Different Interactions between MT7 Toxin and the Human Muscarinic M₁ Receptor in Its Free and N-Methylscopolamine-Occupied States

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ABSTRACT

Muscarinic MT7 toxin is a highly selective and potent antagonist of the M₁ subtype of muscarinic receptor and acts by binding to an allosteric site. To identify the molecular determinants by which MT7 toxin interacts with this receptor in its free and NMS-occupied states, the effect on toxin potency of alanine substitution was evaluated in equilibrium and kinetic binding experiments as well as in functional assays. The determination of the crystallographic structure of an MT7-derivative (MT7-diiodoTyr51) allowed the selection of candidate residues that are accessible and present on both faces of the three toxin loops. The equilibrium binding data are consistent with negative cooperativity between N-methylscopolamine (NMS) and wild-type or modified MT7 and highlight the critical role of the tip of the central loop of the toxin (Arg34, Met35 Tyr36) in its interaction with the unoccupied receptor. Examination of the potency of wild-type and modified toxins to allosterically decrease the dissociation rate of [³H]NMS allowed the identification of the MT7 residues involved in its interaction with the NMS-occupied receptor. In contrast to the results with the unoccupied receptor, the most important residue for this interaction was Tyr36 in loop II, assisted by Trp10 in loop I and Arg52 in loop III. The critical role of the tips of the MT7 loops was also confirmed in functional experiments. The high specificity of the MT7-M₁ receptor interaction exploits several MT7-specific residues and reveals a different mode of interaction of the toxin with the free and NMS-occupied states of the receptor.

Muscarinic neurotoxins, small peptides of 64 to 66 residues derived from the venom of African mambas (Dendroaspis angusticeps and Dendroaspis polylepis), are well known for their ability to interact with different muscarinic receptor subtypes. NMR and X-ray studies of the MT2 toxin have shown that muscarinic toxins have the three-finger fold structure, characteristic of the large superfamily of toxins that act at cholinergic synapses (Ménez and Ducruix, 1993; Ségalas et al., 1995; Servent and Ménez, 2001). Ten different muscarinic toxins have been isolated so far. Despite their high sequence homology, they show divergent interaction profiles with the different muscarinic receptor subtypes (for review, see Bradley, 2000; Karlsson et al., 2000). For instance, the MT7 toxin, purified from the venom of the green mamba, binds to the M₁ receptor subtype with a potency in the low or subnanomolar range, with at least 10,000-fold selectivity relative to the other receptor subtypes (Max et al., 1993; Carsi and Potter, 2000; Olianas et al., 2000; Mourier et al., 2003). Pharmacological and functional studies of the MT7-hM₁ interaction have demonstrated that it 1) inhibits ACh-stimulated [³²P]GTPγS binding to CHO cell membranes expressing M₁ receptors, 2) dissociates very slowly from M₁ receptors, and 3) strongly decreases or increases the rate of dissociation of [³H]N-Methylscopolamine (NMS) and [³H]ACh, respectively (Olianas et al., 2000, 2004; Krajewski et al., 2001; Mourier et al., 2003). In summary, MT7 acts as a highly selective antagonist of M₁ receptors by establishing a strong and stable interaction with an allosteric binding site on the receptor.

There is a limited understanding of the specificity, selectivity, and mechanism of action of the muscarinic toxins at
muscarinic receptor subtypes, and the location of the binding sites used by the toxins has proved to be quite challenging. Because of its high affinity and highest subtype selectivity several studies have investigated the MT7-human M1 receptor (hM1) interaction using wild-type and modified muscarinic toxins. Expression of recombinant MT7 in the baculovirus vector-Sf9 insect cell system (Nasman et al., 2000) and in Pichia pastoris (Krajewski et al., 2001) has resulted in relatively low yields. However, using a one-step solid-phase chemical synthesis, we have been able to achieve good yields of muscarinic toxins (Mourier et al., 2003; Fruchart-Gaillard et al., 2006) and have shown that synthetic MT1 and MT7 toxins possess physicochemical and pharmacological profiles identical to those of purified natural toxins. Furthermore, we have been able to demonstrate that the two toxins seem to recognize the hM1 receptor differently (Mourier et al., 2003).

Like other members of family A of G-protein coupled receptors, the hM1 muscarinic receptor is characterized by its having at least two distinct binding sites. Agonists and competitive antagonists bind to the orthosteric site, located inside the transmembrane domain, whereas allosteric agents induce a significant perturbation of the kinetics of binding of ligands to the primary site by interacting with an allosteric site, located more extracellularly (Stockton et al., 1983, Matsui et al., 1995; Ellis et al., 1993; Christopoulos et al., 1998; Christopoulos and Kenakin, 2002). The effects of allosteric ligands, such as the MT7 toxin, are consistent with a ternary complex model in which the orthosteric and allosteric agents can bind simultaneously to the receptor and modify each other's affinity (Ehler, 1988; Lazarenco and Birdsall, 1995). We have characterized, directly and indirectly, using equilibrium and kinetic experiments, the MT7 affinity for the free and NMS-occupied receptor, using an iodinated MT7 derivative and [3H]NMS, respectively (Fruchart-Gaillard et al., 2006). The similar results obtained using the two approaches confirm the reciprocity of the NMS-MT7 cooperative interaction and validate the allosteric ternary complex model used to study the MT7-hM1 interaction.

In this article, we have attempted to delineate the interaction sites by which the MT7 toxin recognizes the hM1 receptor and have tried to explain the molecular origin of the high affinity and selectivity of this interaction. We first solved the crystallographic structure of a diiodo-MT7 analog, which revealed particular structural properties that were considered to possibly be relevant to the unique pharmacology of MT7. This structure highlighted candidate toxin residues to be modified. Thus, we synthesized 12 MT7 variants, each with a single amino acid modification on one of the three toxin loops and analyzed the effect of these modifications on the MT7 potency for the free or NMS-occupied hM1 receptor using equilibrium and kinetic binding experiments as well as functional assays.

Materials and Methods

Materials. [3H]NMS (78 Ci/mmol) was from PerkinElmer Life Sciences (Courtaboeuf, France). Atropine was from Sigma-Aldrich (St Quentin-Fallavier, France). Automated chain assembly was performed on a standard Applied Biosystems 433 peptide synthesizer (Applied Biosystems, Courtaboeuf, France). FLEXstation calcium assay kit (Molecular Devices Ltd., Wokingham, UK) was the calcium-specific fluorescent dye used in this study and the calcium flux measurement was done on a FLEXstation machine (Molecular Devices Ltd.).

Peptide Synthesis, Disulfide Bond Formation, and Protein Purification. Chemical synthesis, refolding, and purification of the modified toxins were carried out using the methods described previously for the synthesis of the muscarinic MT1 and MT7 toxins (Mourier et al., 2003). In brief, the toxins were synthesized using the 9-fluorenylethoxycarbonyl-stepwise solid-phase method with dicyclohexycarbodiimide/1-hydroxy-7-azabenzotriazole as coupling reagents and N-methyl pyrroldione as the solvent. After TFA deprotection and purification of the crude material, the reduced form of the pure modified synthetic toxins was subjected to an oxidative reaction using a mixture of reduced and oxidized glutathione in the presence of guanidine hydrochloride (0.5 M). Finally, the toxins were purified by reversed-phase HPLC using a Yvdac C18 column (250 × 10 mm) with a gradient of 40 to 60% of solvent B in 40 min (A, 0.1% TFA in H2O; B, 60% acetonitrile and 0.1% TFA in H2O) and analytically characterized. Mass determinations were performed on a micromass platform II (Micromass, Altrincham, UK).

Circular Dichroism Analysis. CD spectra were recorded on a spectropolarimeter (CD6; HORIBA Jobin Yvon Inc., Edison, NJ). Measurements were routinely performed at 20°C in 0.1-cm path length quartz cells (Hellma) with a peptide concentration of 5 × 10⁻⁶ M in 5 mM sodium phosphate buffer, pH 7.4. Spectra were recorded over the 186- to 260-nm wavelength range. Each spectrum represents the average of four spectra.

Crystalization and X-Ray Structure of MT7 Derivatives. Crystals of diiodo-MT7 (labeled at Tyr51) were grown by vapor diffusion from diphosphilic toxi, which was resuspended at 5 mg/ml in 50 mM sodium acetate, pH 4.6. The reservoir solution used was 1.25 M ammonium sulfate, 90 mM sodium citrate, and 16% methyl pentanediol, pH 5.5, with drops consisting of 1 μl of toxin and 1 μl of reservoir solution. After 6 months' equilibration in a cooled incubator at 20°C, the crystals were harvested. Data to 1.38 Å resolution were collected at the ESRF beamline ID14-2 in Grenoble (France) from crystals cryo-cooled in 80% saturated Li2SO4 as a cryoprotectant. The HKL suite (Otwinowski and Minor, 1997) of programs was used for data reduction, and the structure was solved by molecular replacement using MOLREP ( CCP4; Vagin and Teplyakov, 1997) using the coordinates of the muscarinic toxin MT2 [Protein Data Bank (PDB) ID 1FF4; R. Ménez and E. Stura, manuscript in preparation]. The XtalView suite (McRee, 1999) was used for electron density map interpretation and other graphical analysis. The structure was refined with REFMAC (Murshudov et al., 1997) to 1.42 Å and gave Rwork and Rfree of 20.3% and 22.1%, respectively. The PDB ID code of the diido-MT7 derivative is 2vlw.

Cells and Membrane Preparation. Pros. P. O. Couraud and A. D. Strosberg (ICGM, Paris, France) kindly provided CHO cells stably expressing the cloned human muscarinic M1, M2, and M3 receptors and the M2 clone was from the Guthrie cDNA resource center (http://www.cdna.org). The cells were grown in plastic Petri dishes (Falcon; BD Discovery Labware, Bedford, MA) that were incubated at 37°C in an atmosphere of 5% CO2 and 95% humidified air in Ham's F12 medium supplemented with 10% fetal calf serum and 1% penicillin/streptomycin (Sigma). At 100% confluence, the medium was removed, and the cells were harvested using a Versene buffer (PBS + 5 mM EDTA). The cells were washed with ice-cold phosphate buffer and centrifuged at 1700g for 10 min (4°C). The pellet was suspended in ice-cold buffer (1 mM EDTA, 25 mM sodium phosphate, and 5 mM MgCl2, pH 7.4) and homogenized using an Elvehjem-Potter homogenizer (Thermo Fisher Scientific Labosi, Elancourt, France). The homogenate was centrifuged at 1700g for 15 min (4°C). The sediment was resuspended in buffer, homogenized, and centrifuged at 1700g for 15 min (4°C). The supernatant was centrifuged at 35,000g for 30 min (4°C), and the pellet was resuspended in the same buffer (0.1 ml/dish). The membrane preparations were aliquotted
and stored at −80°C. Protein concentrations were determined by the Lowry method using bovine serum albumin as standard.

[^3H]NMS Binding Assays. All binding experiments were carried out at room temperature in 10 mM sodium phosphate, pH 7.2, 135 mM NaCl, 2.5 mM KCl, pH 7.4, 0.1% bovine serum albumin (PBS- bovine serum albumin). The effect of various toxins on the equilibrium binding of a fixed concentration of [^3H]NMS was determined in heterologous inhibition experiments. With [^3H]NMS as tracer at a protein concentration at which no more than 10% of added radioligand was bound (approximately 1500 cpm), membranes were incubated in PBS-bovine serum albumin at 25°C for 18 to 22 h, with [^3H]NMS (0.5 nM) and varying concentrations of toxin, in a final assay volume of 300 μl. Non-specific binding was determined in the presence of 50 μM atropine. The reaction was stopped by addition of 3 ml of ice-cold buffer (10 mM Tris), immediately followed by filtration through Whatman GF/C glass fiber filters presoaked in 0.5% polyethyleneimine. The filters were washed once with 3 ml of ice-cold buffer (PBS) and dried, and the bound radioactivity was counted by liquid scintillation spectrometry. Each experiment was done at least three times.

The binding data from individual experiments (n ≥ 3) were analyzed by nonlinear regression analysis using Kaleidagraph 4.0 (Abelbeck/Synergy Software, Reading, PA) or Prism 4 (GraphPad Software Inc., San Diego, CA). After subtraction of the nonspecific binding through Whatman GF/C glass fiber filters presoaked in 0.5% polyethyleneimine. The filters were washed once with 3 ml of ice-cold buffer (PBS) and dried, and the bound radioactivity was counted by liquid scintillation spectrometry. Each experiment was done at least three times.

The binding data from individual experiments (n ≥ 3) were analyzed by nonlinear regression analysis using Kaleidagraph 4.0 (Abelbeck/Synergy Software, Reading, PA) or Prism 4 (GraphPad Software Inc., San Diego, CA). After subtraction of the nonspecific binding and normalization, competition data obtained with [^3H]NMS were analyzed using the allosteric ternary complex model (Ehlert, 1988). We used the following equation (Gharagozloo et al., 2002):

\[ B_{1X} = B_0 \times \left( \frac{(1 + K_x \times [L]) \times (1 + \alpha \times (K_X \times [X])^{\alpha})}{1 + (K_x \times [X])^{\alpha} + K_L \times [L] \times (1 + \alpha \times (K_X \times [X])^{\alpha})} \right) \]

where \( B_{1X} \) denotes the specific binding of the respective radioligand L ([^3H]NMS) in the presence of the cooperatively interacting agent X (allosteric agent). \( B_0 \) is the binding of L in the absence of X. \( K_x \) and \( K_L \) are the affinity constants for the binding of L and X, respectively, to the unliganded receptors. \( \alpha \) denotes the cooperativity factor for the allosteric interaction between X and L (with \( \alpha > 1, \alpha < 1 \), and \( \alpha = 1 \), indicating positive, negative, and neutral cooperativity, respectively). \( S_{1H} \) represents the slope factor of the binding curve of X. The value of \( K_X \) used in the data analyses was 10^{-10} M. Unconstrained fits for inhibition curves for R34A and Y36A had estimated slope factors less and greater than 1, respectively. However, these and all the other inhibition curves could be fitted satisfactorily with slope factors of 1, and the derived parameters (expressed as log values) are those given in the Tables. When the slope factors were constrained to 1, the estimates of \( \alpha \) and \( K_X \) (and \( \alpha \times K_X \)) were independent of whether the data were analyzed using the above equation to give values of \( \alpha \) and \( \log K_X \), or the equation recast to give values of \( \log \alpha \) and \( \log K_X \) or \( \log \alpha \) and \( \log K_X \).

Single time-point experiments were performed to measure the inhibition of the [^3H]NMS dissociation by wild-type and modified toxins, assuming a monoeponential dissociation kinetic as previously shown for MT7 (Mourier et al., 2003). These off-rate assays have been used previously to estimate the affinity of allosteric agents for the [^3H]NMS-occupied receptor (Lazareno et al., 2000). A high concentration of hM1 membranes (2–3 mg/ml) was incubated for 15 min with a high concentration of [^3H]NMS (5 nM). Then, 10-μl aliquots were added to tubes that were empty or contained 1 ml of 10 μM atropine, alone and with various concentrations of toxins. Non-specific binding was measured by incubating 10-μl aliquots with 10 μM atropine and [^3H]NMS (5 nM). After 45 min of incubation (approximately 2.5 dissociation half-lives), the samples were filtered and rinsed twice. The data were transformed to rate constants using the formula \( k_{sel} = \ln B_0/B_t/v_t \), where \( B_0 \) and \( B_t \) represent the [^3H]NMS bound initially and after \( v_t \) min of dissociation, respectively. The apparent rate constants for the dissociation of [^3H]NMS were determined in the presence of each concentration of toxins (k_{obs}) and divided by the true rate constant, \( k_{true} \), determined in the presence of atropine only (Gnagey et al., 1999). The resulting number, less than 1, indicates a slower dissociation of [^3H]NMS in the presence of the toxins, compared with the control rate constant, 0.04 min⁻¹. The values from triplicate assays were fitted using a nonlinear regression analysis to simple inhibition curves with slope factors of 1. The −log concentration inducing a half-maximal effect on [^3H]NMS dissociation (−logK_{diss}) according to the allosteric ternary complex model, is the log affinity constant of the toxin for the [^3H]NMS-occupied receptor.

Intracellular Calcium Assays. CHO cells stably expressing hM1 receptors were plated (3–5 × 10⁴ cells/well in 100 μl) on black-walled 96-well plates (Greiner Bio-One, Longwood, FL). Twenty-four hours after coating, the cells were first incubated with the toxins for 75 min and then for 45 additional min with the CaKit dye resuspended in Hanks’ balanced salt solution complemented with 20 mM HEPES, at pH 7.4 (Molecular Devices). The fluorescence was recorded using a FlexStation II plate reader (Molecular Devices Ltd, Wokingham, UK) with excitation and emission wavelengths fixed at 485 and 525 nm, respectively. Drug dilutions in assay buffer were prepared in a separate 96-well plate. Parameters for drug addition to the cell plate were preprogrammed, and delivery of the agonist carbamylcholine was automated through an eight-tip head pipettor 20 s after the beginning of the recording. Dose-response curves were constructed by measuring the fluorescence intensity after normalization to the maximal response to carbachol, measured in the absence of toxin. All data points were measured in duplicate. The measured EC_{50} for carbachol was 24 ± 4 nM.

Results

Crystallographic Structure of the Diiodo-MT7 Toxin Derivative. Crystallographic screening of the wild-type MT7 and its diiodo-Tyr51 derivative clearly showed the superiority of the modified toxin in yielding crystals suitable for high-resolution structural studies. Although the wild-type crystals suffered from a high degree of polymorphism, with the best crystals diffraction to only 3 Å, the MT7 diiodo derivative gave crystals that diffracted to atomic resolution. The high-resolution structure of this derivative was considered to be more informative than a low-resolution structure of the wild type, given that the addition of two iodine atoms is characterized by only a minor loss in affinity for the hM1 receptor (data not shown). The crystallographic structure of this modified MT7 was solved by molecular replacement and refined to 1.38 Å resolution with good stereochemistry (statistics in Table 1). The lattice includes two molecules in the asymmetric unit stabilized by the diiodo-tyrosines stacking against each other (Fig. 1A), explaining the superior crystallographic quality of this derivative compared with the wild-type toxin. The overall fold conforms to that of other three finger toxins (Fig. 1B). The electron density maps are excellent for the whole structure, only the tip of loop I (residues Ser8-Ile9-Trp10-Phe11) being less well defined than the other stretches.

Selection of the Modified Toxin Positions. The choice of the MT7 residues to be modified was based on both the sequence alignments of the different muscarinic toxins and on the crystallographic structure of the diiodo-MT7 toxin (Fig. 1B). The 12 amino acids selected to study the functional site by which MT7 interacts with the hM1 receptor are on the toxin surface and are mainly located at the tips of the three loops, regions previously identified as major functional determinants for other three-finger fold toxins interacting with nicotinic receptors (Servent and Ménez, 2001; Fruchart-Gail-
The selected residues are spread over the three toxin loops (Lys5, Ser8, Trp10, and Phe11 in loop I; Tyr30, Ser32, Arg34, Met35, and Tyr36 in loop II; Lys48, Tyr51, and Arg52 in loop III) and are located on both faces of the toxin (Fig. 1B, Table 1). In addition, these residues include those highly conserved in all the muscarinic toxin sequences (Ser8, Tyr30, Arg34, and Lys48), more variable (Tyr36), or specific to the MT7 toxin (Lys5, Trp10, Phe11, Ser32, Met35, Tyr51, and Arg52).

### Chemical Synthesis, Refolding, Purification, and CD Analyses of Wild-Type and Modified MT7 Toxins

The one-step solid phase chemical synthesis of the modified MT7 toxins was performed using a program previously developed for the synthesis of nicotinic and muscarinic toxins (Mourier et al., 2000, 2003). For each modified toxin, the synthesis proceeded smoothly, and the final TFA cleavage yielded a crude mixture in which the main component corresponded to the reduced form of each protein. These major products constituted 30 to 40% of the total reaction mixture, as analyzed by reversed-phase HPLC. Refolding of these toxins was performed in the presence of reduced and oxidized glutathione and 0.5 M guanidine hydrochloride (Mourier et al., 2003). The final yield varied from 18 to 35% for the different toxins. The refolded modified synthetic MT7 toxins were purified to homogeneity on a C18 reversed-phase HPLC column. Finally, the purity and identity of the different toxins was evaluated using [3H]NMS as tracer. To avoid kinetic artifacts caused by the ability of the MT7s to inhibit [3H]NMS association (and dissociation) the assays were carried out overnight to ensure that equilibrium was established. All the wild-type and modified toxins caused a concentration-dependent reduction of specific [3H]NMS binding to the hM1 receptor (illustrated for selected mutants in Fig. 3).

**TABLE 1**

<table>
<thead>
<tr>
<th>Statistics from crystallographic analysis of the 3D structure of the MT7 diiodo Y51-derivative.</th>
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<tbody>
<tr>
<td><strong>Data Collection</strong></td>
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<tr>
<td>Resolution (Å)</td>
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<tr>
<td>Total observations</td>
</tr>
<tr>
<td>Unique reflections</td>
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<tr>
<td>Completeness (%)</td>
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<tr>
<td>Space group</td>
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<tr>
<td>Unit cell parameters (Å; °)</td>
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<tr>
<td>a = 27.76, b = 56.57, c = 45.30, b = 95.17</td>
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<tr>
<td>Matthews’s coefficient (V_M, Å³/Da)</td>
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<tr>
<td><strong>Refinement</strong></td>
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<td>Outer shell (Å)</td>
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<td>Number of water molecules</td>
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<td>R cryst (%)</td>
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<td>Ramachandran Plot</td>
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<td>Residues in additional regions (%)</td>
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<td>Residues in disallowed regions (%)</td>
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<tr>
<td>Average B value</td>
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<tr>
<td>(main chain, side chain Å²)</td>
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<tr>
<td>Molecule A: 13.3, 15.0</td>
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<tr>
<td>Molecule B: 13.1, 15.0</td>
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</table>

**R cryst = \( \Sigma_{hkl} | F_{hkl} | - | F_{m std} | / \Sigma_{hkl} | F_{hkl} | \), where \( I_{hkl} \) is the mean intensity of the \( i \) observations of symmetry-related reflections of \( h \).

**R free = \( \Sigma_{hkl} | F_{hkl} | - | F_{calc} | / \Sigma_{hkl} | F_{hkl} | \), where \( F_{calc} \) is the calculated protein structure factor from the atomic model (\( R_{free} \) was calculated using 5% of the reflections).

**Root-mean-square deviation in bond lengths and angles are the deviations from ideal values.

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**Fig. 1.** Crystallographic structure of the MT7 diiodo derivative. A, the lattice includes two toxin molecules in the asymmetric unit stabilized by the diiodo-tyrosines stacking against each other. B, three-dimensional structure of this modified MT7 refined to 1.38-Å resolution. Location of all the residues studied in this article is indicated on the three toxin loops.
When analyzed using the allosteric ternary complex model, the different toxins showed negative cooperativity with [³H]NMS. Their affinity constants for the unoccupied receptor, (log $K_x$), calculated as described under Materials and Methods section, are reported in Table 2. The value for the wild-type toxin was $10.47 \pm 0.04$ (n = 20). The single amino acid modifications K5A, S8A, F11A, Y30A, S32A, K48A, Y51A, and R52A individually induced no, or small, decreases (less than 4-fold) in the affinity of MT7 for the free hM₁ receptor, whereas weak but significant $K_x$ decreases (6-20-fold) were associated with mutations W10A, M35A, and Y36A. Finally, a major effect was observed at the tip of the central loop, where the R34A modification decreased the affinity of MT7 for the hM₁ receptor approximately 200-fold (Fig. 3) as found previously (Mourier et al., 2003). The effects of these modifications were confirmed by direct competition binding experiments with [¹²⁵I]MT7ox as tracer, where the same order of decrease in toxin affinity generated by the different modifications was observed (data not shown).

As illustrated in Fig. 3 there is a residual binding of [³H]NMS which is not displaceable by high concentrations of toxin. This is a characteristic of the inhibition curves of negatively cooperative allosteric ligands and represents the toxin-receptor-[³H]NMS complex. The fractional residual binding was dependent on nature of the modification to the toxin, being small for wt and R52A and larger for S32A, Y36A and Y36A. All the inhibition curves were analyzed according to the ternary complex model (equation details under Materials and Methods) to give estimates of the log affinities of the toxins for the [³H]NMS occupied receptor (log $o/k_{o}$) and are reported in Table 2. All the inhibition curves could be fit satisfactorily with slope factors of 1. Although the precision of the data are good (S.E.M. values generally in the 1-5% range), the values of log $o/k_{o}$ are quite difficult to estimate accurately, because for most modified toxins, the part of the inhibition curve that determines this value is barely above nonspecific binding values and may be only 3 to 5% of the specific [³H]NMS signal (e.g., Fig. 3) because of the 100-fold or greater negative cooperativity between NMS and the toxins. Where there is high negative cooperativity, small changes in the estimates of nonspecific binding result in large changes in the estimates of log $o/k_{o}$ ($\alpha$ and their 95% confidence limits) but very small changes (<0.03 log unit) in the estimates of log $K_x$ (data not shown).

To evaluate the potential effect of the different modifications introduced into the toxin sequence on the selectivity of the MT7-hM₁ interaction, the inhibitory potency of a high concentration of each toxin (1 μM) was examined on the M₂-M₅ receptor subtypes. At this concentration, none of the modified toxins was able to displace [³H]NMS from these receptor subtypes, indicating that their potencies at these subtypes were above 1 μM (data not shown). Thus, the various modifications introduced into the MT7 sequence have no detectable effect on the large selectivity of MT7 for the hM₁ receptor.

**Dissociation Experiments to Measure the Binding of Wild-Type and Modified MT7 Toxins to the NMS-Occupied hM₁ Receptor.** Because the formation of ternary complexes is the characteristic of allosteric interactions, we have studied the effect of wild-type and modified toxins on the kinetics of binding of [³H]NMS to the hM₁ receptor. As described previously (Gnagey et al., 1999; Lazareno et al., 2000; Mourier et al., 2003), single-time-point experiments measuring the effect of allosteric ligands on the [³H]NMS dissociation, can be used to characterize the interaction of these allosteric agents with the NMS-occupied receptor. Thus, we studied the concentration-dependence of the effect of wild-type and modified MT7 toxins on [³H]NMS dissociation, as shown in Fig. 4. All data could be fitted satisfactorily to simple inhibition curves with slope factors of 1 and extrapolating to a $k_{dis}/k_{in}$ value of zero at high toxin concentrations. The log $K_{dis}$ calculated for the wild-type toxin was $8.59 \pm 0.02$ (n = 10). Modifications of Lys5 and Phe11 in loop I, Ser32 in loop II, or Lys48 and Tyr51 in loop III only had no, or small, effects (≤4-fold) on log $K_{dis}$. However, 6- to 700-fold affinity decreases were associated with the modifications M35A, R34A, S8A, Y30A, R52A, W10A, and Y36A (Table 2). Therefore, all three loops of the MT7 toxin seem to be in-

Fig. 2. Far-ultraviolet CD spectra of the wild-type and different modified toxins. The spectra were monitored in 5 mM sodium phosphate buffer, pH 7.4, at 20°C at a peptide concentration of $5 \times 10^{-6}$ M.

![Fig. 2](image-url)

Fig. 3. Inhibition of [³H]NMS binding to hM₁ receptor by wild-type and modified MT7 toxins. Binding experiments were performed by incubating membrane fractions of receptor with [³H]NMS (0.5 nM) and varying concentrations of toxin at room temperature, overnight. The total specific binding in each experiment was 1500 ± 300 cpm. The results are expressed as the ratio of the specific [³H]NMS binding measured with (B) or without toxin (B₀) (n = 3 to 10). The curve fits were based on the ternary complex equation reported under Materials and Methods.
volved in its binding to the [\(^3\)H]NMS-liganded receptor with critical roles for Trp10 and Tyr36 in loops I and II, respectively, because they produced the largest changes.

According to the allosteric ternary complex model, the affinity of the allosteric ligand for the NMS-occupied receptor obtained from the dissociation studies (log\(K_{\text{diss}}\)) should be equal to the estimate from the equilibrium data (log\(a\times K_X\)) if the two manifestations of an allosteric interaction are to have a common mechanism and explanation (Lazareno and Birdsall, 1995). The primary criteria used in this study are that these two estimates should differ by less than 3-fold or that the 95% confidence limits of the two estimates should overlap. The results in Table 2 show these criteria to be satisfied for MT7 and most (8 of 12) of the modified toxins. However there are \(\approx\)10-fold discrepancies for W10A, Y30A, Y36A, and R52A. As noted in the previous section, small changes, for example in the nonspecific binding estimates, can generate large changes in the values of log\(a\times K_X\) and their confidence limits, and the data for the four constructs then satisfy the criteria. Further possible explanations for these discrepancies are considered under Discussion.

It is possible to calculate two estimates of the negative cooperativity from the binding data, log\(a^\circ\) = log\(K_{\text{diss}}\) - log\(K_X\) and log\(a^\circ\) = log\(a\times K_X\) - log\(K_X\). These are presented in Table 2. Many of the values of log cooperativity are in the range \(-1.8\) to \(-2.1\), reflecting those modifications to the toxin that do not differentially affect their binding to the [\(^3\)H]NMS-occupied and free receptor. Both data sets show higher values of log cooperativity for S32A, R34A and M35A, illustrating a less inhibitory effect of the binding of these residues of loop II to the [\(^3\)H]NMS-occupied receptor, relative to the free receptor. Log\(a^\circ\) shows a large effect on S8A, W10A, Y30A, Y36A, and R52A, showing a selective inhibitory effect of these modifications on the binding of the three loops of the toxins to the NMS occupied receptor, as measured by their effects on [\(^3\)H]NMS dissociation.

**Functional Assays.** The effects of wild-type and modified toxins on the function of hM\(_1\) receptors stably expressed at the surface of CHO cells was evaluated by measuring changes in the intracellular calcium concentration using a calcium fluorescent dye. First, the effect of micromolar concentrations of all the toxins was evaluated directly on the cells and the signal was compared with the control obtained with carbamylcholine (CCh; 300 nM). As shown in Fig. 5A, neither the wild-type nor the modified MT7 toxin was able to activate the hM\(_1\) receptor, demonstrating that these toxins are not agonists. Precubination of the cells with increasing concentrations of MT7 was, however, able to abolish, in a dose-dependent manner, the CCh (100 nM) activation of the hM\(_1\) receptor, as shown in Fig. 5B. The potency of MT7 was estimated to 2.2 ± 0.5 nM. Analogous experiments were performed with the modified toxins. The dose-response curves show that all these toxins were able to inhibit the CCh-induced hM\(_1\) activation, but with variable potencies, as reported in Table 3 and illustrated in Fig. 6. All inhibition curves were fitted to a slope factor of 1. There was evidence that the slopes of the inhibition curves for MT7 and Y36A were significantly greater than 1 but fitting all the inhibition curves with the slope factor as a variable resulted in no changes in pIC\(_{50}\) greater than 0.03 log unit. Four modifications induced greater than 4-fold decreases in functional potency, varying from 5-fold for S32A to 10-, 20-, and 50-fold for R52A, R34A, and Y36A, respectively.

**Discussion**

Current evidence points to the high affinity, large subtype selectivity, and allosteric nature of the interaction of the MT7 toxin with the hM\(_1\) receptor (Max et al., 1993; Carsi and Potter, 2000; Olianas et al., 2000; Mourier et al., 2003; Fruchart-Gaillard et al., 2006). To better understand how MT7 binds to the free and NMS-occupied hM\(_1\) receptor and to progress toward the structural modeling of these interactions, we have investigated how different single amino acid modifications introduced into the MT7 sequence altered its potency, determined using both equilibrium and dissociation kinetics experiments as well as functional assays. The modified residues include amino acids present specifically in the MT7 toxin or highly conserved in all the muscarinic toxin sequences and located on both faces of the three toxin loops. The CD spectra of all the purified chemically synthesized toxins, compared with that of the wild-type toxin, suggested that the modifications introduced into the MT7 sequence had little effect, if any, on the overall structure of the toxin.

**Table 2**

Affinity constants of wild-type and modified MT7 toxins calculated from equilibrium binding experiments with [\(^3\)H]NMS using the ternary complex equation (log\(K_X\) and log\(a\times K_X\)) and from their potency to inhibit the dissociation of [\(^3\)H]NMS (log\(K_{\text{diss}}\))

<table>
<thead>
<tr>
<th>Modified MT7</th>
<th>Loop</th>
<th>Face</th>
<th>(\log K_X)</th>
<th>(K_{\text{diss}}/K_{\text{diss}(\text{wt})})</th>
<th>(\log(K_{\text{diss}}))</th>
<th>(K_{\text{diss}}/K_{\text{diss}(\text{wt})})</th>
<th>(\log a^\circ)</th>
<th>(\log a^\circ)</th>
</tr>
</thead>
<tbody>
<tr>
<td>MT7</td>
<td></td>
<td></td>
<td>10.47 ± 0.04</td>
<td>1</td>
<td>8.66 ± 0.12</td>
<td>1</td>
<td>−1.9</td>
<td>−1.8</td>
</tr>
<tr>
<td>K5A</td>
<td>I</td>
<td>Concave</td>
<td>10.15 ± 0.08</td>
<td>2</td>
<td>8.08 ± 0.30</td>
<td>8.20 ± 0.05</td>
<td>−2.0</td>
<td>−2.1</td>
</tr>
<tr>
<td>S8A</td>
<td>I</td>
<td>Concave</td>
<td>10.18 ± 0.05</td>
<td>2</td>
<td>8.12 ± 0.20</td>
<td>7.52 ± 0.08</td>
<td>−2.7</td>
<td>−2.1</td>
</tr>
<tr>
<td>W10A</td>
<td>I</td>
<td>Concave</td>
<td>9.68 ± 0.11</td>
<td>6</td>
<td>7.90 ± 0.07</td>
<td>6.62 ± 0.07</td>
<td>−3.0</td>
<td>−1.8</td>
</tr>
<tr>
<td>F11A</td>
<td>I</td>
<td>Convex</td>
<td>10.43 ± 0.11</td>
<td>1</td>
<td>8.55 ± 0.40</td>
<td>8.02 ± 0.04</td>
<td>−2.4</td>
<td>−1.9</td>
</tr>
<tr>
<td>Y30A</td>
<td>I</td>
<td>Concave</td>
<td>10.21 ± 0.05</td>
<td>2</td>
<td>8.36 ± 0.17</td>
<td>7.37 ± 0.03</td>
<td>−2.8</td>
<td>−1.9</td>
</tr>
<tr>
<td>S32A</td>
<td>I</td>
<td>Concave</td>
<td>9.97 ± 0.06</td>
<td>3</td>
<td>8.74 ± 0.08</td>
<td>8.60 ± 0.09</td>
<td>−1.4</td>
<td>−1.2</td>
</tr>
<tr>
<td>R34A</td>
<td>I</td>
<td>Plan</td>
<td>8.22 ± 0.07</td>
<td>200</td>
<td>7.35 ± 0.10</td>
<td>7.63 ± 0.04</td>
<td>−0.6</td>
<td>−0.8</td>
</tr>
<tr>
<td>M35A</td>
<td>I</td>
<td>Convex</td>
<td>9.55 ± 0.04</td>
<td>8</td>
<td>7.94 ± 0.07</td>
<td>7.82 ± 0.08</td>
<td>−1.7</td>
<td>−1.6</td>
</tr>
<tr>
<td>Y36A</td>
<td>I</td>
<td>Convex</td>
<td>9.16 ± 0.06</td>
<td>80</td>
<td>7.66 ± 0.11</td>
<td>7.57 ± 0.24</td>
<td>−3.4</td>
<td>−1.5</td>
</tr>
<tr>
<td>K48A</td>
<td>I</td>
<td>Concave</td>
<td>10.36 ± 0.06</td>
<td>1</td>
<td>8.61 ± 0.13</td>
<td>8.37 ± 0.06</td>
<td>−2.0</td>
<td>−1.8</td>
</tr>
<tr>
<td>Y51A</td>
<td>III</td>
<td>Concave</td>
<td>10.29 ± 0.07</td>
<td>2</td>
<td>8.33 ± 0.25</td>
<td>8.38 ± 0.02</td>
<td>−2.0</td>
<td>−1.9</td>
</tr>
<tr>
<td>R52A</td>
<td>I</td>
<td>Concave</td>
<td>10.20 ± 0.03</td>
<td>2</td>
<td>8.35 ± 0.09</td>
<td>7.10 ± 0.07</td>
<td>−3.1</td>
<td>−1.8</td>
</tr>
</tbody>
</table>

\(\log K_X\), log affinity of the toxin for the unoccupied receptor; \(\log a\times K_X\), estimate of the log affinity of the toxins for the NMS-occupied receptor obtained by fitting the inhibition binding curves with the ternary complex equation described under Materials and Methods; \(\log K_{\text{diss}}\), log potency of the toxins to inhibit [\(^3\)H]NMS dissociation; \(\log a^\circ\), estimate of the log cooperativity factor, expressed as log\(K_{\text{diss}}\) - log\(K_X\); \(\log a^\circ\), estimate of the log cooperativity factor, expressed as log\(a\times K_X\) - log\(K_X\).
The determination of the crystallographic structure of a diiodo-MT7 analog, confirms the previous assumption that MT7 belongs to the three-finger fold structural family, which includes several toxins acting at the cholinergic synapse. Loops II and III are similar to those found in nicotinic toxins, whereas loop I is somewhat different, being more distant from the other two loops and appearing to be unusually flexible at its extremity. The side chain of Phe11 is poorly defined, a rare occurrence for this type of residue at this resolution. The surrounding residues, also hydrophobic in nature, are similarly affected. The side chain of Ile9 is better ordered than that of Phe11 but less than that of Trp10. Loop I of MT2, another muscarinic toxin, shares such disorder, but this is not particularly unusual given its sequence: Ile9-Gly10-Gly11 (R. Ménez and E. Stura, unpublished observations). Considering these two structures together, it is plausible that such an unusual feature may have a functional role in these toxins. One should also be aware that the extra stabilization offered in the crystal structure by the interaction of the diiodo-tyrosines could mask some additional flexibility of loop III both in the wild type toxin and in the diiodo analog in solution. The lower conformational constraints of muscarinic toxins, together with the variations in sequence,
may explain the broad pharmacological diversity that exists within the muscarinic toxin family.

Identification from equilibrium binding experiments of the MT7 residues involved in its interaction with the free hM1 receptor highlights the crucial role of the tip of the toxin loop II (Arg34, Met35, Tyr36), the highly conserved Arg34 being particularly important (Table 2). The high affinity and selectivity of the MT7 for the hM1 receptor may be associated with the involvement of residues specific to this toxin, such as Trp10 and Met35, located at the tips of the toxin loops I and II, respectively. The MT7 specific Lys65 residue, previously reported not to play a role in determining selective binding of this toxin (Krajewski et al., 2001) is located in the upper part of the toxin structure, a region probably too far from the toxin interaction site to be important.

MT7 has been reported to decrease the dissociation rate of [3H]NMS (Olianas et al., 2000; Krajewski et al., 2001; Mourier et al., 2003). This effect confirms the ability of this toxin to form a ternary complex with NMS and M1 receptor, which is a hallmark of an allosteric interaction. The ability of MT7 derivatives to decrease [3H]NMS dissociation allowed us to identify toxin residues involved in its binding to the allosteric site when the orthosteric site is occupied by NMS. Alanine substitution introduced into the toxin’s sequence induced up to 700-fold decreases in the toxin IC50diss values (Table 2) and revealed major roles for Ser8, Trp10, Tyr30, and Arg52, as well as Arg34, Met35, and Tyr36, in this interaction. It is interesting to note that these residues (excluding Arg34) are all specific to MT7 and may be related to its allosteric property, which is not observed in the interaction of the homologous MT1 with the hM1 receptor (Mourier et al., 2003).

The wild-type and modified MT7 toxins showed negative cooperativity with [3H]NMS, as indicated by the log cooperative factors ($\log \alpha$ in Table 2) derived from the equilibrium binding curves when analyzed with the ternary complex equation. These values were similar for all the toxins (−1.6 to −2.1) with the exception of the S32A, and R34A toxins (−0.8 to −1.2). Furthermore, according to the ternary model, the affinity of the MT7 for the NMS-occupied receptor can be calculated as 8.66, a value similar to the one obtained from the single-time point experiments (8.59). This agreement also occurs for most but not all of the modified toxins, notably W10A, Y30A, Y36A, and R52A.

There are a number of possible explanations for the discrepancies. Some of the inherent problems of determining accurate values of $\log \alpha \times K_X$ because of the high negative cooperativities between NMS and the toxins have been stated under Results. For two of these modified toxins, W10A and Y36A, the overall toxin binding affinities have also been strongly decreased; therefore, high concentrations of toxins were required to observe the manifestations of negative cooperativity in the equilibrium or kinetic experiments. If, during the long duration equilibrium experiments, there were aggregation problems with high concentrations of some modified toxins (e.g., Y36A), this could result in an overestimation of $\log \alpha \times K_X$. On the other hand, if there were slow isomerizations of the toxin-receptor-NMS complex, then the values of $\log K_{diss}$ might be underestimated relative to $\log \alpha \times K_X$ because of the relatively short duration of the dissociation experiments. In general, we feel that the effects of the toxins on [3H]NMS dissociation provide a more direct and accurate

![Fig. 7. Comparison of the binding residues through which MT7 toxin interacts with the free (A) or NMS-occupied (B) hM1 receptor. The three-dimensional structure of diiodo-MT7 is shown as a space-filling representation. Residues in which alanine substitution caused a decrease in potency of less than 5-fold are shown in green, between 5- and 10-fold in yellow, between 10- and 90-fold in orange, and higher than 90-fold in pink. The violet group shows the iodide molecules of the diiodo-Tyr51. Ser32, green in A and B, is not visible in this view.](image-url)
estimate of the affinities of the modified toxins for the [3H]NMS-occupied receptor.

The interaction of MT7 with the hM1 receptor was also studied in functional assays based on the measurement of the calcium release in CHO-M1 cells. First, the lack of agonism of the MT7 was observed by its inability to generate a signal, and its antagonist property was confirmed by its concentration-dependent inhibition of the CCh activation with nanomolar potency (Fig. 5, Table 3). This value is compatible with those reported for the functional characterization of MT7 by measuring the effect of MT7 on CCh-induced calcium release (Kukkonen et al., 2004), CCh-induced [3H]IPs accumulation (Olianas et al., 2000), or ACh stimulation of the [35S]GTP[S] binding (Olianas et al., 2000). The effect of alanine-substitution on MT7 pIC50 confirms the major role of the tip of the toxin loop II (Ser92, Arg54, Tyr36) in its inhibitory actions. The potency estimate of MT7 in this assay is approximately 60-fold lower than its dissociation constant found in the binding studies. This comparison is confounded by the fact that the IC50 value may be influenced by the cooperativity between CCh and the MT7 and the method of the functional assay [i.e., use of a concentration of carbamycholine well above its EC50, the fact that the equilibrium between the CCh and MT7 was probably not established during the time course of the assay because of the slow dissociation of the MT7 from the receptor (hemiequilibrium), and the possibility of slow isomerization of the MT7-receptor complex]. Hemi-equilibrium and isomerization may or may not be present in the modified toxins, especially when the binding of the toxins is substantially weaker than the wild-type toxin. Given these caveats, there is excellent agreement with the fold-change in functional potency and the fold-change in binding to the unoccupied receptor relative to wild-type MT7 (<3-fold discrepancy) for most modified toxins. Somewhat larger differences are observed for the two toxins (R34A and R52A) where the modifications induce a large change in binding affinity and potentially, as alluded to above, to their kinetic properties.

Comparison of the two binding sites to which MT7 interacts, the free and the NMS-occupied hM1 receptor, points to a major role for the tip of the central loop of MT7, which is responsible in both cases for a major part of the interaction energy (Fig. 7, A and B). This observation agrees with results reporting the interaction of structurally related toxins with other cholinergic targets, such as nicotinic acetylcholine receptors (Fruchart-Gaillard et al., 2002; Bourne et al., 2005) and acetylcholinesterase (Bourne et al., 1995). Furthermore, the fact that both interaction sites include mainly aromatic (Tyr36, Trp10, Tyr30) and positively charged residues (Arg34, Arg52), suggest that cation-π and electrostatic interactions might occur in the MT7-hM1 interactions, as described previously in the Cbtx-AChBP and fasciculin-acetylcholinesterase crystallographic structure complexes (Bourne et al., 1995, 2005). The toxin interaction site on the hM1 receptor might be located at the outer loops or at the beginning of the transmembrane domains of the receptor, assuming that the toxin doesn’t penetrate deep inside the receptor (Kukkonen et al., 2004).

The variable energetic contribution of each residue of the MT7 loop II in its interaction with free or occupied receptor (e.g., Arg54 and Tyr36) and the fact that some MT7 residues seem specific for its interaction with only one receptor state, such as Ser8, Tyr30, and Arg52 for the NMS-occupied receptor, show that the interaction of MT7 with M1 receptor is not identical in the presence or absence of NMS at the orthosteric site. Thus, it seems that the presence of NMS in the acetylcholine-binding pocket induces some conformational modifications in the receptor’s transmembrane domains and extracellular regions, leading to appropriate adjustments of the MT7 location compared with its interaction with the unoccupied receptor. Site-directed mutagenesis of receptor residues is in progress to identify the toxin’s interaction site on the receptor, allowing the use of the mutant-cycle approach and construction of a structural model of the toxin-muscarinic receptor interaction. Such a model should be useful to better understand the different pharmacological and functional profiles of muscarinic toxins and in the future design of molecules with predetermined subtype and allosteric specificities.

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References


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