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Defining Nicotinic Agonist Binding Surfaces through Photoaffinity Labeling†

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Abstract

Nicotinic acetylcholine (ACh) receptor (nAChR) agonists are potential therapeutic agents for neurological dysfunction. In the present study, the homopentameric mollusk ACh binding protein (AChBP), used as a surrogate for the extracellular ligand-binding domain of the nAChR, was specifically derivatized by the highly potent agonist azidoepibatidine (AzEPI) prepared as a photoaffinity probe and radioligand. One EPI-nitrene photoactivated molecule was incorporated in each subunit interface binding site based on analysis of the intact derivatized protein. Tryptic fragments of the modified AChBP were analyzed by collision-induced dissociation and Edman sequencing of radiolabeled peptides. Each specific EPI-nitrene-modified site involved either Tyr195 of loop C on the principal or (+)-face or Met116 of loop E on the complementary or (−) face. The two derivatization sites were observed in similar frequency, providing evidence of the reactivity of the azido/nitrene probe substituent and close proximity to both residues. $[{}^{3}H]AzEPI$ binds to the α 4 β 2 nAChR at a single high-affinity site and photoaffinity-labels only the α 4 subunit, presumably modifying Tyr225 spatially corresponding to Tyr195 of AChBP. Phe137 of the β 2 nAChR subunit, equivalent to Met116 of AChBP, conceivably lacks sufficient reactivity with the nitrene generated from the probe. The present photoaffinity labeling in a physiologically relevant condition combined with the crystal structure of AChBP allows development of precise

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Protein sequence alignment between *Aplysia* AChBP and chick α4 and β2 nAChR subunits. This material is available free of charge via the Internet at<http://pubs.acs.org>.

structural models for the AzEPI interactions with AChBP and α 4 β 2 nAChR. These findings enabled us to use AChBP as a structural surrogate to define the nAChR agonist site.

> Neurotransmitter receptors and their intrinsic or associated channels or coupling proteins are the targets for many drugs of therapy and abuse in the central nervous system. Nicotinic acetylcholine (ACh) receptors (nAChRs) are prototypical agonist-gated ion channels responsible for rapid excitatory neurotransmission. Vertebrate nAChRs consist of diverse subtypes assembled from different sets of subunits expressed in skeletal muscle [α 1, β], γ (or ε), and δ , neurons (α2–α10 and β 2– β 4), and sensory epithelia (α9 and α10) (1,2). Among them the α 4 β 2 nAChR, consisting of two α 4 and three β 2 subunits, is the most abundant and widespread in brain (3–6). The α 4 β 2 nicotinic receptor is a major target for potential therapeutic agents for neuropathic diseases, cognitive enhancement, analgesia, and smoking cessation (2, 7) and is also important in considering the selective toxicity of insecticides (8).

> Electron microscopy, X-ray crystallography, and fluorescence and NMR spectroscopy approaches give important but limited understanding of the recognition properties of the binding sites and their capacity to elicit a signal. Affinity-labeling and site-directed mutation studies on *Torpedo* or skeletal muscle nAChR have identified several amino acid residues delineating the ligand-binding pocket (loops A–F) that are localized at the interfacial region between subunits (1, 9–12). Specific subunit combinations confer differences in sensitivity to ACh and pharmacological profiles among the nAChR subtypes. The ligand-binding site in all subtypes contains a conserved core of aromatic amino acid residues (1, 13–16). Neighboring variable residues are considered to confer individual pharmacological properties to each nAChR subtype (1, 10).

> The knowledge of agonist-nAChR interactions was greatly advanced by the discovery of epibatidine (EPI) isolated from the skin of an Ecuadoran frog (17) which has exceptionally high affinity to the nAChRs, including α 4 β 2, and profound analgesic activity (18). Another major step forward was the identification and crystallization of soluble ACh binding proteins (AChBPs) from the snail *Lymnaea stagnalis* (19, 20) and apo and EPI-bound forms from a saltwater mollusk, *Aplysia californica* (21, 22), as surrogates for the extracellular ligand-binding domain of the nAChR. EPI and AChBP offer a signature high-affinity complex that prompts this study using the photoaffinity-labeling approach combined with mass spectrometry (MS), which is a physiologically compatible chemical method for threedimensional structural investigations. The first objective of this research is to develop 5 azidoepibatidine (AzEPI) as a radioligand and photoaffinity probe. The second is to identify the AzEPI-labeled site by specific derivatization of AChBP and collision-induced dissociation (CID) analysis. The third employs $[3H]$ AzEPI to photoaffinity label the α 4 β 2 nAChR and predict potential modification sites. Finally, structural models are presented for the agonist binding domains of AChBP and α 4 β 2 nAChR.

EXPERIMENTAL PROCEDURES

Chemicals

Sources were as follows: (±)-EPI HCl from TOCRIS (Ellisville, MO); carbachol (Carb) chloride, (−)-nicotine (NIC) hydrogen tartrate, and α-bungarotoxin (R-BGT) from Sigma (St. Louis, MO); unlabeled (\pm) -5-azidoepibatidine (AzEPI) from our previous study (23) using a different preparation methodology from the radiosynthesis of βH]AzEPI described below. [3H]EPI was purchased from Amersham Biosciences (Piscataway, NJ).

Synthesis of AzEPI Intermediate 3 (Scheme 1)

N-Boc-2-tosyl-7-azahepta-2,5-diene (1) was synthesized by the Diels-Alder reaction of *N*-Boc-pyrrole and tosylacetylene (24). Into a solution of **1** (700 mg, 2.0 mmol) in THF and methanol (1:1, 40 mL) was added Na₂HPO₄ (4.25 g, 30 mmol). The solution was cooled to -15 °C, and 5% sodium amalgam (2 × 2.0 g) was added with further stirring for 1 h. The reaction was quenched with saturated $NH₄Cl$ solution and then extracted with EtOAc. Removal of the solvent provided **2** (125 mg, 30%). Into a vial with **2** (125 mg, 0.65 mmol) were added sequentially 3-amino-2-chloro-5-iodopyridine (23) (149 mg, 0.58 mmol), Pd(OAc)₂ (14 mg, 0.062 mmol), PPh₃ (34 mg, 0.13 mmol), DMF (1.0 mL), piperidine (166 mg, 193 *μ*L, 1.95 mmol), and formic acid (75 mg, 61 *μ*L, 1.63 mmol). The solution was then stirred at room temperature for 4 days. Workup as usual (23) followed by column chromatography on silica gel with ether and hexane (2:3) provided $3(30 \text{ mg}, 19\%)$. ¹H NMR: δ7.70 (s, 1H), 7.13 (s, 1H), 6.45 (br s, 2H), 4.84 (br s, 1H), 4.54 (br s, 1H), 4.05 (s, 2H), 2.59 (dd, *J* = 4.1, 8.2 Hz), 1.86–1.74 (m, 2H), 1.41 (s, 9H).

Conversion of 3 to AzEPI and [3H]AzEPI (Scheme 1)

Into a solution of **3** (12 mg, 0.037 mmol) in EtOAc (1.5 mL) was added 10% Pd/C (2.5 mg). The atmosphere was replaced with hydrogen, and the solution was stirred at room temperature for 4 h. Hydrogen was then evacuated, and the solution was passed through a syringe membrane filter into a vial. The solvent was removed by flushing with nitrogen, providing a residue of **4** which was treated with 37% HCl (0.16 mL, 1.92 mmol) and stirred at room temperature for 30 min followed by addition of $H₂O$ (0.4 mL). The mixture was then cooled to 0 °C, and NaNO₂ (125 μ L of 100 mg/mL solution precooled to 0 °C, 0.18 mmol) was added with continuous stirring for 15 min at 0 $^{\circ}$ C and an additional 15 min at room temperature. Solid NH₂OH·HCl salt (15 mg, 0.22 mmol) was added, and the mixture was cooled to 0° C again. NaOH (2 N, 1.0 mL, 2.0 mmol) was added dropwise, and the solution was stirred at 0° C for 4 h. HPLC indicated the desired product AzEPI (23). Tritiation was performed by ViTrax (Placentia, CA). Following the above hydrogenation procedure, replacing hydrogen with tritium gas, and properly modifying the reactor to handle the radioactive atmosphere provided $[{}^{3}H]AzEPI$, which was purified by HPLC with an ODS1 150×4.6 mm C₁₈ reverse-phase column (mobile phases, 0.01% trifluoroacetic acid in water and acetonitrile; gradient, 10–90% acetonitrile over 10 min at 5 mL/min). The specific activity and radiochemical purity were determined by HPLC cochromatography and radioflow chromatography.

Preparation of AChBP

Aplysia AChBP (flanked with an N-terminal FLAG epitope) was expressed as a soluble exported protein from stably transfected HEK293S cells lacking the *N*acetylglucosaminyltransferase I gene and selected for geneticin resistance (21, 22). Culture medium containing the secreted AChBP was harvested from multi-layered flasks to obtain several liters of media over a 1–2 week period. Media were stored at 4 °C with 0.02% sodium azide. AChBP was purified on immobilized anti-FLAG antibody and dialyzed against 50 mM Tris-HCl buffer (pH 7.4) containing 150 mM NaCl and 0.02% sodium azide. The dialysate was concentrated by ultrafiltration using the YM-50 Centricon unit (Millipore, Bedford, MA), and the Tris buffer was finally exchanged by phosphate-buffered saline (PBS) [50 mM sodium phosphate buffer (pH 7.5) containing 50 mM NaCl] using the Centricon to give 2–3 mg of protein/mL.

Preparation of the α**4**β**2 Receptor**

Chick α4β2 nAChR was prepared from the M10 cell line of mouse fibroblasts stably transfected with chick α 4 and β 2 subunits under the control of a dexamethasone-sensitive promoter (25). The 40000*g* pellet (30 min, 4 °C) from a homogenate of the harvested cells was resuspended in PBS and immediately used for radioligand binding. Alternatively, the pellet for photoaffinity labeling was solubilized with 2% Triton X-100 in PBS containing 2 mM phenylmethanesulfonyl fluoride (solubilization medium) for 20 min at 4 °C on a rocking platform. Unsolubilized material was removed by centrifugation as above.

Radioligand Binding

Different types of procedures were used for assays with AChBP and α 4 β 2 nAChR. AChBP determination of AzEPI potency involved an adaptation of a scintillation proximity assay with $\left[\right]^{3}$ H]EPI (26). The α 4 β 2 receptor preparation (200–300 μ g of membrane protein) in PBS (500 μ L final volume) was incubated for 60 min at 25 °C with appropriate concentrations of $\binom{3}{1}$ AzEPI alone or with an unlabeled displacer. The assay was terminated by rapid filtration and three rinses with ice-cold saline and then scintillation counting. Nonspecific binding was determined with 2 *μ*M EPI. Binding parameters [dissociation constant (K_D) , maximum binding capacity (B_{max}) , and Hill coefficient (n_H)] were obtained from saturation experiments. IC_{50} values, molar concentrations for 50% displacement, were converted to inhibition constant (K_i) values using the equation of Cheng and Prusoff (27).

Photoaffinity Labeling

AChBP has five interfacial binding sites in the homopentamer. A solution of AChBP (38 or 76 pmol sites) in PBS [50 or 100 *μ*L final volume, respectively (760 nM sites final concentration)] was incubated with 1.2 equiv of $\lceil 3H\rfloor$ AzEPI for 60 min at 25 °C in the dark without or with 10000 pmol (10 *μ*M final concentration) of EPI (for nonspecific labeling). After irradiation with 300 nm lamps for 20 s (AzEPI completely photoreacted in this condition), the radiolabeled AChBP recovered by ultrafiltration with the YM-30 Centricon unit was subjected to SDS–PAGE with detection of proteins by Coomassie dye and labeled bands with the Amplify fluorographic reagent (Amersham Biosciences) followed by gel slicing and scintillation counting. For MS analysis AChBP (200 pmol sites, 2 *μ*M sites final

concentration) in PBS (100 μ L) was treated with 1.5 equiv of nonradioactive AzEPI as above. Derivatized AChBP was purified by the YM-30 Centricon unit and PBS exchanged to 25 mM ammonium bicarbonate buffer (pH 7.8).

For α 4 β 2 nAChR, the solubilized receptor preparation (10 pmol of [³H]AzEPI binding sites, about 10 mg of total protein) in solublization medium (1 mL) was incubated at 25 °C for 60 min in the dark with 2 equiv of $[3H]AzEPI$ (20 nM final concentration) without or with 10000 pmol (10 *μ*M final concentration) of EPI (for nonspecific labeling). The reaction mixture was irradiated as above, incubated overnight at 4 °C with monoclonal antibody mAb270 against the β 2 subunit (Sigma) (dilution ratio 1:100), and finally reacted with protein G PLUS-agarose (Santa Cruz Biotechnology Inc., Santa Cruz, CA) for 120 min at 4 °C. After recovery of the agarose beads and five washes with solubilization medium, the proteins were subjected to SDS–PAGE followed by fluorographic detection and radioactivity quantification.

Edman Sequencing of Radiolabeled Peptides

[³H]AzEPI-labeled AChBP was denatured with urea, reduced with dithiothreitol, and alkylated with iodoacetamide. The sample was desalted by the YM-3 Centricon column and washed twice with 25 mM ammonium bicarbonate buffer (pH 7.8). The radiolabeled protein was deglycosylated with peptide *N*-glycosidase F (Sigma) and then digested with trypsin from Promega (Madison, WI). The digested fragments were separated by HPLC with a Vydac 150×1 mm C₁₈ reverse-phase column (Sorbent Technologies, Atlanta, GA). The radioactive peptide fractions defined by liquid scintillation counting were analyzed with the ABI 494-HT Procise Edman sequencer (Applied Biosystems, Foster City, CA).

MS Analysis

Molecular weights of the intact AChBP subunit were measured on an Applied Biosystems MDS Sciex QSTAR hybrid quadrupole time-of-flight mass spectrometer. Samples in 10 mM ammonium bicarbonate were injected onto a Phenomenex 150×0.1 mm Onyx monolithic column (Torrance, CA) flowing at 500 nL/min using 50% acetonitrile in water containing 0.1% formic acid as the mobile phase. Data were acquired and the charge state distributions converted to a normalized zero charge spectrum using the Analyst 1.1 software. Protein standards of about the same size showed a mass accuracy of approximately 1–2 Da.

For CID analysis, intact proteins in buffer with an equal volume of acetonitrile were reduced with dithiothreitol, alkylated with iodoacetamide, and digested with trypsin (Promega) for 4 h at pH 7.8 and 37 °C. After removal of the acetonitrile, digested samples were analyzed by nanoflow LC/MS/MS on the QSTAR instrument as above with a self-packed Ultro 120 (5 μ m particle size) (Peeke Scientific, Redwood City, CA) 150 \times 0.1 mm capillary column. The flow rate was 300–330 nL/min with a gradient from 5% to 50% acetonitrile in 60 min (all solvents contained 0.1% formic acid). The Information-Dependent Acquisition mode was used with a 1.0 s survey scan from *m*/*z* 310 to *m*/*z* 1400, followed by one 3 s CID scan on the most abundant multiply charged ion in the survey scan. Data were analyzed with Protein Prospector v4.23.4, an in-house developed program, to identify peptides that shifted in mass reflecting the photolabel incorporation.

Accurate masses of the labeled peptides were obtained from LC/MS/MS analysis on a Thermo Electron LTQ FT hybrid linear ion trap FTICR mass spectrometer (Waltham, MA).

Chromatography was as described above for LC/MS/MS on the QSTAR. The datadependent acquisition method consisted of one full mass survey scan in the FTICR at 25000 resolution. The three most intense multiply charged target ions were isolated for accurate mass measurement in the FTICR in SIM mode, using a 10 Da mass window at 50000 resolution. The accuracy of the mass measurements is typically within 2 ppm under such conditions. These three ions were then fragmented in the linear ion trap using CID. Target values for the three different scan modes were 3000000, 50000, and 30000, respectively.

Modeling and Calculations

The X-ray structure for *Aplysia* AChBP (22) as the EPI-bound (PDB code 2BYQ) form was used as template for building the α 4 β 2 nAChR homology model. The fasta protein sequences of the α4 and β2 subunits for *Gallus gallus* (5, 28, 29) were downloaded from UniProt (30) (accession numbers P09482 for α 4 and P09484 for β 2) and then aligned with 2BYQ using the CLUSTAL W (31) web server at the European Bioinformatics Institute (32, 33). Chain A of the PDB structure was chosen for sequence alignment with the ligandbinding domain (residues α 4 1–240 versus 2BYQ:A, β 2 1–240 versus 2BYQ:A). These alignments were input to the SWISS-MODEL protein homology server (34). The resulting model subunit structures were imported into Maestro 6.5 (Schrodinger, L.L.C., Portland, OR). The α 4 and β 2 model structures were aligned via comparison of backbone atoms onto the corresponding subunits in 2BYQ, yielding a complete pentameric α 4 β 2 model structure. Further optimization was limited to a single $\alpha/2/2$ subunit pair and, hence, a single binding interface. This α 4 β 2 model was subjected to cycles of energy minimization using the OPLS_2005 force field implemented in Macromodel (35, 36). Up to 5000 steps per minimization were run to achieve a gradient of 0.5 with respect to energy. In the initial minimization cycles, the backbone was held constant. In subsequent minimization cycles, the region within 15 \AA of the binding pocket was free to move with progressive constraints on the remainder of the structure. Docking calculations were carried out using the GALS algorithm in AutoDock 3 (37, 38). Docked molecules were constrained to a 40 point grid in the region of the binding pocket.

RESULTS

Radiosynthesis

The reaction sequence to prepare [3H]-AzEPI via **1**–**4** was designed to introduce the tritium label at the penultimate step and finally convert aminopyridine $[{}^{3}H]$ -4 to azidopyridine [³H]AzEPI. Detosylation of **1** with sodium–mercury amalgam (a) provided diene **2**, which was treated with 3-amino-2-chloro-5-iodopyridine under reductive Heck coupling conditions (b) to provide the desired *exo* isomer **3**. Compound **3** was hydrogenated or tritiated with tritium gas using Pd/C as the catalyst (c). The amino group was converted to azido by sequential treatment with concentrated HCl and NH₂OH (d), which also removed the Boc group to provide $[{}^{3}H]$ AzEPI after HPLC with a specific activity of 36 Ci/mmol and radiochemical purity of >99% by radioflow chromatography. Little or no radiochemical decomposition was observed for $\binom{3}{1}$ AzEPI stored in ethanol at −20 °C for at least 3 years.

Ligand-Binding Measurements (Figure 1)

For *Aplysia* AChBP, affinities for EPI and AzEPI were 8.6 and 10 nM, respectively. For α 4 β 2 nAChR expressed in M10 cells, specific [³H]AzEPI binding was saturable and exhibited a single high-affinity interaction with K_D 1.0 nM, B_{max} 1.0 pmol/mg of protein, and n_H 0.99. In competition experiments, EPI, NIC, and Carb displaced $[^3H]$ AzEPI binding with K_i values of 0.2, 14, and 820 nM, respectively, while α -BGT failed to show any inhibition even at 10000 nM.

AChBP Binding Site. Photoaffinity Labeling at Two Distinct Sites in Similar Frequency

[³H]AzEPI photoreacted with 19.5 \pm 1.2 pmol sites with nonspecific labeling of 0.43 \pm 0.002 pmol (specific labeling 98%), occupying 24% of the total AChBP (76 pmol sites) (Figure 2). EPI protected against specific incorporation of $\binom{3}{1}$ AzEPI into AChBP by 50% at approximately 500 pmol. Tryptic fragments of $[3H]$ -AzEPI-derivatized AChBP were subjected to HPLC separation, revealing only two labeled peptides with nearly equivalent radioactivity (Figure 3). Edman sequencing then established that the two radioactive peptides were Ser189−Lys203 (a chymotryptic fragment from trace chymotrypsin impurity) and Thr80−Arg122 (a tryptic fragment) (data not shown), establishing that there were two sites of labeling in similar frequency.

Labeling at the Subunit Level

Mass measurement of AChBP treated with AzEPI yielded 25352 Da for the unmodified protein and 25574 Da for the derivative (Figure 4). The calculated MW for the FLAGtagged [nine amino acids numbered as (−8)DYKDDDDKL(0)] protein is 25944 Da. MS-Nonspecific Search in the ProteinProspector software package revealed that 10 different sequences may correspond to this mass within 0.35%, where the N-terminus could be any amino acid from 1 to 6 [Asp(−8) to Asp(−3)] and the C-termini vary from 223 to full length, i.e., 228. At the same time LC/MS/MS analysis of the tryptic digest verified the presence of the N-terminal peptide (data not shown), and Arg215 was the last residue detected for the Cterminus. Considering the mass accuracy achieved for standard proteins, it was concluded that most likely a mixture with ragged termini was present. Tyr (-7) to Arg215 features the closest mass to the measured value: 25324 Da (−0.1% deviation). There were other components observed at 25368 and 25385 Da, i.e., with 16 and 33 Da higher mass, respectively. Since the sequence contains five Met, four Trp, and four Cys residues, it is reasonable to assume that these masses indicate partial oxidation most likely to methionine mono- and bissulfoxides. The second molecular mass cluster (with 25574 and 25590 Da) clearly indicates the incorporation of a single EPI-nitrene. The added mass relative to the unmodified protein was 222 Da, while the expected mass increment was 221 Da; this difference was verified at the peptide level (see below) where it could be more accurately measured. Very weak ions yielding an intact MW of 25795 Da indicated the possibility of trace double labeling.

Labeled Positions Are Tyr195 and Met116

Once the presence of the label was ascertained at the protein level, the mixture was reduced, alkylated, and digested with trypsin. The resulting digest was subjected to LC/MS/MS

analysis in a data-dependent fashion, where the computer selected the multiply charged ions for CID analysis. All MS/MS data were subjected to a two-step database search using the inhouse ProteinProspector v4.23.4. First, with very strict search parameters the identity of the protein was verified, as well as the lack of contaminating proteins established. Then a second search was conducted that permitted a +300 Da modification on any amino acid in the sequence. The +221 Da shift identified three EPI-nitrene-modified peptides: (1) Gln184−Lys203 and two overlapping peptides, (2) Pro98−Arg122, and (3) Thr80−Arg122. Accurate mass measurements were obtained for the peptide precursor ions (*m*/*z* 668.1 and 970.3) for the two peptides, Gln184−Lys203 and Pro98−Arg122 (Table 1). The CID data identified the modified residues as Tyr195 (Figure 5) and Met116 (Figure 6).

α**4**β**2 nAChR Binding Site. Photoaffinity Labeling Exclusively at the** α**4 Subunit**

The α 4 β 2 receptor was selectively labeled by [³H]AzEPI at the α 4 subunit (as a doublet), but no labeling was evident on the β 2 subunit (Figure 2). The labeling amount on the α 4 subunit was 0.23 ± 0.01 and 0.008 ± 0.0002 pmol in the absence and the presence of unlabeled EPI, respectively, establishing 97% specific incorporation. The occupancy rate was 2.2% of the total sites (10 pmol). The existence of the α 4 subunit in the corresponding region on the SDS−PAGE gel was confirmed by LC/MS/MS analysis of an unlabeled sample after immunoprecipitation (data not shown).

Binding Site Models

The structure of AzEPI was first optimized in Gaussian 03 at B3LYP/6-31G. The N=N=N angles for phenyl azides in the Cambridge Structural Database average around 170°, presumably due to resonance forms wherein the central nitrogen is not entirely *sp.* The electronic and steric environment also affects the angle, and AzEPI has a calculated N=N=N angle of 165°. The azido moiety can rotate about the C–N bond, resulting in two optimized conformers with similar energy levels, suggesting that both exist at room temperature. This bond was allowed to rotate in the docking calculations.

AzEPI is calculated to dock into the AChBP (2BYQ structure without EPI itself) and the present α4β2 nAChR model with energies of −9.69 and −7.87 kcal/mol, respectively (Figure 7). The AzEPI-bound AChBP structure is superimposable to the EPI-bound crystal (2BYQ). Two critical interactions are predicted for the AzEPI ammonium with Trp147 of AChBP and Trp177 of α4, i.e., hydrogen bonding with the backbone carbonyl oxygen (1.8 and 2.0 Å, respectively) and cation– π contact to the indole side chain (3.2 and 3.6 Å, respectively). The 6-chloro substituent of AzEPI makes van der Waals contacts to the backbone oxygen atoms of Ile106 (3.6 Å) and Met116 (3.6 Å) in AChBP and to Asn127 (3.7 Å) and Phe137 (3.3 Å) in α 4 β 2. The AzEPI pyridine nitrogen atom probably undergoes hydrogen bonding with Ile118 (4.1 Å) and Trp147 (3.7 Å) via a solvent bridge in AChBP, and the equivalent targets would be Leu139 (4.6 Å) and Trp177 (3.5 Å) in α 4 β 2. The AzEPI azido group is positioned between Tyr195/Met116 of AChBP and Tyr225 of α4 and Phe137 of β2, allowing the derivatization by the EPI-nitrene (e.g., the distance between the EPI-nitrene nitrogen to the hydroxyl oxygen of Tyr195 or sulfur of Met116 is presumably about $3-4$ Å), although the β 2 subunit was not modified. The azido conformer pointing toward the chlorine atom is closer to Met116, and the one pointing away is closer to Tyr195 (Figure 7). Further, Tyr195

and Met116 in the AzEPI-bound holo structure are 3–4 Å more closely positioned relative to the agonist molecule than those in the apo form (Figure 8).

DISCUSSION

[³H]AzEPI, an Optimized Photoaffinity Radioligand

The photoreactive 5-azido substituent was introduced at a structural position selected to minimize any change in potency $(23, 40, 41)$. AzEPI and $\binom{3}{1}$ AzEPI show similar affinity to EPI itself in the $1-10$ nM range, and fortunately $[{}^{3}H]$ AzEPI has excellent radiochemical stability.

AChBP Agonist Binding Site

Structural information regarding the binding domains and subunit interfaces was expanded by the discovery and crystallization of AChBPs liganded with several agonists and antagonists (19, 20, 22, 42–44). The refined 4 Å resolution electron microscopy structure of the heteropentameric *Torpedo* nAChR established considerable structural similarity of AChBP with this nAChR (45). *Aplysia* AChBP is shown here to bind specifically and photoreact with AzEPI, giving the incorporation ratio of one probe molecule (EPI-nitrene) per AChBP subunit (binding site), and both Tyr195 on loop C and Met116 on loop E (in similar frequency) are revealed as the sites for derivatization. The nearly equivalent labeling on the primary loop C face and the complementary loop E on the neighboring subunit shows the close proximity of these residues on the two subunits. These results precisely define the orientation of EPI in the binding pocket under a physiologically compatible condition.

Comparing *Aplysia* AChBP–ligand complexes with an apo structure provides direct evidence of differential conformational rearrangements (particularly of loop C) induced by antagonist and agonist occupation (22). The agonist-elicited conformational changes are also supported by fluorescence spectroscopy, deuterium-hydrogen exchange MS, and NMR spectroscopy analyses of AChBP (46–48). The present photoaffinity labeling clearly indicates that loop C Tyr195 is oriented toward loop E Met116, and upon agonist binding loop C draws inward to cap the binding pocket and envelop the agonist molecule (Figure 8). A similar proposal has been made from photolabeling studies for tightening the interfacial binding domain (involving primary and complementary subunits) after ligand interaction in *Torpedo* nAChR (12, 49, 50). This conformational change serves as an initial event for the ligand-induced channel opening mechanism of the nAChR, but it remains unclear how agonist occupation is coupled to the distant transmembrane-localized ion gating site.

α**4**β**2 nAChR Binding Sites**

A major quest in developing nicotinic agents is the discovery of highly subtype-selective agonists and antagonists particularly for the α 4 β 2 nAChR. The present study for the first time succeeds in photoaffinity labeling the specific binding site of the α 4 β 2 receptor, and [³H]AzEPI exclusively labels the α 4 subunit. Tyr225 of the α 4 subunit is equivalent to Tyr195 of AChBP loop C. Phe137 of the β2 subunit corresponds to Met116 of AChBP loop E. Therefore, even though they are similarly positioned, the difference between Met116 of AChBP and Phe137 of the β 2 subunit appears to determine whether loop E is labeled or not,

presumably depending on reactivity of the side chain. For instance, the Met residue is a preferred site for photoderivatization by the benzophenone radical (51, 52). The first $\alpha 4\beta 2$ nAChR binding site model was constructed on the basis of the crystal structure of HEPESbound *Lymnaea* AChBP (53, 54). An improved α4β2 model docked with dechloro-EPI was generated using the NIC-bound *Lymnaea* AChBP template (55). The present model combines our photoaffinity labeling result and the X-ray structure of EPI-bound *Aplysia* AChBP. The cationic moiety of AzEPI interacts at a subsite surrounded by aromatic amino acid residues including the critical Trp177 on the α 4 β 2 loop B segment. The backbone carbonyl oxygen and aromatic residue of Trp make hydrogen-bonding and cation– π contacts, respectively, on analogy with the findings on the R1-containing nAChR subtype and the *Lymnaea* and *Aplysia* AChBPs (14, 16, 20, 22). Substitution of Tyr55 (loop D) in *Aplysia* AChBP by Trp, as found in the β 2 subunit (Figure 7), improves the affinity of EPI, suggesting that the nature or size of the aromatic side chain at this position is an important factor for EPI binding (22).

Concluding Remarks

Photoaffinity labeling in a physiologically relevant condition combined with MS analysis and the crystal structure of AChBP offers a precise molecular picture of the drug–receptor interaction and allows development of a structural model for the agonist binding site on the α 4 β 2 subunit interface. These findings further define the surfaces with which agonists associate and should help to facilitate rational design of therapeutic agents for neurological disorders and analgesia.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

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FIGURE 1.

Ligand binding with AChBP (a) and α 4 β 2 nAChR (b and c). (a) Concentration-dependent response curves for direct titration of $[^3H]EPI$ and for AzEPI competing with $[^3H]EPI$ binding to AChBP. K_D for [³H]EPI is 8.6 ± 2.0 nM (n_H 0.90 \pm 0.05), and K_i for AzEPI is 10 \pm 1 nM (n_H 1.13 \pm 0.08) (\pm SD, $n = 3-4$). (b) Saturation isotherm and Scatchard plot (insert) for specific [³H]AzEPI binding to α 4 β 2 nAChR. The binding parameters (\pm SD, *n* = 3) are K_{D} 1.0 \pm 0.1 nM, B_{max} 1.0 \pm 0.02 pmol/mg of membrane protein, and n_{H} 0.99 \pm 0.03. (c) Displacement curves of specific $[3H]$ AzEPI binding to α 4 β 2 nAChR by three nicotinic agonists and the antagonist α -BGT. K_i values (and n_H) are as follows (\pm SD, $n = 3$): 0.17 \pm 0.02 nM (1.02 \pm 0.02) for EPI, 14 \pm 2 nM (0.96 \pm 0.01) for NIC, and 820 \pm 100 nM (0.98 \pm 0.01) for Carb.

FIGURE 2.

[³H]AzEPI photoaffinity labeling of AChBP and α 4 β 2 nAChR. (a) Fluorography of [³H]AzEPI-labeled AChBP compared with Coomassie blue-stained protein and quantification by scintillation counting (bar graph with SD of three experiments). AChBP (76 pmol sites) was photoaffinity labeled by $[3H]AzEPI$ (92 pmol) alone (–) and protected with EPI (10000 pmol) (+). (b) Quantification for protection of photoaffinity labeling of AChBP by EPI. AChBP (38 pmol sites) was reacted with $[3H]$ AzEPI (46 pmol). The titration curve is plotted by radioactivity determination for the protein band as above in separate experiments from the fluorography image. (c) The α 4 β 2 receptor (10 pmol sites) was subjected to photoaffinity labeling by $[3H]AzEPI$ (20 pmol) in the absence (−) and the presence (+) of EPI (10000 pmol). Fluorography shows exclusive labeling of the α 4 subunit. Photoincorporation is defined by radioactivity in the corresponding region on the SDS– PAGE, and the labeled amount is the summation of the doublet which is due to different states of glycosylation (25).

FIGURE 3.

Liquid chromatogram of tryptic fragments of deglycosylated AChBP derivatized by $[3H]$ AzEPI. The entire chromatogram (except for an area around the column void volume) contained two significant radioactive peaks in the positions illustrated. Peak areas are based on UV (214 nm) and radioactivity measurements (dpm) on equal sized aliquots. All other regions of the chromatogram have the 100–600 dpm basal level of radioactivity, i.e., the same in the absence (specific labeling) and the presence of 10000 pmol of EPI (nonspecific labeling).

FIGURE 4.

Molecular mass measurement of an AChBP subunit after labeling with AzEPI. The 25352 cluster represents the unmodified protein (AChBP) within which the mass increases of 16 and 33 Da may indicate the presence of methionine mono- and bissulfoxides. The second cluster corresponds to the singly EPI-nitrene-modified protein (EPI-N-AChBP) and a monooxidized derivative. The third very weak cluster indicates trace double labeling.

FIGURE 5.

Low-energy CID spectrum of the EPI-nitrene-modified Gln184−Lys203 tryptic peptide, precursor ion at *m*/*z* 668.1 (4+). Assignments were made according to the nomenclature of Biemann (39). The modification occurred at Tyr195 as indicated by the unmodified Nterminal b ions (up to b_6 in black) and C-terminal y fragments (up to y_8 in blue) and the modified immonium y ion and internal fragments (in red). For simplicity labeling involves mostly sequence ions and internal fragments containing the modified residue. Carbamidomethylcysteines are designated as C*.

FIGURE 6.

Low-energy CID spectrum of the EPI-nitrene-modified Pro98−Arg122 tryptic peptide, precursor ion at *m*/*z* 728.4 (4+). Met116 is modified. The C-terminal y ions were detected unmodified up to y_6 (fragments in blue), while all y fragments from y_7 (in red) show the 221 Da mass shift. Labeling involves the sequence ions and only a few other fragments.

FIGURE 7.

AzEPI-bound structures of *Aplysia* AChBP and chick α4β2 nAChR. Relevant residues in yellow are from the (+)-face of AChBP or the α4 subunit while relevant residues in aquamarine are from the (-)-face of AChBP or the β 2 subunit. The AzEPI-docked AChBP model was calculated on the basis of the corresponding crystal structure (2BYQ) (22). The α 4 β 2 interface was constructed using 2BYQ as the scaffold.

FIGURE 8.

Comparisons of geometries for Tyr195 and Met116 between the AzEPI−AChBP complex and the apo form. Tyr195 and Met116 (in aquamarine) in the AzEPI-bound conformation (holo) are overlaid with those in the apo structure (in yellow) (2BYN) (22).

Scheme 1. Radiosynthesis of [3H]AzEPI*^a*

^aConditions: (a) 5% Na/Hg, MeOH; (b) 3-amino-2-chloro-5-iodopyridine, Pd(OAc)₂, PPh₃, DMF, piperidine, HCOOH, room temperature, 4 days; (c) T_2 or H_2 , Pd/C (10%), EtOAc, room temperature, 4 h; (d) (i) 37% HCl; (ii) NaNO₂, 0 °C to room temperature, 30 min; (iii) NH₂OH·HCl, 0 °C; (iv) 2 N NaOH, 0 °C, 4 h. Asterisks indicate positions of tritium.

Accurate Masses of the Labeled Peptides Accurate Masses of the Labeled Peptides

 $\frac{a_{m/z}}{2284}$ (+4 ion) for the peptide Pro98–Arg122 (see legend for Figure 6) was not observed in the LTQFT LCMSMS run. However, m/z 970.2933 (+5 ion) for the peptide Thr80-Arg122 was seen. The CID obtained on the LTQ *am/z* 728.4 (+4 ion) for the peptide Pro98−Arg122 (see legend for Figure 6) was not observed in the LTQFT LCMSMS run. However, *m/z* 970.2933 (+5 ion) for the peptide Thr80-Arg122 was seen. The CID obtained on the LTQFT for 970.2933 confirmed its identity.