Location of the Polyamine Binding Site in the Vestibule of the Nicotinic Acetylcholine Receptor Ion Channel

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To map the structure of a ligand-gated ion channel, we used the photolabile polyamine-containing toxin MR44 as photoaffinity label. MR44 binds with high affinity to the nicotinic acetylcholine receptor in its closed channel conformation. The binding stoichiometry was two molecules of MR44 per receptor monomer. Upon UV irradiation of the receptor-ligand complex, 125I-MR44 was incorporated into the receptor α-subunit. From proteolytic mapping studies, we conclude that the site of 125I-MR44 cross-linking is contained in the sequence aHis-186 to a.Leu-199, which is part of the extracellular domain of the receptor. This sequence partially overlaps in its C-terminal region with one of the three loops that form the agonist-binding site. The agonist carbachol and the competitive antagonist α-bungarotoxin had only minor influence on the photocross-linking of 125I-MR44. The site where the hydrophobic head group of 125I-MR44 binds must therefore be located outside the zone that is sterically influenced by agonist bound at the nicotinic acetylcholine receptor. In binding and photocross-linking experiments, the luminal noncompetitive inhibitors ethidium and triphenylmethylphosphonium were found to compete with 125I-MR44. We conclude that the polyamine moiety of 125I-MR44 interacts with the high affinity noncompetitive inhibitor site deep in the channel of the nicotinic acetylcholine receptor, while the aromatic ring of this compound binds in the upper part of the ion channel (i.e. in the vestibule) to a hydrophobic region on the α-subunit that is located in close proximity to the agonist binding site. The region of the α-subunit labeled by 125I-MR44 should therefore be accessible from the luminal side of the vestibule.

The nicotinic acetylcholine receptor (nAChR), 1 a prototypical member of the superfamily of ligand-gated ion channels, is an integral transmembrane protein with the subunit stoichiometry α2βγδ (1–3). Each receptor subunit contains four hydrophobic sequences, which are presumed to span the plasma membrane (4, 5). The large N-terminal domain and the relatively short C-terminal part of the subunits are oriented toward the extracellular side. A large connecting loop, which is found between transmembrane sequences M3 and M4, is assumed to extend into the cytoplasm. The five subunits contribute their homologous M2 sequences to the formation of the ion channel (6), which is permeable for cations upon agonist binding. A selectivity filter formed by the five M2 helices contributes to the cation conductance properties of the channel. Three rings of negatively charged amino acid residues (7, 8) located at the constriction of the channel and on the cytoplasmic and the extracellular side of this constriction, respectively, in particular are of functional importance.

In the absence of crystals suitable for x-ray analysis, the three-dimensional structure of nAChR is investigated mainly by three approaches: electron microscopy (9, 10), site-directed mutagenesis in combination with patch clamp electrophysiology (e.g. Refs. 7, 11, and 12), and affinity labeling (e.g. Refs. 1 and 13–16). Two binding sites for agonists and competitive antagonists are located in the extracellular region, mainly on each of the two α-subunits (1) at the α-δ and α-γ interfaces (14, 17, 18). A binding site for noncompetitive inhibitors (NCIs), such as chlorpromazine, ethidium bromide, and triphenylmethylphosphonium (TPMP+), has been found within the channel lumen (6, 19). These NCIs are assumed to enter the ion channel from the extracellular side and to bind deep in the channel lumen, thereby inhibiting the ion flow. Photoaffinity labels derived from well characterized NCIs have been developed to characterize the structure of the nAChR ion channel. [3H]Chlorpromazine and [3H]TPMP are preferentially photocross-linked to amino acid residues within the M2 transmembrane sequence of the desensitized receptor, thus demonstrating that these compounds bind deep in the ion channel and close to the selectivity filter (6, 19).

Phianthotoxin-433 (PhTX-433) is a neuroactive, polyamine-containing toxin found in the venom of the digger wasp Philanthus triangulum (20). Synthetic analogues of this polyamine amide, such as PhTX-343, have been shown to noncompetitively antagonize a range of ionotropic receptors (21), including nAChR (22–26). These low molecular weight compounds have a hydrophobic head group linked to a polyamine tail. At physiological pH, they are highly positively charged and, therefore, should bind to any surface with a corresponding distribution of anionic functionalities (21). The binding affinities of these compounds to nAChR are significantly influenced by modifying

1 The abbreviations used are: nAChR, nicotinic acetylcholine receptor; TPMP+, triphenylmethylphosphonium; NCI, noncompetitive inhibitor; PhTX-433, phosphanothix-433; α-BTX, α-bungarotoxin; ACh, acetylcholine; Tricine, N-[2-hydroxy-1,1-bis(hydroxymethyl)ethyl]glycine; HPLC, high pressure liquid chromatography; PAGE, polyacrylamide gel electrophoresis.

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‡ The abbreviations used are: nAChR, nicotinic acetylcholine receptor; TPMP+, triphenylmethylphosphonium; NCI, noncompetitive inhibitor; PhTX-433, phosphanothix-433; α-BTX, α-bungarotoxin; ACh, acetylcholine; Tricine, N-[2-hydroxy-1,1-bis(hydroxymethyl)ethyl]glycine; HPLC, high pressure liquid chromatography; PAGE, polyacrylamide gel electrophoresis.

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their structural elements (26–28). We have synthesized a series of polyamine-containing analogues of PhTX-343 in the search for a ligand with high affinity and specificity for \textit{Torpedo californica} nAChR for photocross-linking studies. This approach resulted in the discovery of a photoactivatable compound, MR44, which binds to the nAChR with high affinity (26, 29).

In the present work, we showed that two molecules of MR44 bind with high affinity in the lumen of the nAChR ion channel. Using \textsuperscript{125}I-MR44 as a photoaffinity label, we localized the site of interaction of the aromatic head group of MR44 in the vestibule of the ion channel. The sequence that was labeled by \textsuperscript{125}I-MR44 was found on the \( \alpha \)-subunit close to, but not over, the agonist-binding site. In addition, we found that bound MR44 was displaced by luminal NCIs and calcium, suggesting that the positively charged polyamine moiety of MR44 binds deep in the channel lumen at the high affinity NCI site.

**EXPERIMENTAL PROCEDURES**

**Materials**—Liquid nitrogen-frozen tissue from \textit{T. californica} was supplied by C. Winkler (Aquatic Research Consultants, Sa Pedro). 

Carbachol, calcium chloride, ethidium bromide, and HEPES were from Sigma. Dithiothreitol, TPMP\(^*\), and chloramphenicol were from Aldrich. K\(^{125}\)I was from Amersham Pharmacia Biotech. \textsuperscript{125}I-Labeled \( \alpha \)-bungarotoxin (\( \alpha \)-BTX) was purchased from PerkinElmer Life Sciences. ArgC protease, AspN protease, LysC protease, endoglycosidase H (Endo H), and V8 protease were obtained in sequencing grade from Roche Molecular Biochemicals (Mannheim, Germany).

**Synthesis and Purification of \textsuperscript{125}I-Labeled MR44**—\textsuperscript{125}I-MR44 was radioactively iodinated with \textsuperscript{125}I using the chloramine T method (31). The monoo-\textsuperscript{125}I derivative was isolated by reverse-phase HPLC (Waters model 626, Eschborn, Germany) on a Vydac C\(_{18}\) column applying the following linear gradient (1 ml/min): solvent A (aqueous solution containing 0.1% trifluoroacetic acid) and solvent B (acetonitrile containing 0.085% trifluoroacetic acid). The UV absorption of the eluent was determined at 305 nm, and the radioactivity of each fraction was detected using a \( \gamma \)-counter. \textsuperscript{125}I-MR44 was characterized by matrix-assisted laser desorption-ionization mass spectrometry.

**\textsuperscript{125}I-MR44 Binding Assay**—\( \alpha \)-ChR-rich membranes were prepared from frozen \textit{T. californica} electric organ as described earlier (32). Increasing concentrations of \textsuperscript{125}I-MR44 (5,000 cpm/nmol) were added to a constant amount of \( \alpha \)-ChR-rich membranes (10 mg of nAChR [0.25 mg/ml in 0.1 M NaPi buffer, pH 7.4; total volume per sample, 200 \( \mu \)l) containing 0.5 mM protein, diluted in 100 mM NaPi, pH 7.4; total volume per sample, 200 \( \mu \)l) and were incubated for 45 min at room temperature. Bound ligand was separated from the free ligand by ultracentrifugation in a Beckmann tabletop ultracentrifuge for 10 min at 80,000 \( \times \) g and 4 \( ^\circ \)C. Aliquots were withdrawn prior to centrifugation to determine the total radioactivity, and duplicate aliquots of the supernatant were removed after centrifugation (as the free ligand in supernatant) to determine the nonspecific binding. As a control, buffer was added instead of NCI or carbachol (500 \( \mu \)l of 10 mM NaPi buffer). 

**Photocross-linking Experiments**—\( \alpha \)-ChR-rich membranes (50 \( \mu \)g) were diluted in 0.1 M NaPi, pH 7.4, to a final receptor concentration of 140 nM. After the addition of carbachol (500 \( \mu \)l), the samples were incubated for 30 min at room temperature. Subsequently, TPMP\(^*\), ethidium, \( \alpha \)-BTX, or unlabeled MR44 was added, and the samples were incubated for further 30 min at room temperature. The radioactive \textsuperscript{125}I-MR44 (10 \( \mu \)M; 250,000 cpm/nmol) was mixed with the sample solution and irradiated with UV light at 254 nm (distance, 15 cm; quartz lamp; Desaga, Heidelberg, Germany) for 15 s. Longer irradiation times resulted in a significant loss of label, presumably because the aromatic group of MR44 releases \textsuperscript{125}I, and in irreversible damage of the nAChR, resulting in high molecular weight aggregates of the receptor (data not shown). Unbound \textsuperscript{125}I-MR44 was separated from \textsuperscript{125}I-MR44 bound to \( \alpha \)-ChR-rich membranes by centrifugation (15,000 \( \times \) g, 20 min, 4 \( ^\circ \)C). The pellet was dissolved and separated by SDS-PAGE using a 10% SDS-PAG (35). The stained gel was dried, and radioactive receptor subunits were visualized by autoradiography.

**Deglycosylation of nAChR Using Endo H**—180 \( \mu \)g of \( \alpha \)-ChR-rich membranes were centrifuged after labeling with \textsuperscript{125}I-MR44 and resuspended in 50 \( \mu \)l of 100 mM NaPi buffer, pH 6.5. 5 \( \mu \)l of 1% SDS was added to this suspension, followed by 10 milliunits of Endo H in 3 \( \mu \)l of 0.5 M NaPi buffer. The pellet was dissolved and separated by SDS-PAGE using a 10% SDS-PAGE (35). The stained gel was dried, and radioactive receptor subunits were visualized by autoradiography.

**Cell Culture**—TR671 cells were maintained in Dulbecco’s modified Eagle’s medium containing 4.5 \( \mu \)liter glucose and supplemented with 10% (v/v) fetal calf serum, 1 mM pyruvic acid, 4 mM glutamine, 10 units/ml penicillin, and 10 \( \mu \)g/ml streptomycin and incubated at 37 \( ^\circ \)C in a 5% CO\(_2\) atmosphere. Cells were divided 1:10 when they were 75% confluent. For electrophysiology, cells were grown on glass coverslips (5–20 mm) in 35-mm Petri dishes and perfused with a perfusion buffer that mounted on the stage of an inverted microscope.

**Electrophysiology**—The whole-cell patch-clamp configuration was used to record whole cell currents evoked by acetylcholine (ACh). Patch pipettes were fabricated from borosilicate glass capillaries (GC150–10; Clarke Electromedical Instruments) using a Sutter (P-97) programmer. Pipette resistances were ~5 megohms when filled with 140 mM NaCl, 1 mM CaCl\(_2\), 1 mM MgCl\(_2\), 11 mM EGTA, and 5 mM HEPES (pH 7.2, adjusted with CsOH). The cells were constantly perfused with rat saline containing 135 mM NaCl, 5.4 mM KCl, 1 mM CaCl\(_2\), 1 mM MgCl\(_2\), and 5 mM HEPES (pH 7.4, adjusted with NaOH). Membrane currents were monitored using a List Electromedical LM-EPCT patch clamp amplifier. The patch clamp and DAD-12 Superfusion system were controlled by pClamp 5.7.2 software (Axon Instruments), which simultaneously acquired data to the hard disc of an IBM-compatible PC. Concentration-inhibition relationships for MR44 were measured to determine the IC\(_{50}\) value for inhibition of the peak current evoked by ACh (10 \( \mu \)M) at holding potentials (V\(_{h}\)) of −25, −50, and 100 mV. Experiments were performed at 18–22 \(^\circ\)C. All data analyses were performed on PC using pClamp software (Axon Instruments). Curve fitting was performed using Graphpad Prism software. IC\(_{50}\) values were determined by fitting a four-parameter logistic equation to the concentration-inhibition/response data. p values were determined by the unpaired Student’s t test, and differences were considered to be significant for p < 0.05.

**Polyamine Binding Site at the nAChR**

Polyamine-containing analogues of PhTX-343 in the series that was labeled by \textsuperscript{125}I-MR44 was found on the \( \alpha \)-subunit close to, but not over, the agonist-binding site. In addition, we found that bound MR44 was displaced by luminal NCIs and calcium, suggesting that the positively charged polyamine moiety of MR44 binds deep in the channel lumen at the high affinity NCI site.
Polyamine Binding Site at the nAChR

RESULTS

High Affinity Binding of Two Molecules of $^{125}$I-MR44 per nAChR Monomer—Binding of the radioactively labeled polyamine-containing toxin MR44 (Fig. 1A) to the nAChR was investigated (Fig. 1, B and C). $^{125}$I-MR44 was purified by reverse-phase HPLC and analyzed by matrix-assisted laser desorption-ionization mass spectrometry (data not shown). Specific binding of $^{125}$I-MR44 to nAChR-rich membranes from T. californica was determined by subtracting the nonspecific component from the total binding curve (Fig. 2B). Nonspecific binding was measured in the presence of a 100-fold molar excess of nonradioactive I-MR44. As shown in Fig. 1B, $B_{\text{max}}$ was obtained at 2.11 ± 0.38 μM (n = 4). The Scatchard plot shows a straight line indicating that $^{125}$I-MR44 binds to a single class of noninteracting binding sites with a $K_{\text{app}}$ value of (175 mM Tris/HCl, 0.1% (w/v) SDS, 5% (v/v) glycerol, pH 6.8), and incubated for 10 min at 70 °C. 4 μg of V8 protease were dissolved in 6 μl of loading buffer and added to the α-subunit. The entire sample was loaded immediately onto the stacking gel. Stacking and proteolysis were carried out for 30 min at 12 mA. Then the current was shut off for 30 min to allow further digestion by the protease, after which electrophoresis was continued. The separating gel contained 15% (w/v) acrylamide to allow adequate separation of the low molecular weight cleavage products. After Coomassie staining, the gel was dried, and radioactive peptides were visualized by autoradiography.

Proteolytic Mapping Using LysC Protease—Since the proteolytic activity of AspN protease is drastically reduced in the presence of 0.1% (w/v) SDS in the electrophoresis buffer (37), proteolytic cleavage was carried out in solution prior to electrophoretic separation of the proteolytic peptides using the Tricine gel system described by Schagger and von Jagow (38). 25 μg of $^{125}$I-MR44-labeled α-subunit in 100 mM NaF, pH 7.5, were incubated with 2 μg of LysC protease for 27 h at 37 °C (40). Sample preparation, Tricine gel electrophoresis, staining of the peptides, and detection of labeled peptides were performed as described above.

Peptide Sequencing by Edman Degradation—For determination of N-terminal amino acid sequences, the peptides obtained by V8 digestion or AspN digestion and subsequent SDS-PAGE or Tricine-polyacrylamide gel electrophoresis, respectively, of $^{125}$I-MR44-labeled α-subunit were blotted onto polyvinylidene difluoride membrane. After Coomassie staining, the peptides of interest were excised and submitted to Edman degradation. Protein sequence analysis was performed using a Type 473A protein sequencer (Applied Biosystems).
AChR. nAChR-rich membranes were incubated with 125I-MR44 (9 μM) in the presence of increasing concentrations of the luminal NCI ethidium. The IC₅₀ value was determined to be 20.4 ± 3.6 μM (n = 3), B, calcium prevented binding of 125I-MR44 to the nAChR. nAChR-rich membranes were incubated with 125I-MR44 (9 μM) in the presence of increasing concentrations of calcium with (●) and without (▲) the agonist carbachol (n = 3). According to a logarithmic equation described by Herz et al. (34), the K₅₀ values for calcium binding were calculated to be 12.3 ± 1.8 μM, and in the presence of carbachol they were 14.6 ± 2.2 μM.

0.82 ± 0.22 μM. In the presence of the agonist carbachol, which results in nAChR desensitization, the binding affinity of 125I-MR44 was increased by a factor of 1.4, but the P_max value was not changed (data not shown). Using 125I-labeled α-BTX in a binding assay to determine the number of receptor monomers per μg of protein (33), the binding stoichiometry of 125I-MR44 was calculated to be 2.16 ± 0.18 mol of 125I-MR44/mol of nAChR monomer, demonstrating that two molecules of 125I-MR44 bind per receptor monomer.

Voltage-independent Inhibition of nAChR by MR44—The influence of MR44 on the agonist-mediated ion conductance of the nAChR was investigated electrophysiologically using cells that express muscle-type nAChR (cell line TE671). MR44 inhibited ACh-mediated whole-cell currents of TE671 cells with IC₅₀ values of 16.1 ± 4.6 μM (n = 5), 17.5 ± 7.1 μM (n = 6), and 16.9 ± 4.0 μM (n = 7) at V₁₅₀ of -25, -50, and -100 mV, respectively (Fig. 1D).

The NCI Ethidium Displaced Reversibly Bound 125I-MR44—The well characterized luminal NCI ethidium interacts with a binding affinity of 1 μM (41) with the high affinity NCI site of the nAChR. The binding of 125I-MR44 was determined in the presence of increasing concentrations of ethidium bromide. Fig. 2A shows that bound 125I-MR44 was displaced by ethidium with an IC₅₀ value of 20.4 ± 3.6 μM (n = 3). The competitive antagonist α-BTX had no influence on the binding of 125I-MR44 (data not shown).

Calcium Displaced Reversibly Bound 125I-MR44—It was shown previously that the NCI ethidium could be completely displaced by cations, indicating that NCIs and channel-permeating cations bind to the nAChR ion channel in a competitive manner (42). As shown in Fig. 2B, the divalent cation calcium displaced 125I-MR44 from its binding site with an IC₅₀ value of 2.4 mM. According to a logarithmic formula described by Herz et al. (34), the K₅₀ value for calcium binding was calculated to be 12.3 ± 1.8 μM (n = 3). In the presence of carbachol, the affinity of calcium for the MR44 binding site was not significantly changed (K₅₀ = 14.6 ± 2.2 μM).

125I-MR44 Photolabeled the nAChR α-Subunit—For photocross-linking, 125I-MR44 (10 μM) was incubated with nAChR-rich membranes. 125I-MR44 was covalently cross-linked to AChR rich membranes (30 μg each lane) were photolabeled with 10 μM 125I-MR44. Samples were incubated without Endo H (lane 2) and with Endo H (lane 3) and separated on an 8% SDS-polyacrylamide gel. The gel was stained with Coomassie Blue (lanes 1–3), and radioactive protein bands were visualized by autoradiography (lanes 4 and 5). Exposure time was typically 2 days. B, photofinity labeling of nAChR with 125I-MR44. nAChR-rich membranes (50 μg each lane) were photolabeled with 10 μM 125I-MR44 in the absence (lanes 6–9) or in the presence of calcium (lanes 6–9) of 500 μM carbachol and separated on an 10% SDS-polyacrylamide gel. The gel was stained with Coomassie Blue (lane 2 shows 50 μg of nAChR-rich membranes), and radioactive protein bands were visualized by autoradiography (lanes 3–10). Samples were preincubated with 560 nM α-BTX (lanes 4 and 7) or 5 mM TPMP+ (lanes 5 and 8) or 100 μM ethidium (lane 9) or 1 mM MR44 (lane 10). The exposure time was typically 2 days. A and B, lane 1 shows the molecular mass markers in kDa: phosphorylase b (97 kDa), bovine serum albumin (67 kDa), ovalbumin (43 kDa), and carbonic anhydrase (30 kDa). Exposure time was typically 1 day.
Incubation of the 125I-MR44-labeled nACHr with Endo H quantitatively converted the labeled α-subunit to its high mobility form (Fig. 3A, lane 5), indicating that 125I-MR44 was incorporated exclusively into the α-subunit. A 100-kDa protein (Fig. 3A, lane 2; Fig. 3B, lane 2) that most likely represents the Na⁺/K⁺-ATPase, was faintly labeled by 125I-MR44 (Fig. 3A, lane 4; Fig. 3B, lane 3). This protein is found in nACHr-rich membrane preparations as a contamination. It has been shown previously that polyamines modulate the Na⁺/K⁺-ATPase and that other photolabile polyamine derivatives also photolabel this enzyme to a minor extent (26).

In the absence or presence of the agonist carbachol, photocross-linking of 125I-MR44 with nACHr-rich membranes occurred with the receptor being in one of the two closed conformations, i.e., in the resting or in the desensitized state, respectively. In both experiments, the nACHr α-subunit was labeled (Fig. 3B, lanes 3 and 6). In the presence of well-characterized luminal NCIs, such as TPMP⁺ and ethidium bromide, 125I-MR44 bound at the NCI site was completely displaced, with the labeling intensity reduced to background levels (Fig. 3B, lanes 5, 8, and 9). The competitive antagonist α-BTX did not significantly affect the binding of 125I-MR44 (Fig. 3B, lanes 4 and 7). In the presence of α-BTX (lane 4) or carbachol (lane 6), the labeling intensity was reduced by 10–15%. Cross-linking 125I-MR44 in presence of a 100-fold molar excess of the nonradioactive analogue resulted in the complete loss of labeling showing the specificity (saturability) of MR44 binding (Fig. 3B, lane 10). Without irradiation, no cross-linking was observed (data not shown).

Mapping the 125I-MR44-labeled α-Subunit Using V8 Protease—The binding site of MR44 was mapped using the method described by Cleveland et al. (36) using S. aureus V8 protease, which specifically cleaves peptide bonds at the carboxyl side of glutamate residues. Limited in-gel digestion of the α-subunit with V8 protease produces preferentially four nonoverlapping fragments (44). The largest fragment, a 20-kDa peptide (V8-20), begins at αSer-173 and contains the first three membrane-spanning regions, M1, M2, and M3. The 18-kDa peptide (V8-18) is part of the extracellular domain and carries at αAsn-141 a carbohydrate residue of ~4 kDa. The 10-kDa peptide (V8-10) that begins at αAsn-339 contains the fourth membrane-spanning region, M4. The smallest fragment of 4 kDa (V8-4) represents the N-terminal part of the α-subunit.

nACHr-rich membranes were photocross-linked with 125I-MR44, and the labeled α-subunit was isolated by preparative tube gel electrophoresis. The 125I-MR44 labeled α-subunit was cleaved in the gel with V8 protease. Peptides generated were separated using 15% SDS-PAGE, and those fragments carrying a radioactive label were identified by autoradiography (Fig. 4A). Fig. 4A, lanes 2–6, show Coomassie staining of the uncleaved radiolabeled α-subunit (lane 2), a V8 protease digest of 125I-MR44-labeled α-subunit (lanes 3 and 4) and of unlabeled α-subunit (lane 5), and V8 protease alone (lane 6). Using varying enzyme/substrate ratios, the limited proteolysis of the radioactively labeled and the unlabeled α-subunit reproducibly yielded identical peptide patterns. V8 cleavage of the labeled and of the unlabeled α-subunit revealed two prominent cleavage products with apparent molecular masses of about 19 kDa (V8-20) and 17 kDa (V8-18) and a smaller peptide of ~10 kDa (V8-10) (Fig. 4A, lanes 3–5). Additional minor poorly resolved cleavage products were found near the dye front of the gel. V8 protease and its proteolytic fragments appear as two major protein bands with apparent molecular masses of 29 and 27 kDa and as a 14-kDa fragment of lower intensity (Fig. 4A, lane 6). The autoradiograph of the V8-protease digest revealed that the V8-20 fragment carried the majority of the radioactive label (Fig. 4A, lanes 8 and 9). No radioactivity was detected in the αV8-18 and αV8-10 peptides. As expected, in the absence of V8 protease the uncleaved radioactive α-subunit was found in the range of 41 kDa (Fig. 4A, lane 7).

N-terminal Amino Acid Sequencing of 125I-MR44-labeled V8-20 and of Unlabeled V8-18 and V8-10—For the characterization of the peptides generated by V8 digestion, their N-terminal amino acid sequences were identified using Edman degradation. The peptides generated by in-gel V8-protease digestion were transferred to polyvinylidene difluoride membrane and visualized by Coomassie staining. Single peptide bands were excised and submitted to N-terminal amino acid sequencing. Table I shows the results of the first five sequencing cycles for each of the peptides examined. The most prominent N-terminal sequence found in the 125I-MR44-labeled V8-20 peptide band started from αVal-46. As a minor signal, a...
second peptide sequence was observed that most likely begins with αSer-173. The sequence was difficult to detect, since the first two amino acids of the N terminus of this peptide showed barely visible signals in the chromatograph (Tables I and II). Previous studies using limited in-gel proteolysis of the nAChR α-subunit with V8 protease (45) clearly demonstrated that V8-20 reproducibly contained two peptides beginning from αVal-46 and αSer-173, respectively, confirming the presence of the two proteolytic fragments detected in V8-20. The N-terminal sequence of unlabeled V8-18 was found to be identical with one of the V8-20 peptides (Table I). Microsequencing of unlabeled V8-18 revealed an N terminus starting from Asp-200 (Table I). Consequently, 125I-MR44 must be photoincorporated into the 2.5-kDa peptide designated by PEDersen et al. (45). These authors could show that V8-15 corresponded most likely to the deglycosylated form of an incompletely cleaved form of V8-18 that comigrated with V8-20. Also demonstrated that V8-12 was the deglycosylated form of V8-18. The autoradiograph revealed that Endo H incubation had no influence on the mobility of the 125I-MR44 labeled peptide in the gel (Fig. 4B, lanes 4 and 5), demonstrating that the label and the carbohydrate moiety were associated with different V8 proteolytic fragments. These findings indicate that 125I-MR44 was most likely cross-linked to the V8-20 peptide starting from αSer-173.

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**Table I**

**N-terminal amino acid sequence analysis of 125I-MR44-labeled nAChR α-subunit peptides obtained by proteolytic cleavage using V8, AspN, ArgC, or LysC proteases.** Peptides were prepared and sequenced as described under "Experimental Procedures."

"x, no unambiguous assignment possible; amino acid in parentheses, only weak signal detected.
Mapping the 125I-MR44-labeled α-subunit Using LysC Protease—The sequence labeled by 125I-MR44 contains at position 185 a lysine residue. LysC protease, an protease that cleaves the peptide bond after lysine residues (40), was therefore used to cleave the labeled α-subunit. The peptides generated were subsequently separated using Tricine gel electrophoresis. The Coomassie-stained gel showed peptides in the molecular mass range of 7–14 kDa (Fig. 6B). The autoradiograph identified a single radioactive peptide of about 8 kDa (Lys-C-8), while the other peptide generated carried no radioactivity (Fig. 6B, lane 3).

N-terminal Amino Acid Sequencing of 125I-MR44-labeled LysC-8 and of Unlabeled LysC-12, LysC-10, LysC-7, and LysC-6—Sequencing of the N terminus of the radioactively labeled LysC-8 fragment yielded a single sequence starting from His-186 (Tables I and II). The other unlabeled fragments LysC-12, LysC-10, and LysC-7 were found to begin from Asp-C-210 and Asp-C-116, respectively (Table I). The unlabeled 15-kDa peptide was identified as a proteolytic fragment of the ArgC protease. The specificity of ArgC is primarily to arginine residues, although hydrolysis proceeds to a minor degree also after lysine residues (39) and occasionally after aromatic residues (47). As a result, the labeled α-subunit was cleaved at the C-terminal side of α-Lys-116 and α-Tyr-181 (N-terminal side of α-Arg-182). Taken together with the results obtained with V8 and AspN protease, this observation allows us to locate the site of 125I-MR44 cross-linking to the sequence α-Arg-182 to α-Leu-199.

N-terminal amino acid sequence analysis of 125I-MR44-labeled peptides obtained by cleavage of the labeled α-subunit using V8, AspN, ArgC, or LysC proteases. The αV8–20, αAspN-16, αArgC-4, and αLys-C-8 proteolytic peptide carried a radioactive label. Peptides were prepared and sequenced as described under "Experimental Procedures."

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* x, peak was not integrated.
* Indication to one of these two sequences not possible.

Note: The molecular mass markers in kDa on the ordinate: globine II (6.2 kDa), glucagon (3.5 kDa).

Peptide idiotypic mapping of the 125I-MR44-labeled α-subunit using AspN protease. After cross-linking of 125I-MR44 (10 μM) with nAChR (0.2 mg/ml receptor), the 125I-MR44-labeled receptor subunits were separated by preparative SDS-PAGE. The isolated 125I-MR44 labeled α-subunit was incubated overnight with AspN protease, and the generated peptide fragments were separated using Tricine-polyacrylamide gel electrophoresis with 3% spacer and 16% separation. The isolated 125I-MR44-labeled α-subunit was incubated overnight with AspN protease, and the generated peptide fragments were separated using Tricine-polyacrylamide gel electrophoresis with 3% spacer and 16% separation gel. The autoradiograph identified a single radioactive peptide of about 8 kDa (Lys-C-8), while the other peptide generated carried no radioactivity (Fig. 6B, lane 3).
Polyamine Binding Site at the nAChR

The architecture of ligand-gated ion channels can be explored using photoaffinity derivatives of high affinity ligands. The aim of the present studies was to identify amino acid residues facing the ligand-binding site of the nAChR. The novel photoaffinity label MR44 was first characterized using photoaffinity derivatives of high affinity ligands. 125I-MR44 was incubated with nAChR-rich membranes and subsequently irradiated with UV light to generate a reactive nitrile species that efficiently cross-links to any amino acid residues nearby that are facing the binding site. The finding that 125I-MR44 photolabeled the receptor a-subunit suggests that the aromatic head groups of the two MR44 bound in the channel lumen most likely interact each with one of the a-subunits, since this part of MR44 carries the photolabile group.

To localize the polyamine-binding site at the nAChR in its resting closed state, the regions of the a-subunit that incorporated 125I-MR44 were mapped by proteolytic cleavage using V8 (36, 45), AspN (37), ArgC (39), and LysC proteases (40). The N termini of peptides generated were microsequenced and localized in the known primary structure of the Torpedo nAChR a-subunit (4). When using limited V8 proteolysis, the majority of the radioactivity was detected in a 19-kDa proteolytic fragment. Microsequencing and Endo H treatment identified two fragments in the V8-20 peptide band beginning with aSer-Val6 and oSer-173, respectively. These data correspond to earlier findings of Pedersen et al. (45) demonstrating that the V8-20 band contained two comigrating fragments: a carbohydrate-containing peptide starting from aVal-46 and an unglycosylated peptide beginning from oSer-173. Removal of the oligosaccharide moiety by Endo H had no influence on the mobility of the radioactively labeled V8-20 peptide, indicating that 125I-MR44 photolabeled the fragment beginning with oSer-173. To confirm this finding, AspN protease was used to cleave the 125I-MR44 labeled a-subunit at sites different from that cleaved by V8 protease. An AspN proteolytic fragment of 16 kDa was found to carry the radioactive label. Microsequencing of this peptide revealed a single sequence beginning from oAsp-180, confirming the results obtained with V8 protease. The N-terminal sequence of the unlabeled proteolytic fragment migrating at 14 kDa was identified to start from oAsp-200. Thus, the cross-linked 125I-MR44 must be located within a stretch of 20 amino acid residues from oAsp-180 to oLeu-199. Cleavage of the 125I-MR44-labeled a-subunit with ArgC protease generated a peptide pattern with only one peptide labeled of about 4 kDa. N-terminal amino acid sequencing identified two proteolytic fragments starting with Arg-182 and Leu-80, respectively. The peptide beginning with oLeu-80 was shown to be nonradioactive using V8, AspN, and LysC proteases. Consequently, 125I-MR44 must be cross-linked to the ArgC-4 fragment starting with Arg-182, confirming the region of 125I-MR44 photoincorporation suggested from the V8 and AspN digest experiments. The sequence oArg-182 to oLeu-199 labeled by 125I-MR44 contains a lysine residue in position 185. Therefore, LysC protease was used to cleave the labeled a-subunit. In mapping experiments using LysC protease, we further narrowed down the site of modification to the amino acid stretch oHis-186 to oLeu-199.

FIG. 6. Proteolytic mapping of the 125I-MR44-labeled nAChR a-subunit using ArgC protease (A) and LysC protease (B). After cross-linking of 125I-MR44 (10 μM) with nAChR (0.2 mg/ml receptor), the 125I-MR44-labeled receptor subunits were separated by preparative SDS-PAGE. The isolated 125I-MR44-labeled a-subunit was incubated overnight with ArgC protease (A) or LysC protease (B), and the generated peptide fragments were separated using Tricine-polyacrylamide gel electrophoresis with 3% spacer and 16% separation gel. Lane 2 (A and B) shows the Coomassie-stained peptides that were generated by incubation of the 125I-MR44-labeled a-subunit (25 μg) with ArgC protease (A; 2 μg) or LysC protease (B; 2 μg). Lane 3 (A and B) shows the corresponding autoradiograph to locate radioactive peptides (exposure time was typically 2 days). Lane 1 (A and B) shows the molecular mass markers in kDa on the ordinate: globine (16.9 kDa), globine I + II (14.4 kDa), globine I + III (10.7 kDa), globine I (8.2 kDa), globine II (6.2 kDa), and glucagon (3.5 kDa).

DISCUSSION

The architecture of ligand-gated ion channels can be explored using photoaffinity derivatives of high affinity ligands. The aim of the present studies was to identify amino acid residues facing the ligand-binding site of Torpedo californica nAChR and thereby to obtain information on the structure of the luminal surface of the channel gated by this receptor. We have used a newly developed polyamine-containing toxin carrying a hydrophobic head group in photocross-linking experiments. The novel photoaffinity label MR44 was first characterized electrophysiologically and by binding studies and was subsequently used to map the ligand binding site in the lumen of the nAChR ion channel.

Using fluorescent titration, we have shown recently that PhTX analogues including MR44 interact with the fluorescent NCI ethidium bound to the high affinity NCI site in the desensitized state of the nAChR (26). Like most amine NCIs containing aromatic or aliphatic rings, such as chlorpromazine, meproadifen, and TPMP+, MR44 was found to bind with higher affinity in the presence of the agonist carbachol, i.e. when the nAChR is in its desensitized closed state, as compared with the resting closed conformation. Since MR44 causes a voltage-independent block of AChR-induced ion flow, MR44 presumably acts as inhibitor of the closed channel conformation rather than as an open channel blocker. Unlike other classical NCIs, which show a binding stoichiometry of 1:1, we found that two molecules of MR44 were bound per receptor monomer. In a recent study, the same binding stoichiometry was observed for another PhTX derivative, N2-phenyl-125I-PbTX-343-lysine (26). MR44 has two functional moieties; a long positively charged polyamine tail on the one side and an aromatic head group on the other. It is most likely that these two structural elements of the molecule, although both contribute to the binding affinity, bind at different sites within the receptor lumen.

For photocross-linking, 125I-MR44 was incubated with nAChR-rich membranes and subsequently irradiated with UV light to generate a reactive nitrile species that efficiently cross-links to any amino acid residues nearby that are facing the binding site. The finding that 125I-MR44 photolabeled the receptor a-subunit suggests that the aromatic head groups of the two MR44 bound in the channel lumen most likely interact each with one of the a-subunits, since this part of MR44 carries the photolabile group.

To localize the polyamine-binding site at the nAChR in its resting closed state, the regions of the a-subunit that incorporated 125I-MR44 were mapped by proteolytic cleavage using V8 (36, 45), AspN (37), ArgC (39), and LysC proteases (40). The N termini of peptides generated were microsequenced and localized in the known primary structure of the Torpedo nAChR a-subunit (4). When using limited V8 proteolysis, the majority of the radioactivity was detected in a 19-kDa proteolytic fragment. Microsequencing and Endo H treatment identified two fragments in the V8-20 peptide band beginning with oVal-46 and oSer-173, respectively. These data correspond to earlier findings of Pedersen et al. (45) demonstrating that the V8-20 band contained two comigrating fragments: a carbohydrate-containing peptide starting from oVal-46 and an unglycosylated peptide beginning from oSer-173. Removal of the oligosaccharide moiety by Endo H had no influence on the mobility of the radioactively labeled V8-20 peptide, indicating that 125I-MR44 photolabeled the fragment beginning with oSer-173. To confirm this finding, AspN protease was used to cleave the 125I-MR44 labeled a-subunit at sites different from that cleaved by V8 protease. An AspN proteolytic fragment of 16 kDa was found to carry the radioactive label. Microsequencing of this peptide revealed a single sequence beginning from oAsp-180, confirming the results obtained with V8 protease. The N-terminal sequence of the unlabeled proteolytic fragment migrating at 14 kDa was identified to start from oAsp-200. Thus, the cross-linked 125I-MR44 must be located within a stretch of 20 amino acid residues from oAsp-180 to oLeu-199. Cleavage of the 125I-MR44-labeled a-subunit with ArgC protease generated a peptide pattern with only one peptide labeled of about 4 kDa. N-terminal amino acid sequencing identified two proteolytic fragments starting with Arg-182 and Leu-80, respectively. The peptide beginning with oLeu-80 was shown to be nonradioactive using V8, AspN, and LysC proteases. Consequently, 125I-MR44 must be cross-linked to the ArgC-4 fragment starting with Arg-182, confirming the region of 125I-MR44 photoincorporation suggested from the V8 and AspN digest experiments. The sequence oArg-182 to oLeu-199 labeled by 125I-MR44 contains a lysine residue in position 185. Therefore, LysC protease was used to cleave the labeled a-subunit. In mapping experiments using LysC protease, we further narrowed down the site of modification to the amino acid stretch oHis-186 to oLeu-199.
This sequence is located in the large N-terminal domain of the α-subunit; it is found close to one of the three loops that contribute to the agonist-binding site. Photoaffinity reagents and site-directed mutagenesis have been utilized in previous studies to identify amino acid residues facing the agonist-binding domain. Three discrete regions on the α-subunit primary structure have been identified: αTrp-86 to αTyr-93 (loop A), αTrp-147 to αTyr-150 (loop B), and αTyr-190 to αTyr-198 (loop C; Refs. 50–54). Two additional regions on each of the neighboring δ- and γ-subunits were found to contribute to the binding site (loop D and E; Refs. 55–57). Based on these data, a spatial model has been developed according to which the sequences of the α-subunit and two sequences of the neighboring δ- and γ-subunits form the agonist binding pockets of nAChR (58, 59). The amino acid residues forming loop C overlap in the C-terminal region at least partially with the sequence αHis-186 to αLeu-199 labeled by 125I-MR44. Since the agonist carbachol or the competitive antagonist α-BTX had only minor influence on the photocross-linking yield of 125I-MR44, the site of interaction between the aromatic ring of 125I-MR44 and the nAChR should be found outside the zone that is sterically influenced by any bound agonist contacting αTyr-190, αCys-192, αCys-193, and αTyr-198. Therefore, the site of interaction of the aromatic head group of 125I-MR44 lies presumably within the hydrophobic sequence HWYY (residues α186–189) containing three aromatic amino acid residues. Receptor desensitization induced by carbachol might influence the accessibility of those residues that react in the resting channel state with 125I-MR44 without affecting the binding affinity of the whole molecule.

From various structure-activity relationship studies, it is obvious that the aromatic moiety of PhTX derivatives has a significant influence on the binding affinity of these compounds (26, 27). Their binding properties were considerably improved by increasing the size and hydrophobicity of the head group. The finding that 125I-MR44 photolabeled a sequence of the α-subunit in which aromatic amino acid residues are accumulated is in line with the observation that PhTX derivatives that carry a large, hydrophobic head group bind with increased affinity to the nAChR (26, 27).

Since the photolabile azido residue of MR44 is located at its aromatic head group, the site of 125I-MR44 cross-linking identifies the region to which this hydrophobic part of the molecule binds. In contrast, the site of interaction of the conformationally flexible carbon chain can be less exactly determined. Due to its positively charged -NH₂ groups interspersed with hydrophobic -CH₂ groups, the polyamine chain is expected to preferentially interact with acidic and hydrophobic amino acid side chains, respectively. This is similar to the active site of bacterial polyamine binding proteins (60). In previous studies, it was suggested that the positively charged polyamine tail binds in the lumen of the nAChR channel to the negatively charged amino acids that are part of the selectivity filter (27). To examine the site to which the polyamine moiety of MR44 binds, the well characterized luminal NCI ethidium was used in various displacement assays. Previous studies located the ethidium-binding site at the high affinity NCI site deep in the ion channel. Labeled isomer of ethidium was used as a fluorescent probe at the nAChR, we determined recently the binding affinities of various PhTX derivatives including MR44. Increasing concentrations of MR44 reduced the fluorescence of bound ethidium, indicating that MR44 displaced ethidium from its binding site. This observation corresponds well to the results presented here that ethidium competes with bound 125I-MR44 and that photoincorporation of 125I-MR44 was prevented in the presence of ethidium and TPM⁺. Ethidium was still displaced by MR44 even when allosteric transitions within the nAChR had been abolished by covalent cross-linking (26). This finding strongly suggests that MR44 interacts with ethidium in a direct competitive manner at an overlapping luminal binding site. Channel-permeating cations, such as calcium, are known to bind to sites within the nAChR ion channel that sterically overlap with the high affinity NCI site (42). Using calcium in direct binding experiments, we could show that 125I-MR44 bound to the nAChR was completely displaced by calcium. This observation indicates a strong influence of polar electrostatic interactions between 125I-MR44 and the NCI site of the nAChR. A site of negative charges located deep in the lumen of the nAChR ion channel that might interact with the positively charged polyamine tail of 125I-MR44 could provide the acidic amino acid residues of the selectivity filter as suggested in earlier studies (27). To summarize, the polyamine moiety of MR44 interacts with the high affinity NCI site of the nAChR, while the aromatic ring of this compound binds to the upper part of the ion channel, i.e., in the vestibule, and therefore to a hydrophobic region on the α-subunit that is located in close proximity to the loop C of the agonist binding site and is accessible from the water-filled lumen of the channel.

Acknowledgment—We thank Hermann Bayer for help with nAChR preparations and for excellent technical assistance.

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Polyamine Binding Site at the nAChR


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