascual and Karlin, 1998), and these mutants were protected by QX-314 and QX-222 (Fig. 8). On the other hand, the rate constants for the reactions of αS248C and αS252C with MTSEA were not much different in the presence and absence of ACh, and these mutants were also protected. Conversely, αL258C and αE262C were not protected, even though they reacted much faster in the presence of ACh than in its absence. Thus, the blockers did not act by stabilizing an unreactive state from a remote site.

It is also unlikely that the blockers protected the Cys-substituted residues by promotion of the closed state through direct competition with ACh. Although the concentrations of QX-314 and QX-222 used to protect against MTSEA were high enough that some binding to sites in addition to the open channel was possible (Neher, 1983), there was no protection of αL258C or αE262C, which were far less reactive in the absence than presence of ACh.

The mechanism of protection also could not have been solely electrostatic repulsion. QX-314 protected αT244C against the reaction of the neutral reagent MTSEH. Furthermore, the protection of αS252C by QX-222 and not by QX-314 indicates that the mechanism of protection was not primarily electrostatic.

The most likely mechanism of protection was the steric hindrance of access to the substituted Cys by QX-314 and QX-222. Since αV255C was the protected residue closest to the extracellular end of M2, this residue marks the extracellular end of the binding site for the two blockers. If we compare the dimensions of QX-314 and αM2, configured in the open state as an α helix (Karlin and Akabas, 1995), and if the xylyl moiety of QX-314 overlaps αV255, the triethylammonium group overlaps αS248 (Fig. 9). QX-314 or QX-222 bound in this location would protect the residues from αV255C to αS248C by occluding them directly and would protect αT244C and αE241C by blocking the pathway to these residues from the extracellular medium.

The orientation of the blocker with its positively charged quaternary ammonium group toward the intracellular end of the channel would be favored by the electrostatic potential profile in the channel. The net electrostatic potential at each point in the channel is the sum of the electrical distance times the transmembrane potential and of the intrinsic potential due to fixed charges and dipoles in the protein and lumen of channel. The electrostatic potential in the open channel is ∼75 mV more negative at αS248 than at αV255 (Pascual and Karlin, 1998), so that QX-314 or QX-222 would bind ∼20× more tightly with its charged ammonium group near αS248 than near αV255. The proposed location and orientation of the blocker in the channel (Fig. 9) are the same as those proposed by Charnet et al. (1990).

Assuming the location and orientation of the blocker is as shown in Fig. 9, we can estimate the electrostatic contribution to the binding energy. The net electrostatic potential in the open channel in the vicinity of αS248, with a transmembrane potential of −50 mV, was estimated to be ∼−100 mV relative to the extracellular medium (Pascual and Karlin, 1998). Therefore, the electrostatic contribution to the binding energy per mole due to the quaternary ammonium group of QX-222 or QX-314 binding close to αS248 would be approximately four RT.

The apparent affinity constant for QX-222 estimated from burst durations and open times (Neher, 1983) can be extrapolated to ∼2.5 × 10^4 M^−1 at −50 mV. The affinity of QX-314 is considerably greater than that of QX-222 (Neher and Steinbach, 1978). The free energy per mole of QX-222 binding would be ∼10 RT. Thus, hydrophobic and van der Waals interactions would contribute approximately six RT to the binding of QX-222 and more to the binding of QX-314. The side chains of αL251 and αV255, the aligned side chains in β, and possibly the aligned side chains in γ and δ, are all more exposed in the open than in the closed channel (Akabas et al., 1994; Pascual and Karlin, 1998; Zhang and Karlin, 1998). These hydrophobic side chains are likely to interact with bound blocker (Fig. 9).

Although the mutation of (α7)L247 (aligned with
Figure 9. A model of QX-314 binding in the channel between two αM2 segments. The M2 segments are drawn as α helices. The dots represent the van der Waals surfaces. The drawing was made on a Silicon Graphics workstation running Insight II.
αL251) to Thr rendered the neuronal, homopentameric α7 complex insensitive to QX-222 (Revah et al., 1991), the mutation in mouse-muscle receptor of the two αL251 to Cys, and the mutation of the two αV255 to Cys, increased the IC₅₀ for QX-314 but decreased the IC₅₀ for QX-222 (Table I). The effects of these mutations on IC₅₀ may have been confounded by underlying effects of the mutations on the kinetics of state transitions.

The patterns of protection indicate that QX-314 and QX-222 bind at the same depth in the channel. Yet for wild-type receptor, the electrical distances estimated from the fits of the IC₅₀s to the Woodhull equation were quite different, 0.35 for QX-314 and 0.80 for QX-222. The IC₅₀, however, depends not only on the kinetics of binding to the open channel but also on the kinetics of the transitions of the receptor among states. Also, the IC₅₀ was determined after considerable fast desensitization had occurred. Thus, the voltage dependence of the IC₅₀ does not have a simple interpretation. The complexity of the IC₅₀ could partly account for the variation we observed in electrical distance from 0.28 to 0.57 for QX-314 and from 0.34 to 0.68 for QX-222 among the Cys-substitution mutants. Two different single-channel analyses of QX-222 blocking gave electrical distances of 0.78 (Neher and Steinbach, 1978) and ~0.4 (Neher, 1983). Voltage-jump–relaxation analysis yielded electrical distances from 0.65 to 0.75 for QX-222 (Charnet et al., 1990) and 0.72 for QX-314 (Horn and Brodwick, 1980). In general, quite disparate electrical distances for different ACh receptor channel blockers have been reported (e.g., Table I in Sanchez et al., 1986); whether these reflect different binding sites in the channel or other complications is not known. The electrical distance estimated by the Woodhull (1973) equation may not be a consistent marker of position in the channel.

The protection of αS252C by QX-222 and not by QX-314 (Fig. 8) was unexpected and may indicate that the two blockers do not bind in exactly the same configuration. The orientation of the xylyl ring structure shown in Fig. 9 is drawn roughly in the plane containing the two αM2 segments; the M2 segments and β, γ, and δ subunits are not shown. Because the quaternary ammonium group is symmetrical, the somewhat rigid blocker molecule can rotate around its axis, which would maintain similar interactions of the ammonium group but change the subunit interactions of the xylyl ring. Different distributions of axial rotation for the two blockers could be the basis for the different protection of αS252C. In addition, the rotation of the xylyl group and the presumed widening of the channel toward the extracellular end could account for the incompleteness of the protection of αL251C and αV255C, as well as of αS252C.

In the absence of ACh, QX-314 did not protect αT244C or αL251C. αT244C is the mutant that reacts most rapidly with MTSEA, both in the presence and absence of ACh, and the inhibition due to the reaction is complete (Pascual and Karlin, 1998). αT244C is at the intracellular end of the channel, so that QX-314 binding above would protect αT244C from extracellular MTSEA coming through the channel. Thus, αT244C affords a sensitive test for protection. The lack of protection in the absence of ACh indicates that QX-314 did not bind in the closed channel at the concentrations tested.

The poor binding of QX-314 to the closed channel is not due to the obstruction of the channel by the activation gate, which we have located between αG240 and αT244, at the intracellular end of the channel (Wilson and Karlin, 1998). Notwithstanding, there could be a partial obstruction at the extracellular end of the closed channel that would limit access by channel blockers but not by the smaller methanethiosulfonates used to locate the gate or by inorganic cations. Alternatively, the blockers could bind poorly in the closed channel because the net electrostatic potential is ~100 mV less negative in the closed than in the open channel (Pascual and Karlin, 1998), and the side chains of αL251 and αV255 and of the aligned residues in the other subunits are less exposed (Akabas et al., 1994; Pascual and Karlin, 1998; Zhang and Karlin, 1998). Restating this, we suggest that more favorable electrostatic and hydrophobic interactions, rather than greater accessibility, could account for the stronger binding of blockers in the open than in the closed channel. Some of the complexities of local anesthetic action on the voltage-gated Na⁺ channel have been explained on the basis of different intrinsic local-anesthetic binding affinities in different states (the modulated receptor model); however, in contrast to the ACh receptor, in the Na⁺ channel, accessibility to the binding site controlled by the activation and inactivation gates is crucial for the binding of quaternary ammonium derivatives of local anesthetics like QX-314 (Hille, 1977; Ragsdale et al., 1994).

We thank Drs. Myles Akabas, Jonathan Javitch, and Gary Wilson for comments on the manuscript. We are also grateful to Gilda Salazar-Jimenez for oocyte culture.

This research was supported by research grants to A. Karlin from the National Institutes of Health (NIH) (NS-07065), the Muscular Dystrophy Association, and the McKnight Endowment Fund for Neuroscience. During part of this work, J.M. Pascual...
REFERENCES

Adams, P.R. 1976. Drug blockade of open end-plate channels. J. Physiol. (Camb.), 260:539–552.
Akabas, M.H., and A. Karlin. 1995. Identification of acetylcholine receptor channel-lining residues in the M1 segment of the alpha subunit. Biochemistry, 34:12496–12500.
Akabas, M.H., C. Kaufmann, P. Archdeacon, and A. Karlin. 1994. Identification of acetylcholine receptor channel-lining residues in the entire M2 segment of the alpha subunit. Neuron, 13:919–927.
Akabas, M.H., D.A. Stauffer, M. Xu, and A. Karlin. 1992. Acetylcholine receptor channel structure probed in cysteine-substitution mutants. Science, 258:307–310.
Anand, R., W.G. Conroy, R. Schoepfer, P. Whiting, and J. Lindstrom. 1991. Neuronal nicotinic acetylcholine receptors expressed in Xenopus oocytes have a pentameric quaternary structure. J. Biol. Chem., 266:11192–11198.
Ascher, P.R., A. Marty, and T.O. Neild. 1978. The mode of action of antagonists of the excitatory response to acetylcholine in Aplysia neurones. J. Physiol. (Camb.), 278:207–235.
Boyd, N.D., and J.B. Cohen. 1984. Desensitization of membrane-bound Torpedo acetylcholine receptor by amine noncompetitive antagonists and aliphatic alcohols: studies of [3H]acetylcholine binding and Na+ ion fluxes. Biochemistry, 23:4029–4033.
Clapham, D.E., and E. Neher. 1984. Trifluoperazine reduces inward ion currents and secretion by separate mechanisms in bovine chromaffin cells. J. Physiol. (Camb.), 353:541–564.
Colquhoun, D., and R.E. Sheridan. 1981. The modes of action of gallamine. Proc. R. Soc. B Biol. Sci., 211:181–203.
Cooper, E., S. Couturier, and M. Ballivet. 1991. Pentameric structure and subunit stoichiometry of a neuronal nicotinic acetylcholine receptor. Nature, 350:235–238.
DiPaola, M., P.N. Kao, and A. Karlin. 1990. Mapping the alpha subunit site photolabeled by the noncompetitive inhibitor [3H]quinacrine azide in the active state of the nicotinic acetylcholine receptor. Neuron, 4:87–95.
Farley, J.M., J.Z. Yeh, S. Watanabe, and T. Narahashi. 1981. End-plate channel block by guanidine derivatives. J. Gen. Physiol, 77:273–293.
Gage, P.W., and R.E. Wachtel. 1984. Some effects of procaine at the toad end-plate are not consistent with a simple channel-blocking model. J. Physiol. (Camb.), 346:351–359.
Gross, A., and R. MacKinnon. 1996. Agitoxin footprinting the Shaker potassium channel pore. Neuron, 16:399–406.
Heidmann, T., R.E. Oswald, and J.P. Changeux. 1983. Multiple sites of action for noncompetitive blockers on acetylcholine receptor rich membrane fragments from Torpedo marmorata. Biochemistry, 22:3112–3127.
Hille, B. 1977. Local anesthetics: hydrophilic and hydrophobic pathways for the drug-receptor reaction. J. Gen. Physiol, 69:497–515.
Horn, R., and M.S. Brodwick. 1980. Acetylcholine-induced current in perfused rat myoballs. J. Gen. Physiol, 75:397–392.
Horn, R., M.S. Brodwick, and W.D. Dickey. 1980. Asymmetry of the acetylcholine channel revealed by quaternary anesthetics. Science, 210:205–207.
Hucho, F., W. Oberthür, and F. Lotthipe. 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.
Imoto, K., C. Busch, B. Sakmann, M. Mishina, T. Konno, J. Nakai, H. Bujo, Y. Mori, K. Fukuda, and S. Numa. 1988. Rings of negatively charged amino acids determine the acetylcholine receptor channel conductance. Nature, 335:645–648.
Karlin, A. 1989. Explorations of the nicotinic acetylcholine receptor. Harvey Lect, 85:71–107.
Karlin, A., and M.H. Akabas. 1995. Toward a structural basis for the function of nicotinic acetylcholine receptors and their cousins. Neuron, 13:1231–1244.
Karpen, J.W., H. Aoshima, L.G. Abood, and G.P. Hess. 1982. Cocaine and phenacyclidine inhibition of the acetylcholine receptor: analysis of the mechanisms of action based on measurements of ion flux in the millisecond-to-minute time region. Proc. Natl. Acad. Sci. USA, 79:2509–2513.
Kirsch, G.E., M. Alam, and H.A. Hartmann. 1994. Differential effects of sulfhydryl reagents on saxitoxin and tetrodotoxin block of voltage-dependent Na channels. Biophys. J, 67:2305–2315.
Krodel, E.K., R.A. Beckman, and J.B. Cohen. 1979. Identification of a local anesthetic binding site in nicotinic post-synaptic membranes isolated from Torpedo marmorata electric tissue. Mol. Pharmacol, 15:294–312.
Leonard, R.J., C.G. Labarca, P. Charnet, N. Davidson, and H.A. Lester. 1988. Evidence that the M2 membrane-spanning region lines the ion channel pore of the nicotinic receptor. Science, 242:1578–1581.
Lurtz, M.M., M.L. Harelund, and S.E. Pedersen. 1997. Quinacrine and ethidium bromide bind the same locus on the nicotinic acetylcholine receptor from Torpedo californica. Biochemistry, 36:2068–2075.
Magleby, K.L., and B.S. Pallotta. 1981. A study of desensitization of acetylcholine receptors using nerve-released transmitter in the frog. J. Physiol. (Camb.), 316:225–250.
Maleque, M.A., C. Souccar, J.B. Cohen, and E.X. Albuquerque. 1982. Mepropradifen reaction with the ionic channel of the acetylcholine receptor: potentiation of agonist-induced desensitization at the frog neuromuscular junction. Mol. Pharmacol, 22:636–647.
Neher, E. 1983. The charge carried by single-channel currents of rat cultured muscle cells in the presence of local anaesthetics. J. Physiol. (Camb.), 339:663–678.
Neher, E., and J.H. Steinbach. 1978. Local anaesthetics transiently block currents through single acetylcholine-receptor channels. J. Physiol. (Camb.), 277:153–176.
Oswald, R.E., T. Heidmann, and J.P. Changeux. 1983. Multiple affinity states for noncompetitive blockers revealed by [3H]phenacyclidine binding to acetylcholine receptor rich membrane fragments from Torpedo marmorata. Biochemistry, 22:3128–3136.
Pascual, J.M., and A. Karlin. 1998. State-dependent accessibility and electrostatic potential in the channel of the acetylcholine recep-
tor: inferences from rates of reaction of thiosulfonates with substituted cysteines in the M2 segment of the alpha subunit. J. Gen. Physiol. 111:717–739.

Pascual, J.M., C.C. Shieh, G.E. Kirsch, and A.M. Brown. 1995. Multiple residues specify external tetraethylammonium blockade in voltage-gated potassium channels. Biophys. J. 69:428–434.

Pedersen, S.E., S.D. Sharp, W.S. Liu, and J.B. Cohen. 1992. Structure of the noncompetitive antagonist-binding site of the Torpedo nicotinic acetylcholine receptor. [3H]Meproadifen mustard reacts selectively with alpha-subunit Glu-262. J. Biol. Chem. 267:10489–10499.

Ragsdale, D.S., J.C. McPhee, T. Scheuer, and W.A. Catterall. 1994. Molecular determinants of state-dependent block of Na+ channels by local anesthetics. Science. 265:1724–1728.

Revah, F., D. Bertrand, J.L. Galzi, A. Devillers-Thiery, C. Mulle, N. Hussy, S. Bertrand, M. Ballivet, and J.P. Changeux. 1991. Mutations in the channel domain alter desensitization of a neuronal nicotinic receptor. Nature. 353:846–849.

Revah, F., J.L. Galzi, J. Giraudat, P.Y. Haumont, F. Lederer, and J.P. Changeux. 1990. The noncompetitive blocker [3H]chlorpromazine labels three amino acids of the acetylcholine receptor gamma subunit: implications for the alpha-helical organization of regions III and for the structure of the ion channel. Proc. Natl. Acad. Sci. USA. 87:4675–4679.

Ruff, R.L. 1977. A quantitative analysis of local anesthetic alteration of miniature end-plate currents and end-plate current fluctuations. J. Physiol. (Camb.). 264:89–124.

Sanchez, J.A., J.A. Dani, D. Siemen, and B. Hille. 1986. Slow permeation of organic cations in acetylcholine receptor channels. J. Gen. Physiol. 87:985–1001.

Schreibmayer, W., H.A. Lester, and N. Dascal. 1994. Voltage clamping of Xenopus laevis oocytes utilizing agarose-cushion electrodes. Pflügers Arch. 426:453–458.

Sine, S.M., and P. Taylor. 1982. Local anesthetics and histrionicotoxin are allosteric inhibitors of the acetylcholine receptor. Studies of clonal muscle cells. J. Biol. Chem. 257:8106–8114.

Stauffer, D.A., and A. Karlin. 1994. Electrostatic potential of the acetylcholine binding sites in the nicotinic receptor probed by reactions of binding-site cysteines with charged methanethiosulfonates. Biochemistry. 33:6840–6849.

Tamamizu, S., A.P. Todd, and M.G. McNamee. 1995. Mutations in the M1 region of the nicotinic acetylcholine receptor alter the sensitivity to inhibition by quinacrine. Cell. Mol. Neurobiol. 15:427–438.

Wilson, G.G., and A. Karlin. 1998. The location of the gate in the acetylcholine receptor channel. Neuron. 20:1269–1281.

Woodhull, A.M. 1973. Ionic blockage of sodium channels in nerve. J. Gen. Physiol. 61:687–708.

Xu, M., D.F. Covey, and M.H. Akabas. 1995. Interaction of picrotoxin with GABA(A) receptor channel-lining residues probed in cysteine mutants. Biophys. J. 69:1858–1867.

Zhang, H., and A. Karlin. 1997. Identification of acetylcholine receptor channel-lining residues in the M1 segment of the beta subunit. Biochemistry. 36:15856–15864.

Zhang, H., and A. Karlin. 1998. Contribution of the beta subunit M2 segment to the ion-conducting pathway of the acetylcholine receptor. Biochemistry. 37:7952–7964.
