STRUCTURE-FUNCTION STUDIES OF ALLOSTERIC AGONISM AT M₂ MUSCARINIC ACETYLCHOLINE RECEPTORS

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Abbreviations: AC-42, 4-n-butyl-1-[4-(2-methylphenyl)-4-oxo-1-butyl] piperidine hydrogen chloride; C7'/3-phth, heptane-1,7-bis-[dimethyl-3’-phthalimidopropyl]-ammonium bromide; 77-LH-28-1, 1-[3-(4-butyl-1-piperidinyl)propyl]-3,4-dihydro-2(1H)-quinolinone; McN-A-343, 4-[(3-chloro phenyl)carbamoyloxy)-2-butynyltrimethylammmonium chloride; mAChR, muscarinic acetylcholine receptor; [3H]NMS, [3H]N-Methyl-scopolamine; [35S] GTPγS, [35S] Guanosine 5’-(γ-thio)triphosphate; GPCR, G protein-coupled receptor; ACh, acetylcholine; ATCM, allosteric ternary complex model; ERK1/2, extracellular signal regulated kinases 1 and 2; DMEM, Dulbecco’s modified eagle medium; FBS, fetal bovine serum; CHO, Chinese hamster ovary; GDP, Guanosine 5’-diphosphate; Tris, Tris(hydroxymethyl)aminomethane; EGTA, Ethylene glycol-bis(2-amino-ethylether)-N,N,N’, N’-tetra-acetic acid; EDTA, Ethylenediamintetra-acetic acid.
Abstract

The M₂ muscarinic acetylcholine receptor (mACHR) possesses at least one binding site for allosteric modulators that is dependent on the residues ¹⁄₂EDGE¹⁷⁵, Y¹⁷⁷ and T⁴²³. However, the contribution of these residues to actions of allosteric agonists, as opposed to modulators, is unknown. We created mutant M₂ mAChRs where the charge of the ¹⁄₂EDGE¹⁷⁵ sequence had been neutralized and each of Y¹⁷⁷ and T⁴²³ were substituted with alanine. Radioligand binding experiments revealed that these mutations had a profound inhibitory effect on the prototypical modulators gallamine, alcuronium and C₇/₃-phth, but minimal effects on the orthosteric antagonist, [³H]NMS. In contrast, the allosteric agonists, McN-A-343, AC-42 and a novel AC-42 derivative, 77-LH-28-1, demonstrated an increased affinity or proportion of high affinity sites at the combined EDGE-YT mutation, indicating a different mode of binding to the prototypical modulators. Subsequent functional assays of ERK1/2 phosphorylation and [³⁵S]GTPγS binding revealed minimal effects of the mutations on the orthosteric agonists, ACh and pilocarpine, but a significant increase in the efficacy of McN-A-343 and potency of 77-LH-28-1. Additional mutagenesis experiments found that these effects were predominantly mediated by Y¹⁷⁷ and T⁴²³, rather than the ¹⁄₂EDGE¹⁷⁵ sequence. The functional interaction between each of the allosteric agonists and ACh was characterized by high negative cooperativity, but was consistent with an increased allosteric agonist affinity at the combined EDGE-YT mutant M₂ mAChR. This study has thus revealed a differential role of critical allosteric site residues on the binding and function of allosteric agonists versus allosteric modulators of M₂ mAChRs.
Muscarinic acetylcholine receptors (mAChRs) are a group of five Family A G protein-coupled receptors (GPCRs) distributed throughout the body (Christopoulos, 2007; Hulme et al., 1990). Drugs targeting mAChRs are currently used in the treatment of chronic obstructive pulmonary disorder and urinary incontinence. These receptors also represent potential therapeutic targets for conditions such as Alzheimer’s disease, schizophrenia and irritable bowel syndrome (Eglen et al., 2001; Felder et al., 2000). To date, however, the widespread development of highly efficacious mAChR therapeutics with acceptable side effect profiles has been limited by a relative lack of ligands with sufficient subtype selectivity (Eglen et al., 2001; Felder et al., 2000).

Allosteric modulation of GPCRs represents a novel therapeutic avenue for overcoming difficulties associated with selective drug targeting (Christopoulos, 2002; May et al., 2007), and may be particularly amenable to mAChRs. Functional and radioligand binding studies have provided evidence for at least two allosteric binding sites on each mAChR subtype (Christopoulos et al., 1998; Ellis et al., 1991; Lazareno et al., 2000), with one of the sites recognized by prototypical mAChR allosteric modulators such as gallamine, alcuronium and heptane-1,7-bis-[dimethyl-3’-phthalimidopropyl]-ammonium bromide (C7’3-phth) (Lanzafame et al., 1997). The allosteric interactions mediated by these mAChR modulators are for the most part adequately described by a simple allosteric ternary complex model (ATCM; Ehlert, 1988; Lazareno and Birdsall, 1995), with the effects of the modulators largely restricted to either enhancing or inhibiting orthosteric ligand binding affinity; minimal, if any, intrinsic efficacy has been detected for these compounds (but see Zahn et al., 2002)).
It is now recognized, however, that some GPCR allosteric ligands have the capacity to affect receptor signaling in the absence of orthosteric agonist (Langmead and Christopoulos, 2006). Such “allosteric agonists” represent an important expansion in the chemical space surrounding allosteric modulators, since they have the potential to modulate orthosteric ligand pharmacology in addition to perturbing cellular signaling in their own right. Two mAChR ligands suggested to act this way are the functionally selective partial agonists, 4-(N-(3-chlorophenyl)carbamoyloxy)-2-butynyltrimethyl-ammonium chloride (McN-A-343) and 4-n-butyl-1-[4-(2-methylphenyl)-4-oxo-1-butyl] piperidine hydrogen chloride (AC-42) (Figure 1A). Both compounds have been reported to cause incomplete inhibition of the binding of the orthosteric antagonist, [³H]N-methylscopolamine ([³H]NMS), when present at saturating concentrations at rat M₂ (McN-A-343) and human M₁ (AC-42) mAChRs, as well as retarding [³H]NMS dissociation (Birdsall et al., 1983; Langmead et al., 2006; Waelbroeck, 1994); these phenomena are characteristics of the formation of a ternary complex between the receptor and two concomitantly-bound ligands. Despite these observations, it remains to be determined if the allosteric effects of these agonists are receptor subtype or species-dependent, what the relationship is between the common allosteric site recognized by prototypical modulators and that recognized by allosteric agonists, and whether both the agonistic and allosteric modulator properties of these latter compounds are mediated via the same (allosteric) domain on the receptor, or whether they reflect differential interactions with the orthosteric site (agonism) and an allosteric site (modulation).

Using the M₂ mAChR as a model, prior mutagenesis studies have found that the common allosteric site recognized by classic mAChR modulators comprises the second extracellular loop and the interface between the third extracellular loop and the top of transmembrane (TM) domain.
7; specifically, the highly acidic \textsuperscript{172}EDGE\textsuperscript{175} sequence (Leppik et al., 1994), as well as Y\textsuperscript{177} in the second extracellular loop, N\textsuperscript{419} at the junction of the third extracellular loop and TM7, and T\textsuperscript{423} near the top of TM7 (Figure 1B; Huang et al., 2005; Voigtlander et al., 2003). The conserved W\textsuperscript{422} in TM7 has also been implicated in the actions of allosteric modulators of mAChRs (Matsui et al., 1995; Prilla et al., 2006). Importantly, most mutations in these regions that affect allosteric modulator binding have minimal effects on binding of orthosteric ligands.

In contrast to the studies on prototypical modulators, there have been no reports to date on the effect of allosteric site mutations on the actions of allosteric agonists. Thus, the aim of the current study was to investigate the mode of action of McN-A-343, AC-42 and a novel AC-42 derivative, 1-[3-(4-butyl-1-piperidinyl)propyl]-3,4-dihydro-2(1H)-quinolinone (77-LH-28-1; Figure 1A), that we have recently found to be a more potent M\textsubscript{1}-selective agonist than AC-42 (Langmead et al., manuscript on preparation) at the human M\textsubscript{2} mAChR, and to assess the contribution of the “common allosteric site” \textsuperscript{172}EDGE\textsuperscript{175}, Y\textsuperscript{177} and T\textsuperscript{423} epitopes of the M\textsubscript{2} mAChR on the pharmacology of these agents.
Materials and Methods

Materials

Dulbecco’s modified eagle medium (DMEM), penicillin-streptomycin, hygromycin-B, zeocin and geneticin were purchased from Invitrogen (Carlsbad, CA). Fetal bovine serum (FBS) was purchased from ThermoTrace (Melbourne, VIC, Australia). [3H]NMS (82.0 Ci/mmol) and [35S]Guanosine 5’-(γ-thio)triphosphate ([35S]GTPγS) (1250 Ci/mmol) were purchased from PerkinElmer (Boston, MA, USA). McN-A-343 was purchased from Research Biochemicals (Natick, MA). C7/3-phth was synthesized at the Institute of Drug Technology (Boronia, Victoria, Australia) whereas AC-42 and 77-LH-28-1 were synthesized in-house at GlaxoSmithKline, Harlow. Alcuronium chloride was a generous gift from F. Hoffmann-La Roche (Basel, Switzerland) and the SureFire™ cellular ERK1/2 assay kits were a generous gift from TGR BioSciences (Adelaide, Australia). AlphaScreen reagents were from PerkinElmer (Boston, MA). All other reagents were purchased from Sigma-Aldrich (St. Louis, MO) or BDH Merck (Victoria, Australia).

Receptor mutagenesis

The coding sequence of the human M2 mAChR, obtained from the UMR cDNA Resource Centre (www.cdna.org), was cloned into the Gateway recombination Entry vector, pENTR/D-TOPO, using the pENTR directional TOPO® cloning kit (Invitrogen, Mt. Waverley, Australia) according to the manufacturer’s instructions, following amplification of the gene using the following primers 5’-CACCATGAATAACT CAACAAACTCC-3’ (N-terminal forward primer with CACC sequence) and 5’-TTACCTTGTAGCGCCTATGTTC-3’ (C-terminal reverse primer). The native stop codon was subsequently mutated to Lys using the QuickChange Multi mutagenesis kit (Stratagene, La Jolla, CA), according to the manufacturer’s instructions, prior to subcloning of
the receptor into pEFS/FRT/V5-DEST Gateway destination vector. Transfer of the M² mAChR from pENTR/D-TOPO into pEFS/FRT/V5-DEST was achieved using the LR Clonase enzyme mix kit (Invitrogen) and resulted in in-frame insertion of the V5 epitope tag at the C-terminus of the receptor. This receptor sequence is referred to as “wild-type” throughout this study. Analysis of the properties of this clone following stable transfection in CHO cells revealed equivalent pharmacological properties to human M² mAChRs studied previously where the native stop codon was intact (Avlani et al., 2004). To study the influence of specific amino acids in receptor function, mutations were introduced into the wild-type receptor in pENTR/D-TOPO by site-directed mutagenesis using the QuickChange kit or QuickChange Multi kit (Y¹⁷⁷A + T⁴²³A mutant). Mutant receptors were subsequently subcloned into the pEFS/FRT/V5-DEST vector as described above. Oligonucleotides for site-directed mutagenesis and DNA sequencing were purchased from GeneWorks (Hindmarsh, Australia). The primers used for the site-directed mutagenesis reactions are as follows: Wild Type (with V5 tag) - 5' CATAGGCGCTACAAGGAAAAGGTTGGGC 3' (Quick Change Multi; single primer); ¹⁷²EDGE¹⁷⁵-QNGQ  5' GGGGTGAGAACTGTGCAGAATGAGGAGTGCTACATTGAG 3' (forward); 5’ CTGAATGTAGCAGCTGTGCCATTGTGCAGTTCCCC 3’ (reverse); Y¹⁷⁷A - 5’ GAGGATGGGGAGTGCACCTCATTTTTCC 3’ (Quick Change Multi; single primer); T⁴²³A - 5’ CCAACACACTGTGCAGGCAATTGCTGCTGCTT 3’ (Quick Change Multi; single primer); ¹⁷²EDGE¹⁷⁵-QNGQ  +  Y¹⁷⁷A+T⁴²³A;  5' GGGGTGAGAACTGTGCAGAATGAGGAGTGCTACATTGAG 3' (forward); 5’ CTGAATGAGGAATGAGGCAATTGCTGCTGCTT 3’ (reverse). The integrity of all receptor clones was confirmed by cycle-sequencing with the ABI Prism BigDye Terminator
v3.1 ready reaction cycle sequencing kit with reactions analysed on an ABI Prism 373xI 96 capillary automated DNA sequencer (Australian Genome Research Facility, Parkville, Australia).

**Transfections and cell culture**

Wild-type and mutant receptors were isogenically integrated into CHO-FlpIn cells (Invitrogen) as follows: 75 cm$^2$ flasks with CHO FlpIn cells at 70-75% confluency were transfected in serum and antibiotic-free DMEM with 1 µg of pEFS/FRT/V5-DEST vector containing the wild type or mutant $M_2$ mAChR gene and 9 µg POG44 vector (containing Flp recombinase) using lipofectamine (75 µl/75 cm$^2$ flask) according to the manufacturers recommendations. Selection of cells expressing the receptors was achieved by treatment with 400 µg/ml hygromycin-B every two days until resistant flocules were obtained, prior to passaging a further five times. The cells were characterized for receptor expression by radioligand binding assay (see below). Transfected and non-transfected CHO-FlpIn cells were grown and maintained in DMEM containing 20 mM HEPES, 10% fetal bovine serum, 50 U/mL penicillin-streptomycin and 200 µg/mL Hygromycin-B (transfected CHO-FlpIn cells only) at 37°C in a humidified incubator containing 5% CO$_2$, 95% O$_2$.

**$[^3H]$NMS radioligand binding assay membrane preparation**

When cells were approximately 90% confluent, they were harvested using trypsinization and centrifuged (300 g, 3 minutes). The pellet was then resuspended in HEPES homogenization buffer (50 mM HEPES, 2.5 mM MgCl$_2$, 2 mM EGTA) and the centrifugation procedure repeated. The intact cell pellet was resuspended in HEPES homogenization buffer and homogenized using a Polytron homogenizer for two 10-second intervals at maximum setting, with 30 second cooling periods on ice between each burst. The homogenate was centrifuged (1000 g, 10 min, 25°C), the pellet discarded and the supernatant recentrifuged (30,000 g, 30 min, 4°C). The resulting pellet
was resuspended in 5 mL HEPES buffer (110 mM NaCl, 5.4 mM KCl, 1.8 mM CaCl₂, 10 mM MgSO₄, 25 mM glucose, 50 mM HEPES, 58 mM sucrose; pH 7.4) and the protein content determined using the method of Bradford (1976). The homogenate was then divided into 1 mL aliquots and stored frozen at -80°C until required for radioligand binding assays.

[^35S]GTP\gamma S assay membrane preparation

When cells were approximately 90% confluent, they were harvested using lifting buffer (10 mM HEPES, 0.9% NaCl, 0.2% EDTA, pH 7.4 at room temperature) and centrifuged (300 g, 3 minutes). The pellet was then resuspended in Buffer A (10 mM HEPES, 10 mM EDTA; pH 7.4 at 4°C) and homogenized as described above. The homogenate was centrifuged (5 000 g, 10 minutes, 4°C), the pellet discarded and the supernatant recentrifuged (30 000 g, 30 minutes, 4°C). The pellet was resuspended in Buffer B (10 mM HEPES, 0.01 mM EDTA, pH 7.4 at 4°C) and centrifuged (30 000 g, 30 minutes, 4°C). The resulting pellet was resuspended in 3 mL of Buffer B and the protein content determined and aliquots stored as described above.

[^3H]NMS inhibition or potentiation binding assays

Membrane homogenates (25 µg) were incubated in a 1 mL total volume of Tris buffer (50 mM Tris Base, 3 mM MgCl₂, 0.2 mM EGTA; pH 7.4) containing[^3H]NMS (0.5 nM) and a range of concentrations of gallamine, alcuronium, C₇/₃-phth, acetylcholine, McN-A-343, AC-42 or 77-LH-28-1 at 37°C for 60 min (agonists) or 90 mins (prototypical allosteric modulators), unless otherwise indicated in the Results. Non-specific binding was defined using 10 µM atropine. Incubation was terminated by rapid filtration through Whatman GF/B filters using a Brandell cell harvester (Gaithersburg, MD). Filters were washed three times with 3 mL aliquots of ice-cold 0.9% NaCl buffer and dried before the addition of 4 mL of scintillation cocktail (Ultima Gold;
Packard Bioscience, Meriden, CT). Vials were then left to stand until the filters became uniformly translucent before radioactivity was determined using scintillation counting.

**[^3H]NMS dissociation kinetic assay**

CHO-FlpIn cell membranes (25 µg) were equilibrated with[^3H]NMS (0.5 nM) in a 1 mL total volume of Tris buffer (also containing 100 µM Gpp(NH)p for the McN-A-343 experiments) for 60 minutes at 37°C. Atropine (10 µM) alone or in the presence of test (allosteric) ligand was then added at various time points to prevent the reassociation of[^3H]NMS with the receptor. In subsequent experiments designed to investigate the effect of a range of modulator concentrations on[^3H]NMS dissociation rate, a “two-point kinetic” experimental paradigm was used, where the effect of increasing concentrations of allosteric modulator on[^3H]NMS dissociation was determined at 0 and 6 minutes. This approach is valid to determine[^3H]NMS dissociation rate constants if the full time course of radioligand dissociation is monophasic both in the absence and presence of modulator (Kostenis and Mohr, 1996; Lazareno and Birdsall, 1995); this was the case in our current study. The assays involved initial equilibration of CHO-FlpIn cell membranes (125 µg/mL) with[^3H]NMS (2.5 nM) in the presence or absence of atropine (100 µM) at 37°C for 1 hr. For determination of radioligand dissociation at 6 minutes, equilibrated membrane/radioligand (100 µL) was added to Tris buffer containing atropine (100 µM) alone or in the presence of increasing concentrations of allosteric modulator to a 500 µL final volume. To determine radioligand binding at 0 minutes and non-specific radioligand binding, 100 µL of membrane/radioligand equilibrated in the presence of absence of atropine was added to tubes alone. Termination of the reaction and determination of radioactivity were performed as described above.
**[^3]H\textit{NMS} pseudo-equilibrium binding assay**

Two sets of tubes containing Tris buffer in the presence of McN-A-343 (0.3 nM – 3 mM) or atropine (3 pM – 10 µM), in the absence or presence of 100 µM 5′-guanylylimidodiphosphate (Gpp(NH)p), were prepared to a 1 mL total volume. In each case, the first set of tubes was treated as per normal for standard radioligand binding assays, i.e., the reactants were added together and allowed to approach equilibrium. The second set of tubes was treated differently, whereby the orthosteric radioligand and receptor were pre-equilibrated at a high concentration, prior to dilution and exposure to allosteric ligand. Specifically, to the first set of tubes 10 µL of both[^3]H\textit{NMS} (20 nM) and membrane (2500 µg/mL) were added separately (100-fold dilution). For the second set of tubes[^3]H\textit{NMS} (20 nM) and membrane (2500 µg/mL), each representing 100 x the desired final concentration, were first combined in a 1:1 ratio (thus reducing the concentration to 50 x the desired final concentration), equilibrated for 30 mins at 37° at which point 20 µL of the mixture was distributed to each tube, with the final result being a 50-fold dilution to the final desired concentration. Both sets of tubes were then incubated for 20 mins at 37°C. Determination of non-specific binding, termination of the reaction, and determination of radioactivity were performed as described above.

**[^3]H\textit{GTP}γS binding assay**

Membrane homogenates (15 µg) were equilibrated in a 900 µL total volume of[^35]S\textit{GTP}γS assay buffer (10 mM HEPES, 100 mM NaCl, 10 mM MgCl₂; pH 7.4 at 30°C) containing 10 µM guanosine 5′-diphosphate (GDP) and a range of concentrations of acetylcholine (0.3 nM – 100 µM), pilocarpine (3 nM – 300 µM) or McN-A-343 (3 nM – 300 µM) at 30°C for 30 minutes. After this time, 100 µL of[^35]S\textit{GTP}γS (100 pM) was added and incubation continued for another
30 minutes at 30°C. Termination of reaction and determination of radioactivity were performed as described above

**ERK1/2 phosphorylation assay**

Cells were seeded into 96-well plates at a density of 50 000 cells/well. After 4 hours, cells were washed twice with PBS and maintained in DMEM containing 20 mM HEPES and 50 U/mL penicillin-streptomycin for at least 4 hours before assaying. Assays investigating the time course of action and concentration-response curves were generated by the addition of ligand for the indicated time periods (200 µL final volume) at 37°C. The time of stimulation for concentration-response curves represents the time of peak response as determined in time course assays. Agonist stimulation of cells (5 mins unless otherwise specified) was terminated by the removal of media and the addition of 100 µL of *SureFire*™ lysis buffer to each well. The plate was then agitated for 1-2 minutes. A 4:1 v/v dilution of Lysate:*SureFire*™ activation buffer was made in a total volume of 50 µL. A 1:100:120 v/v dilution of AlphaScreen™ beads:activated lysate mixture:*SureFire*™ reaction buffer in a 11 µL total volume was then transferred to a white opaque 384-well Proxiplate™ in diminished light. This plate was then incubated in the dark at 37°C for 1.5 hours after which time the fluorescence signal was measured by a Fusion-α™ plate reader (PerkinElmer), using standard AlphaScreen™ settings. All data was expressed as a percentage of the ERK1/2 phosphorylation mediated after a 6 minutes exposure to DMEM containing 3% FBS.
Data analysis

Data sets of total and non-specific binding obtained from each [³H]NMS saturation binding assay were globally fitted to the following equation using Prism 4.03 (GraphPad Software, San Diego, CA):

\[
Y = \frac{B_{\text{max}} \cdot [A]}{[A]+K_A} + NS \cdot [A]
\]

where \(Y\) represents radioligand binding, \([A]\) denotes the concentration of radioligand, \(B_{\text{max}}\) denotes the maximal density of binding sites, \(K_A\) is the radioligand equilibrium dissociation constant, and \(NS\) is the fraction of non-specific binding. The hyperbolic term in this equation was not used when fitting the non-specific binding data, whereas the parameter, \(NS\), was shared between both total and non-specific binding data sets (Motulsky and Christopoulos, 2004).

Agonist inhibition binding data were empirically fitted to either a one-site (equation 2) or two-site/state (equation 3) inhibition mass action curve using Prism 4.03:

\[
Y = \text{Bottom} + \frac{\text{Top-Bottom}}{1+10^{X-\log IC_{50}}}
\]

where \(\text{Top}\) is the specific binding of the radioligand in the absence of any competing ligand, \(\text{Bottom}\) is the specific binding of the radioligand equal to non-specific binding, \(IC_{50}\) is the concentration of competing ligand that produces radioligand binding half way between the \(\text{Top}\) and \(\text{Bottom}\), and \(X\) is the logarithm of the concentration of the competing ligand.

\[
Y = \text{Bottom} + \left(\text{Top-Bottom}\right) \left(\frac{F_H}{1+10^{X-\log IC_{50}}} + \frac{1-F_H}{1+10^{X-\log IC_{50}}}ight)
\]
Y, Top, Bottom and X are as above for equation 2; FH denotes the fraction of receptors inhibiting radioligand binding with a potency described by IC_{50,1} while IC_{50,2} represents the inhibitory potency of the remaining fraction of receptors. In all instances, an extra-sum-of-squares (F-test) was used to determine whether the data were better described by a one vs a two-site model.

Dissociation kinetic data all followed a monoexponential decay, and were thus fitted to the following equation using Prism 4.03:

\[ B_t = B_0 \cdot e^{-k_{off} \times t} \] (4)

where \( t \) denotes incubation time, \( B_t \) denotes specific radioligand binding at time \( t \), \( B_0 \) denotes the specific radioligand binding at time at equilibrium (time = 0) and \( k_{off} \) represents the observed radioligand dissociation rate constant. For the two-point dissociation experiments, where the effects of a range of concentrations of allosteric modulators were investigated, individual \( k_{off} \) values determined in the presence of modulator were normalized to the control \( k_{off} \) value (absence of modulator) and then plotted as a function of modulator concentration. The EC\(_{50}\) value from the fit of a 3-parameter logistic equation to these data represents the ratio, \( K_B/\alpha \), in the ATCM (Lazareno and Birdsall, 1995). In the case of the prototypical modulators gallamine, C\(_{7/3}\)-phth and alcuronium, these two-point kinetic concentration response curves were globally (simultaneously) fitted with the corresponding pseudo-equilibrium binding curves (see below), with the parameters \( K_B \) and \( \alpha \) shared between the datasets.

Pseudo-equilibrium binding data were analyzed by Prism 4.03 according to a kinetic ATCM, (Avlani et al., 2004; Lanzafame et al., 2006; Lazareno and Birdsall, 1995). This model explicitly incorporates additional kinetic parameters, such as the dissociation rate constant of the
radioligand in the absence and presence of allosteric modulator, and fits the binding as a function of incubation time.

\[ B_t = B_{AB} \left[ 1 - e^{-t \cdot k_{\text{onobs}}} \right] + B_{HI} \cdot e^{-t \cdot k_{\text{onobs}}} \]  \hspace{1cm} (5)

where

\[ k_{\text{onobs}} = k_{\text{offobs}} \cdot \left(1 + \frac{[A]}{K_{\text{APP}}} \right) \]  \hspace{1cm} (6)

\[ k_{\text{offobs}} = \frac{k_{\text{off}} + [B] \cdot k_{\text{offB}}}{1 + \left( \frac{[B]}{K_B} \right) / \alpha} \]  \hspace{1cm} (7)

\[ B_{AB} = \frac{[R] \cdot [A]}{1 + [A] / K_{\text{APP}}} \]  \hspace{1cm} (8)

\[ B_{HI} = \frac{[R]_t \cdot [A] \cdot 50 \cdot \frac{1}{K_A}}{1 + [A] \cdot 50 \cdot \frac{1}{K_A}} \]  \hspace{1cm} (9)

In these equations, \( B_t \) and \( t \) are as previously defined, \([A]\) is the concentration of radioligand, \( k_{\text{onobs}} \) and \( k_{\text{offobs}} \) are the apparent radioligand association and dissociation rate constants in the presence of allosteric modulator respectively and \( k_{\text{off}} \) and \( k_{\text{offB}} \) denote the radioligand dissociation rate constants for the modulator-unoccupied and modulator-occupied receptor, respectively. \([R]_t\) is the total concentration of receptors, \( B_{AB} \) denotes bound radioligand to both free and occupied receptor and \( B_{HI} \) denotes the level of bound radioligand in the pre-equilibrated tubes. For tubes that are not pre-equilibrated, \( B_{HI} \) is equal to zero. \( K_{\text{APP}}, K_A, \alpha \) and \([B]\) are as defined above. It is important to note that all equations (5 - 9) based on the ATCM assume that the measured effects
of the allosteric ligand (i.e., modulation effects and any agonistic properties) arise via binding to a single (allosteric) site; if the ligand actually displays two different modes of binding, e.g., interacting with both the orthosteric and allosteric sites, then the ATCM will not yield correct estimates of the equilibrium binding properties of the modulator (May et al., 2007).

Agonist concentration-response curves were fitted to the following 3-parameter-logistic equation using Prism 4.03.

\[
\text{Response} = \text{Bottom} + \frac{(\text{Top} - \text{Bottom})}{1 + 10^{(\log \text{EC}_{50} - \log [A])}}
\]

(10)

where Bottom and Top are the lower and upper plateaus, respectively, of the concentration-response curve, \([A]\) is the molar concentration of agonist and \(\text{EC}_{50}\) is the molar concentration of agonist required to generate a response half-way between the Top and Bottom.

For combination studies, the interaction between ACh and each of the partial allosteric agonists displayed high negative cooperativity (see Results) and therefore was indistinguishable from a competitive interaction. Consequently, ACh concentration response curves in the absence and presence of the partial agonists, McN-A-343, AC-42 and 77-LH-28-1, were adequately fitted to the following operational model for the competitive interaction between an orthosteric full agonist and orthosteric partial agonist, as derived previously by Leff et al. (1993).

\[
E = E_m \left( \frac{[A][K_B + n[EC_{50}]^n]}{[EC_{50}]^n(K_B + [B])^n + ([A][K_B + n[EC_{50}]^n])} \right)
\]

(11)

where \(E\) is the pharmacological effect, \(E_m\) is the maximal possible response, \(\text{EC}_{50}\) is the molar concentration of the orthosteric full agonist (A) required to achieve half maximal response, \(n\) is
the Hill slope of the orthosteric full agonist concentration-response curve, $K_B$ and $\tau$ represent the equilibrium dissociation constant and the operational index of efficacy of the partial agonist (B) respectively. This analysis assumes that a) the values of $E_m$ and $n$ derived from the orthosteric full agonist concentration-response curve are approximate estimates of system maximal responsiveness and transducer function slope, respectively, and that b) both agonists utilize the same transduction machinery and as such the Hill slope can be shared across the full and partial agonist concentration-response curves.

All affinity, potency, efficacy and cooperativity parameters were estimated as logarithms (Christopoulos, 1998). In all instances, results are expressed as mean ± standard error of the mean. Statistical analyses were performed by a paired $t$ test or by F-test, as appropriate, using Prism 4.03 and statistical significance was taken as $p < 0.05$. 
RESULTS

Characterization of the binding properties of [³H]NMS and prototypical allosteric modulators at wild type and mutant M₂ mAChRs

Previous studies indicating a role for the M₂ mAChR residues, ¹⁷²EDGE¹⁷⁵, Y¹⁷⁷ and T⁴²³, in allosteric modulator binding generally utilized a substitution approach, whereby these amino acids in the M₂ mAChR (highest affinity for prototypical modulators) were swapped to corresponding amino acids in the M₅ mAChR (lowest affinity for prototypical modulators) in order to probe the determinants of modulator subtype-selectivity (see Introduction and references therein). In the current study, we created a different set of mutations in these regions. Specifically, we chose to investigate the impact of neutralizing the charge of the EDGE sequence while maintaining the side chain structure, i.e., QNGQ (“M₂-EDGE” mutant), replacing Y¹⁷⁷ and T⁴²³ with alanine (“M₂-YT” mutant) and a combination of the EDGE-QNGQ and Y¹⁷⁷A, T⁴²³A mutations (“M₂-EDGE-YT” mutant). As expected, saturation binding assays using the orthosteric antagonist, [³H]NMS, found no significant effects of any of the mutations on [³H]NMS binding affinity relative to the wild type (“M₂-WT”) receptor (Table 1). Some variations were noted in the maximal density of binding sites (although all constructs were well expressed), but this was only statistically significant for the M₂-EDGE mutant (Table 1). In dissociation kinetic experiments, no significant differences were found for in the dissociation rate of [³H]NMS at any of the mutants relative to the wild type (0.29 ± 0.02 min⁻¹; n = 3). Thus, it can be concluded that the mutations did not significantly perturb the conformation of the orthosteric binding pocket, nor the access to or egress from that pocket by the radioligand.
In contrast, Figure 2 illustrates the dramatic effects of the mutations on the ability of the prototypical modulators, alcuronium, gallamine and C7/3-phth, to interact with [3H]NMS in pseudo-equilibrium binding assays. Also shown are the effects of the modulators on [3H]NMS dissociation kinetics. Global analysis of both groups of data to a kinetic ATCM yielded the parameters listed in Table 1. Removing the charge of the EDGE sequence while maintaining the essential side chain structure significantly reduced the affinity of the allosteric enhancer, alcuronium, as well as that of the allosteric inhibitors, gallamine and C7/3-phth. The M2-YT mutant had an even more profound inhibitory effect on modulator affinity. Not surprisingly, the combination of both sets of mutations had the strongest inhibitory effect on modulator affinity. In some instances, the cooperativity between each modulator and [3H]NMS was also significantly changed (Table 1). The combined effect of reductions in modulator affinity and, in some cases cooperativity, is manifested in the dissociation kinetic assays, where the potency of the compounds to allosterically retard [3H]NMS dissociation is reduced between 10 – 150 fold (Figure 2). Despite the reduced potency, however, all modulators were still able to completely prevent [3H]NMS dissociation at high concentrations (Figure 2).

Effects of orthosteric and allosteric agonists on [3H]NMS inhibition binding at the M2-WT and M2-EDGE-YT mAChRs

Given that the combined M2-EDGE-YT mutation yielded the most profound reduction in the affinity of prototypical allosteric modulators, this receptor was chosen for comparison with the M2-WT to investigate the effects on agonist binding. As shown in Figure 3, the orthosteric agonist, ACh, as well as McN-A-343, AC-42 and 77-LH-28-1 all caused full inhibition of [3H]NMS (0.5 nM) specific binding at both the M2-WT and M2-EDGE-YT mAChRs. These
experiments were performed in the absence of any guanine nucleotides, and as such it was expected that they could be influenced by the G protein-coupling status of the receptor. At both the M2-WT and M2-EDGE-YT, nonlinear regression of the ACh and McN-A-343 inhibition curves according to empirical binding models yielded Hill slopes that were significantly less then one and therefore preferentially fitted to an empirical two-site binding model (Table 2). Global nonlinear regression analysis of the ACh curves in conjunction with an extra-sum-of-squares test ($F$-test) indicated that these curves were preferentially described by a single shared value for the fraction of receptors exhibiting high affinity binding. At the M2-EDGE+YT, ACh had a reduced apparent dissociation constant for the high affinity, presumably G protein-coupled, form of the receptor; a similar trend was noted for the low affinity binding site, but this was not statistically significant (Table 2). These results suggest that mutations within the common allosteric site may actually have a small inhibitory effect on orthosteric agonist binding.

In contrast, global nonlinear regression analysis of the McN-A-343 binding curves in conjunction with an $F$-test indicated that the data were best described by sharing both the high and low affinity dissociation constants of the two sites, but not the fraction of receptors exhibiting high affinity, which was found to be significantly increased at the M2-EDGE+YT mAChR (Table 2). For AC-42, the inhibition of $[^3]H$NMS specific binding at both the wild type and mutant receptor yielded Hill slopes that were not significantly different from one, but the apparent equilibrium dissociation constant of AC-42 was modestly, but significantly ($p < 0.05$), increased at the M2-EDGE+YT mAChR (Table 2). Interestingly, the inhibition of $[^3]H$NMS binding by 77-LH-28-1 preferentially fitted to a one-site model at the M2-WT but a two-site binding model at the M2-EDGE+YT mAChR (Table 2). Collectively, these results suggest that, in the absence of guanine
nucleotides, the affinity of ACh appears to be slightly reduced, whereas McN-A-343, AC-42 and 77-LH-28-1 trend either towards an increased affinity or increased fraction of receptors exhibiting high affinity at the M2-EDGE-YT mAChR.

Effects of McN-A-343, AC-42 and 77-LH-28-1 on [³H]NMS dissociation kinetics at the M₂-WT and M₂-EDGE-YT mAChRs

To more directly probe the ability of McN-A-343, AC-42 and 77-LH-28-1 to allosterically modulate orthosteric ligand binding, the effect of these compounds on the rate of orthosteric radioligand dissociation was investigated. These experiments were all performed in the presence of 100 µM Gpp(NH)p, to reduce receptor-G protein coupling and thus simplify the interpretation of the data. At the M₂-WT mAChR, the presence of McN-A-343 (300 µM; 0.17 ± 0.02 min⁻¹) and 77-LH-28-1 (100 µM; 0.15 ± 0.02 min⁻¹), but not AC-42 (100 µM; 0.22 ± 0.02 min⁻¹), significantly (p < 0.05) retarded the dissociation rate of [³H]NMS (Figure 4A). These findings suggest that McN-A-343 and 77-LH-28-1 can bind simultaneously with [³H]NMS to the M₂ mAChR to allosterically alter [³H]NMS dissociation. At the M₂-EDGE-YT mAChR, 77-LH-28-1 (100 µM; 0.13 ± 0.06 min⁻¹ vs 0.23 ± 0.02 min⁻¹ for the wild type) retained the ability to significantly retard [³H]NMS dissociation; a trend was noted for McN-A-343 (0.17 ± 0.02 min⁻¹) as well, but this was not statistically significant at the concentration utilized (Figure 4B). To more rigorously investigate the latter, the entire concentration-response relationship of McN-A-343 inhibition of [³H]NMS dissociation kinetics was determined at both the wild type and mutant mAChRs. At the M₂-WT, McN-A-343 caused virtually complete inhibition of [³H]NMS dissociation (Figure 4C) and the pEC₅₀ was determined as 3.23 ± 0.05 (n = 4). In contrast, at the M₂-EDGE-YT mAChR, McN-A-343 was unable to cause complete inhibition of [³H]NMS dissociation kinetics and only had a small, albeit significant (p < 0.05), reduction in potency.
(pEC$_{50}$ = 2.98 ± 0.05; n = 4; Figure 4C). This is in contrast to the effects on the prototypical modulators (Figure 2), which were characterized by substantial reductions in allosteric potency but not maximal effect.

**Quantification of the allosteric binding properties of McN-A-343 at the M$_2$-WT and M$_2$-EDGE-YT mAChRs**

The potency of McN-A-343 to retard the dissociation of [$^{3}$H]NMS in the kinetic assay reflects the combined effects of both modulator affinity (K$_B$) and cooperativity with the radioligand ($\alpha$). To obtain individual estimates of the parameters, a different type of experiment is required. Due to the high negative cooperativity between [$^{3}$H]NMS and McN-A-343 (Figure 3B), we exploited the ability of McN-A-343 to slow [$^{3}$H]NMS dissociation kinetics and to mediate a kinetic artifact in pseudo-equilibrium binding assays with reduced incubation times (20 mins). These assays generate two curves, with the only variable between them being whether or not incubation was initiated by the addition of ‘free’ (non pre-equilibrated) receptor or radioligand-bound (pre-equilibrated) receptor (see Materials and Methods). To ensure that orthosteric ligands do not mediate similar kinetic artifacts, these assays were initially performed using the classic orthosteric antagonist, atropine. As shown in Figure 5A for the M$_2$-WT mAChR, atropine yielded the expected behavior of a simple competitive orthosteric antagonist, causing complete inhibition of [$^{3}$H]NMS specific binding irrespective of the order of ligand-receptor exposure. Nonlinear regression revealed that both curves preferentially fitted to a monophasic isotherm with a Hill slope not significantly different from one. When the Hill slope was constrained to one, the estimated pK$_B$ values for atropine were 8.54 ± 0.03 and 8.47 ± 0.07 for non pre-equilibrated and pre-equilibrated receptors respectively (n = 3). In contrast, high concentrations
of McN-A-343 (in the presence of 100 µM Gpp(NH)p), caused strikingly divergent effects on the approach of \[^3\text{H}\]NMS to equilibrium at both the M2-WT (Figure 5B) and the M2-EDGE-YT (Figure 5C) mAChRs, clearly indicating a stabilization of a ternary complex between receptor, radioligand and high concentrations of McN-A-343 that leads to either an overestimation (pre-equilibration) or underestimation (no pre equilibration) of the level of specific binding over the short assay time period relative to what would be obtained if the system were at equilibrium. A kinetic ATCM, that accounts for additional parameters such as the incubation time and the dissociation rates of \[^3\text{H}\]NMS from the modulator unoccupied and occupied receptor, was globally fitted to the curves to derive values of pK$_B$ and Log$\alpha$. In comparison to the M2 wild type mAChR, the affinity of McN-A-343 for the M2-EDGE-YT mAChR had significantly increased negative cooperativity (Log $\alpha$ of -1.40 ± 0.14 and -2.37 ± 0.05 for the M2 WT and M2-EDGE-YT mAChRs respectively; n = 3) and a trend towards increased affinity (pK$_B$ of 4.70 ± 0.17 and 5.10 ± 0.01 for the M2-WT and M2-EDGE-YT mAChRs respectively; n = 3).

**Effects of allosteric site mutations on agonist functional properties**

For initial functional experiments, the time course of ERK1/2 phosphorylation in response to orthosteric and allosteric agonists was determined in non-transfected, M2-WT and M2-EDGE-YT CHO-FlpIn cells. In addition to ACh, the partial agonist, pilocarpine, was included in these studies as a second well-characterized orthosteric agonist. None of the ligands tested mediated a significant level of ERK1/2 phosphorylation in non-transfected CHO-FlpIn cells at any time points measured (data not shown). At both the M2-WT and M2-EDGE+YT mAChRs, ACh and pilocarpine mediated a robust stimulation, whereas AC-42 and 77-LH-28-1 mediated a very modest stimulation, of ERK1/2 phosphorylation (Figure 6). Interestingly, the level of ERK1/2
phosphorylation mediated by McN-A-343 was markedly different at the M2-WT and M2-EDGE-YT mAChRs, with McN-A-343 causing only modest stimulation at M2-WT mAChRs (Figure 6A) but equivalent stimulation at the M2-EDGE-YT mAChR to that obtained by ACh (Figure 6B).

The time to the peak ERK1/2 phosphorylation was then chosen as the stimulation period for subsequent experiments aimed at characterizing orthosteric and allosteric concentration-response profiles at M2-WT and M2-EDGE-YT mAChRs. The orthosteric agonists, ACh and pilocarpine, mediated a robust stimulation of ERK1/2 phosphorylation at both M2-WT and M2-EDGE-YT mAChRs (Figure 7A, 7B). Estimates of ACh potency and maximal response were not significantly different between the two receptors (Table 3). However, the maximal response elicited by pilocarpine was modestly but significantly (p < 0.05) increased at the M2-EDGE-YT mAChR (Table 3); this likely reflects the slightly higher expression of the M2-EDGE-YT mAChR as compared to the M2-WT receptor (Table 1). At the M2-WT receptor, McN-A-343 and 77-LH-28-1 were weak partial agonists with respect to stimulation of ERK1/2 phosphorylation whereas AC-42 had minimal activity (Figure 7A; Table 3). Interestingly, at the M2-EDGE-YT mAChR, the efficacy of McN-A-343 was profoundly and significantly increased such that at this receptor it behaved as a full agonist (Figure 7B; Table 3). Given the modest effect noted on pilocarpine maximal response, the substantial increase in the efficacy of McN-A-343 is unlikely to simply reflect increased receptor expression. At the M2-EDGE-YT mAChR, AC-42 maintained minimal activity whereas 77-LH-28-1 had a significantly increased potency (Figure 7B; Table 3); since the latter compound remained a partial agonist, this increase in potency suggests an increase in affinity of 77-LH-28-1 at the mutant receptor. Thus, the functional results
support the binding data in suggesting that mutations within the common allosteric site increase the affinity or efficacy of the allosteric agonists McN-A-343 and 77-LH-28-1.

To ensure that the functional effects of the receptor mutations were not influenced by any potential non-equilibrium artifacts associated with the ERK1/2 phosphorylation assay, a second assay measuring ligand mediated [$^{35}$S]GTPγS binding to activated G proteins was performed with the ligands pre-equilibrated prior to addition of the radiolabel. In this assay, ACh was a full agonist and pilocarpine a partial agonist at both the M2-WT and M2-EDGE+YT mAChRs (Figure 7C, 7D). Similar to the ERK1/2 phosphorylation assays, McN-A-343 caused partial stimulation of [$^{35}$S]GTPγS binding at the M2-WT mAChR but was practically indistinguishable from a full agonist at the M2-EDGE-YT mAChR (Figure 7C, 7D).

Finally, to determine which epitopes are primarily involved in mediating the effects of the combined M2-EDGE-YT mutations, ERK1/2 phosphorylation assays were repeated using M2-EDGE and M2-YT mAChRs. At both receptors, ACh mediated a robust stimulation of ERK1/2 phosphorylation with no significant difference in potency or efficacy when compared to the M2-WT mAChR (Figure 8 and Table 3). At the M2-EDGE mAChR, the potency and efficacy of McN-A-343, AC-42 and 77-LH-28-1 were not significantly different from the M2-WT mAChR (Figure 8A and Table 3). However, the efficacy of McN-A-343 and the potency of 77-LH-28-1 were significantly increased at the M2-YT mAChR, similar to the M2-EDGE-YT mAChR (Figure 8B and Table 3), suggesting that it is the two alanine substitutions that account for the majority of the effect. AC-42 had minimal activity at both the M2-EDGE and M2-YT mAChRs (Figure 8).
Functional interaction studies between ACh and allosteric agonists at the M2-WT and M2-EDGE-YT mAChRs

In addition to gaining insight into the nature of the allosteric interaction, investigating the effect of allosteric modulators on orthosteric agonists can provide estimates of modulator affinity and, under appropriate conditions, cooperativity. McN-A-343, AC-42 and 77-LH-28-1 all mediated a parallel rightward shift in ACh mediated-ERK1/2 phosphorylation with no depression in the maximal response at either the M2-WT (Figure 9) or the M2-EDGE-YT mAChR (Figure 10; AC-42 and 77-LH-28-1 only); the interaction between McN-A-343 and ACh at the M2-EDGE-YT mAChR was not investigated due to McN-A-343 acting as a full agonist at this receptor. Global fitting of the data sets to an operational model of competitive antagonism yielded Schild slopes that were not significantly different from 1 (Table 4), indicating that the interactions were characterized by very high negative cooperativity such that they were indistinguishable from a competitive (orthosteric) interaction. Therefore the data were re-fitted to the operational model with the Schild slope constrained to unity. The resulting pK_B estimates (Table 4) show that 77-LH-28-1 had a significantly increased affinity for the M2-EDGE+YT mAChR, which agrees with the increased potency observed for 77-LH-28-1 in ERK1/2 phosphorylation concentration-response assays at the same mutant.
DISCUSSION

We have found that the agonists, McN-A-343 and the novel 77-LH-28-1, have an allosteric mechanism of action at the human M\textsubscript{2} mAChR. A similar mechanism of action of AC-42 at the human M\textsubscript{2} mAChR could not be confirmed. Although we cannot yet conclude whether these agonists mediate all their effects via the allosteric site, or whether they interact with both orthosteric and allosteric sites, the unique sensitivity of each of the compounds investigated to mutations within the common allosteric site on the M\textsubscript{2} mAChR strongly suggests that all these agonists have a different mode of binding in comparison to orthosteric ligands, such as ACh and pilocarpine, as well as prototypical mAChR modulators, such as gallamine, alcuronium and C\textsubscript{7}/3-phth. The enhanced activities of McN-A-343 and 77-LH-28-1 at mutant mAChRs that display reduced potencies for prototypical modulators was mediated mainly by the “common allosteric site” residues, Y\textsubscript{177} and T\textsubscript{423}, rather than 172EDGE\textsuperscript{175}. To our knowledge, this is the first study investigating the effects of allosteric-site mutations on allosteric agonists of a Family A GPCR.

Our finding that removal of the charge on 172EDGE\textsuperscript{175} and substitution of Y\textsuperscript{177} and T\textsuperscript{423} with alanine led to a reduction in the potency of prototypical allosteric ligands (Figure 2) is in agreement with previous studies (Buller et al., 2002; Gnagey et al., 1999; Huang et al., 2005; Leppik et al., 1994; Voigtlander et al., 2003), and confirms that these residues are vital to the binding of modulators structurally related to classic neuromuscular blockers and alkane-bis-onium compounds (Birdsall and Lazareno, 2005). In contrast, binding assays revealed no reduction in the potency of McN-A-343, AC-42 and 77-LH-28-1 to mediate complete inhibition of [\textsuperscript{3}H]NMS binding at both the M\textsubscript{2}-WT and M\textsubscript{2}-EDGE-YT mAChRs (Figure 3). This finding
suggested either that the agonists interacted competitively with \[^3\text{H}\]NMS, or that they interacted allosterically but with very high negative cooperativity. Previous studies of AC-42 and related compounds at M\(_1\) mAChRs found similar evidence for high negative cooperativity with \[^3\text{H}\]NMS (Langmead et al., 2006; Spalding et al., 2002; 2006), as well as differential effects of orthosteric site mutations on these compounds when compared to the orthosteric agonist, carbachol (Spalding et al., 2002; 2006; Sur et al., 2003). In the current study, dissociation kinetic experiments provided conclusive evidence for an allosteric interaction at the human M\(_2\) mAChR by McN-A-343 and 77-LH-28-1, but highlighted significant differences between the effects of allosteric-site mutations on allosteric agonist versus modulator binding. Specifically, mutation of the key EDGE-YT residues in the common allosteric site caused approx. 10 – 150 fold reductions in the potency of prototypical modulators to retard \[^3\text{H}\]NMS dissociation (Figure 3), but did not affect the ability of the modulators to completely retard radioligand dissociation. In contrast, the same mutations caused only a modest reduction in the potency of McN-A-343, (approximately 1.8 fold), but led to a significant reduction in its maximal effect on radioligand dissociation rate (Figure 4C).

In addition to the dissociation kinetic assays, our pseudoequilibrium binding assays also provided evidence for an allosteric mode of interaction by McN-A-343, as well as yielding estimates of the affinity of McN-A-343 for the allosteric site and its cooperativity with \[^3\text{H}\]NMS. With the caveat that the pK\(_B\) and log\(\alpha\) values obtained from these assays are derived from a model that assumes McN-A-343 does not interact with the orthosteric site, the resulting estimates suggested that whilst there was no significant difference in the affinity of McN-A-343 for the mutant M\(_2\) mAChR relative to the wild type, the modulator had significantly higher negative cooperativity
with \[^3\text{H}\]NMS at the M\(_2\)-EDGE-YT mAChR, consistent with a change in the mode of McN-A-343 binding. Collectively, the dissociation kinetic and pseudoequilibrium binding studies suggest that the \(^{172}\text{EDGE}^{175}\text{-QNGQ + Y}^{177}\text{A + T}^{423}\text{A}\) mutations alter the binding of McN-A-343 and 77-LH-28-1 to the receptor in a manner that is different from classic (non agonistic) modulators. It is likely that the allosteric agonists utilize different epitopes on the M\(_2\) mAChR to the prototypical modulators, but it remains to be determined if their binding site(s) overlap at all with the common allosteric site or whether the effects determined in the current study are mediated via indirect conformational changes transmitted between distinct allosteric sites.

Another interesting finding from the initial \[^3\text{H}\]NMS inhibition binding assays conducted in the absence of Gpp(NH)p, was the effect of the combined allosteric-site mutation, EDGE+YT, on agonist affinity estimates. ACh, like many full agonists, displayed two apparent dissociation constants when competing with \[^3\text{H}\]NMS; the effect of the allosteric-site mutations was to reduce the potency of ACh somewhat, but McN-A-343, 77-LH-28-1 and AC-42 all trended towards either a higher affinity or an increase in the proportion of receptors demonstrating high affinity binding. At the moment, it remains difficult to interpret the mechanistic basis of multiple agonist affinity states in competition binding studies, although in most instances it is assumed to reflect the G protein-coupling status to some extent (Christopoulos and El-Fakahany, 1999). If so, the findings from these types of binding assay indicated that mutation of the common allosteric site on the M\(_2\) mAChR may actually increase the efficacy and/or potency of allosteric agonists, a hypothesis that could be directly tested in functional assays.
In experiments measuring M2 mAChR-mediated ERK1/2 phosphorylation, both ACh and pilocarpine mediated a robust response, with potencies that were not significantly different between the M2-WT and M2-EDGE-YT receptors. Different results were obtained with McN-A-343, 77-LH-28-1 and AC-42, however, supporting the hypothesis that the binding mode of these agonists is unlikely to be the same as that of classic orthosteric ligands. The finding that McN-A-343, 77-LH-28-1 and AC-42 mediated minimal, or no, stimulation of ERK1/2 phosphorylation at the M2-WT mAChR is consistent with the high functional selectivity of these compounds for the M1 mAChR (Mitchelson, 1988; Spalding et al., 2002; Langmead et al., manuscript in preparation). At the M2-EDGE-YT mAChR, the trend towards a modestly increased maximal response observed for pilocarpine, AC-42 and 77-LH-28-1, is likely to be a consequence of the slightly increased expression of the M2-EDGE-YT mAChR (Table 1) as opposed to an enhanced intrinsic efficacy of each of the ligands. The most significant changes observed at the M2-EDGE-YT mAChR were a greater than 5 fold increase in the maximal response (efficacy) of McN-A-343, such that at the mutant mAChR it was a full agonist, and a greater than 10 fold increase in the potency of 77-LH-28-1; since the latter remained a partial agonist, the change in its potency must reflect a significant increase in its affinity for the M2-EDGE-YT. These effects are qualitatively consistent with effects noted in the binding assays, and are not assay-specific, since a similar profile was noted for McN-A-343 relative to ACh and pilocarpine in assays of [35S]GTPγS binding to activated G proteins. Together, these results suggest that the conformation of the M2-EDGE-YT mAChR, while having minimal effects on the function of orthosteric agonists, can have a profound effect on the efficacy (McN-A-343) or affinity (77-LH-28-1) of allosteric agonists. They also suggest that differences may exist between the binding/activation modes of McN-A-343, on the one hand, and 77-LH-28-1, on the other.
Experiments focusing on additional mutants, namely M2-EDGE\textsuperscript{175} - versus M2-YT, found that it was the latter alanine substitutions that likely mediate the bulk of the observed effects on the allosteric agonists (Figure 8).

Given that McN-A-343 and 77-LH-28-1 had an allosteric mechanism of action, it was also important to investigate the propensity for functional interaction between these compounds and the endogenous ligand, ACh. In all instances, McN-A-343, 77-LH-28-1 and AC-42 displayed high negative cooperativity ($\alpha$ approaches 0 in the ATCM) with ACh at the M2-WT, and as such were fitted to a competitive (orthosteric) model describing the interaction of a partial agonist against a full agonist (Leff et al., 1993); this model is equivalent to the ATCM when $\alpha = 0$. As expected, the affinity of 77-LH-28-1 as an antagonist of ACh-mediated responses was significantly increased at the M2-EDGE-YT mAChR. Interestingly, the affinity of AC-42 was only slightly increased, suggesting perhaps subtle differences in its binding mode relative to 77-LH-28-1. Unfortunately, the full agonism of McN-A-343 at the mutant receptor precluded the use of this agonist in similar interaction studies.

In conclusion, we have shown that McN-A-343 and 77-LH-28-1 exhibit both allosteric modulation and agonism at the human M2 mAChR. In addition, we identified strikingly different effects that 172\textsuperscript{EDGE}-QNGQ, $Y^{177}A$ and $T^{423}A$ allosteric-site mutations have on allosteric agonist compared to prototypical M2 mAChR modulators, such as gallamine, alcuronium and C7/3-phth. The growing list of allosteric modulators that possess intrinsic efficacy in their own right suggests that GPCRs may be activated from different regions within the receptor in addition to the orthosteric binding site. Given the potentially greater sequence diversity for receptor
regions located outside the endogenous ligand-binding site, such allosteric agonists represent a new and exciting avenue for therapeutic intervention.
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FOOTNOTE

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FIGURE LEGENDS

Figure 1 (A) Structures of the allosteric mAChR agonists used in this study. (B) Snake diagram of the M₂ mAChR, indicating amino acids previously reported to contribute to orthosteric ligand binding (grey) and prototypical allosteric modulator binding (black). Amino acids highlighted by arrows indicate those mutated in our current study.

Figure 2 Mutation of residues in the second extracellular loop/top of TM7 reduces the binding of prototypical M₂ mAChR allosteric modulators. The interaction between each prototypical modulator and the orthosteric ligand, [³H]NMS (0.5 nM), was assessed at the M₂-WT and each of the indicated M₂ mAChR mutants using pseudoequilibrium binding (left panels) or dissociation kinetic (right panels) assays at 37°C on membranes from CHO FlpIn cells. The curves superimposed on the data for each modulator represent the best global nonlinear regression curve fit of a kinetic ATCM to both the pseudoequilibrium and dissociation kinetic experiments. Points represent the mean ± standard error of the mean of of 3-8 experiments performed in triplicate.

Figure 3 Effects of the combined M₂-EDGE-YT mutations on the ability of mAChR agonists to inhibit the binding of [³H]NMS. Inhibition of 0.5 nM [³H]NMS binding at 37°C by (A) ACh, (B) McN-A-343, (C) AC-42 or (D) 77-LH-28-1 at M₂-WT (●) and M₂-EDGE-YT (○) mAChRs in membranes from CHO FlpIn cells. The curves superimposed on the data points represent the best global fit of an empirical one or two-site mass action binding model, as
determined by an F-test (see Table 2). In each case, data points represent the mean + standard error of the mean obtained from three experiments conducted in triplicate.

Figure 4 Allosteric agonists can retard the dissociation rate of $[^3$H]NMS from the M$_2$ mAChR. $[^3$H]NMS dissociation determined in the absence (●) or presence of 300 µM McN-A-343 (○), 100 µM AC-42 (□) and 77-LH-28-1 (■) at 37°C to (A) M$_2$-WT (●) or (B) M$_2$-EDGE-YT (○) mAChRs in membranes from CHO FlpIn cells. (C) Full concentration-response relationship of the effect of McN-A-343 on the dissociation rate of $[^3$H]NMS at 37°C from the M$_2$-WT (●) and M$_2$-EDGE+YT (○) mAChRs. Data represent the mean + standard error of the mean obtained from three to five experiments conducted in duplicate.

Figure 5 McN-A-343 can cause kinetic binding artifacts on $[^3$H]NMS pseudo-equilibrium binding. M$_2$ mAChRs were either pre-equilibrated with $[^3$H]NMS prior to addition of unlabelled inhibitor (○), or exposed simultaneously to a high concentration of $[^3$H]NMS and inhibitor prior to dilution and incubation for an additional 20 minutes (●) at 37°C. Effects of (A) atropine at M$_2$-WT mAChRs, (B) McN-A-343 at M$_2$-WT mAChRs and (C) McN-A-343 at M$_2$-EDGE-YT mAChRs in membranes from CHO FlpIn cells. Curves superimposed on the data represent the best global fit of (A) a monophasic inhibition mass action curve (B, C) a kinetic ATCM with the following parameter values (B): pK$_A$ = 9.17; pK$_B$ = 4.70 ± 0.17; Log $\alpha$ = -1.40 ± 0.14; k$_{off}$ = 0.29 min$^{-1}$; k$_{offB}$ = 0.002 min$^{-1}$; (C) pK$_A$ = 8.7; pK$_B$ = 5.10 ± 0.01; Log $\alpha$ = -2.37 ± 0.05; k$_{off}$ = 0.23 min$^{-1}$; k$_{offB}$ = 0.04 min$^{-1}$. Values not associated with standard errors were determined in separate experiments and fixed as constants in the current analysis. In each case,
data points represent the mean + standard error of the mean obtained from three experiments performed in triplicate.

**Figure 6**  **Time course of M₂ mAChR-mediated ERK1/2 phosphorylation.** Effects of ACh (1 µM; ●), pilocarpine (100 µM; ▲), McN-A-343 (100 µM; ○), AC-42 (100 µM; □) and 77-LH-28-1 (100 µM; ■) on ERK1/2 phosphorylation at 37°C in CHO FlpIn cells stably expressing (A) M₂-WT or (B) M₂-EDGE-YT mAChRs. Data points represent the mean + standard error of the mean obtained from three to six experiments conducted in duplicate, apart from pilocarpine which is the mean from one experiment conducted in duplicate.

**Figure 7**  **Concordance between effects of M₂ mAChR mutation on allosteric agonism at two different signaling pathways.** (A and B) Concentration-response curves to ACh (5 min stimulation; ●), pilocarpine (8 min stimulation; ▲), McN-A-343 (5 min stimulation; ○), AC-42 (8 min stimulation; □) and 77-LH-28-1 (8 min stimulation; ■) mediated ERK1/2 phosphorylation at 37°C in CHO FlpIn cells stably expressing the M₂-WT (A) or M₂-EDGE-YT (B) mAChRs. Data points represent the mean + standard error of the mean obtained from six to twelve experiments conducted in duplicate. (C and D) Concentration-response curves to ACh (●), pilocarpine (▲) and McN-A-343 (○) mediated stimulation of [³⁵S]GTPγS binding to activated G proteins in FlpInCHO cell membranes stably expressing the M₂ wild type (C) and M₂-EDGE+YT (D) mAChRs at 30°C. Data points represent individual means obtained from two experiments conducted in duplicate.
Figure 8  Enhancement of the efficacy of McN-A-343 at mutant M₂ mAChRs is mediated mainly by the residues, Y¹⁷⁷ and T⁴²³. Concentration-response curves to ACh (5 min stimulation; ●), McN-A-343 (5 min stimulation; ○), AC-42 (8 min stimulation; □) and 77-LH-28-1 (8 min stimulation; ■) mediated ERK1/2 phosphorylation at 37°C in CHO FlpIn cells stably expressing the (A) M₂-EDGE or (B) M₂-YT mAChRs. Data points represent the mean ± standard error of the mean obtained from five to twelve experiments conducted in duplicate.

Figure 9  Characterization of the interaction between ACh and allosteric agonists at M₂-WT mAChRs. ACh mediated ERK1/2 phosphorylation in the absence (●) or presence of 10 µM (○), 100 µM (■) or 1 mM (□) McN-A-343 (A) or 1 μM (○), 3 μM (■), 10 μM (□) or 30 μM (♦) AC-42 (B) or 77-LH-28-1 (C) at 37°C in CHO FlpIn cells stably expressing the M₂-WT mAChR. Curves superimposed on the data represent the best global fit of an operational model of competitive antagonism. Data points represent the mean ± standard error of the mean obtained from five experiments conducted in duplicate.

Figure 10  Characterization of the interaction between ACh and allosteric agonists at M₂-EDGE-YT mAChRs. ACh mediated ERK1/2 phosphorylation in the absence (●) or presence of 1 μM (○), 3 μM (■), 10 μM (□) or 30 μM (♦) AC-42 (A) or 77-LH-28-1 (B) at 37°C in CHO FlpIn cells stably expressing the M₂-EDGE-YT mAChR. Curves superimposed on the data represent the best global fit of an operational model of competitive antagonism. Data points represent the mean ± standard error of the mean obtained from four experiments conducted in duplicate.
Table 1  Allosteric model binding parameters for the interaction between the orthosteric antagonist, \([^3\text{H}]\text{NMS}\), and each of three prototypical allosteric modulators at various M2 mAChR mutants. Values represent the mean ± standard error of the mean from 3-8 experiments performed in triplicate.

<table>
<thead>
<tr>
<th>M2 Receptor Construct</th>
<th>Orthosteric Ligand</th>
<th>Allosteric Modulator</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>[^3\text{H}]\text{NMS}\</td>
<td>Alcuronium</td>
</tr>
<tr>
<td></td>
<td>(\text{pK}_B^a)</td>
<td>(\text{B}_{\text{max}}^b) (pmol/mg)</td>
</tr>
<tr>
<td>M2-WT</td>
<td>9.20 ± 0.20</td>
<td>2.30 ± 0.30</td>
</tr>
<tr>
<td>M2-EDGE</td>
<td>9.16 ± 0.36</td>
<td>1.00 ± 0.16*</td>
</tr>
<tr>
<td>M2-YT</td>
<td>8.80 ± 0.12</td>
<td>2.93 ± 0.96</td>
</tr>
<tr>
<td>M2-EDGE-YT</td>
<td>8.77 ± 0.28</td>
<td>3.15 ± 0.15</td>
</tr>
</tbody>
</table>

* Significantly different (p < 0.05) from the wild type receptor as determined by one way ANOVA.

\(a\) Negative logarithm of the radioligand equilibrium dissociation constant.

\(b\) Maximal density of binding sites.

\(c\) Negative logarithm of the allosteric modulator equilibrium dissociation constant.

\(d\) Logarithm of the cooperativity factor governing the allosteric interaction between the modulator and \([^3\text{H}]\text{NMS}\).
Table 2  Empirical inhibition binding parameters for ACh, McN-A-343, AC-42 and 77-LH-28-1 against [³H]NMS at the M₂-WT and M₂-EDGE-YT mAChRs. Values represent the mean ± standard error of the mean from three experiments performed in triplicate.

<table>
<thead>
<tr>
<th></th>
<th>M₂ - WT</th>
<th>M₂-EDGE-YT</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>pKᵢ(High)</td>
<td>pKᵢ(Low)</td>
</tr>
<tr>
<td>ACh</td>
<td>7.08 ± 0.13</td>
<td>5.61 ± 0.17</td>
</tr>
<tr>
<td>McN-A-343</td>
<td>6.14 ± 0.16</td>
<td>4.89 ± 0.08</td>
</tr>
<tr>
<td>AC-42</td>
<td>-</td>
<td>5.42 ± 0.06</td>
</tr>
<tr>
<td>77-LH-28-1</td>
<td>-</td>
<td>6.23 ± 0.03</td>
</tr>
</tbody>
</table>

* Negative logarithm of the apparent high affinity equilibrium dissociation constant.

# Negative logarithm of the apparent low affinity equilibrium dissociation constant.

Fraction of the receptors exhibiting high affinity binding.

Hill slope.

* Significantly different (p < 0.05) from the corresponding control value within the same treatment group.
Table 3  Estimated potency ($pE_{50}$) and maximal agonist effect ($E_{max}$) of mAChR agonists at mediating ERK1/2 phosphorylation. Values represent the mean ± standard error of the mean obtained from 5-12 experiments performed in duplicate.

<table>
<thead>
<tr>
<th></th>
<th>M$_2$-WT</th>
<th>M$_2$-EDGE-YT</th>
<th>M$_2$-EDGE</th>
<th>M$_2$-YT</th>
</tr>
</thead>
<tbody>
<tr>
<td>Acetylcholine $pE_{50}$</td>
<td>7.52 ± 0.12</td>
<td>7.38 ± 0.12</td>
<td>7.64 ± 0.10</td>
<td>7.50 ± 0.09</td>
</tr>
<tr>
<td></td>
<td>$E_{max}$</td>
<td>90.8 ± 6.1</td>
<td>105.3 ± 4.3</td>
<td>77.1 ± 6.1</td>
</tr>
<tr>
<td>Pilocarpine $pE_{50}$</td>
<td>6.16 ± 0.24</td>
<td>5.66 ± 0.06</td>
<td>N/P</td>
<td>N/P</td>
</tr>
<tr>
<td></td>
<td>$E_{max}$</td>
<td>80.2 ± 5.6</td>
<td>108.3 ± 2.4*</td>
<td>N/P</td>
</tr>
<tr>
<td>McN-A-343 $pE_{50}$</td>
<td>5.35 ± 0.05</td>
<td>5.53 ± 0.12</td>
<td>5.10 ± 0.12</td>
<td>5.38 ± 0.11</td>
</tr>
<tr>
<td></td>
<td>$E_{max}$</td>
<td>18.0 ± 4.1</td>
<td>97.1 ± 4.3*</td>
<td>32.8 ± 8.1</td>
</tr>
<tr>
<td>AC-42 $pE_{50}$</td>
<td>N/D</td>
<td>N/D</td>
<td>N/D</td>
<td>N/D</td>
</tr>
<tr>
<td></td>
<td>$E_{max}$</td>
<td>N/D</td>
<td>N/D</td>
<td>N/D</td>
</tr>
<tr>
<td>77-LH-28-1 $pE_{50}$</td>
<td>5.87 ± 0.05</td>
<td>6.98 ± 0.15*</td>
<td>5.83 ± 0.53</td>
<td>6.78 ± 0.13*</td>
</tr>
<tr>
<td></td>
<td>$E_{max}$</td>
<td>16.8 ± 2.2</td>
<td>27.9 ± 4.7</td>
<td>13.43 ± 2.6</td>
</tr>
</tbody>
</table>

*Significantly different ($p < 0.05$) from the corresponding control value within the same treatment group.

N/D- Value could not be accurately determined due to the very modest level of ERK1/2 phosphorylation.

N/P – Not performed
Table 4  Operational model pK$_B$ and Schild slope estimates for McN-A-343, AC-42 and 77-LH-28-1 at the M$_2$-WT and M$_2$-EDGE-YT mAChRs from interaction studies with ACh in assays of ERK1/2 phosphorylation. Values represent the mean ± standard error of the mean obtained from 4-5 experiments conducted in duplicate.

<table>
<thead>
<tr>
<th></th>
<th>M$_2$-WT</th>
<th>M$_2$-EDGE-YT</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>pK$_B^a$</td>
<td>Schild slope</td>
</tr>
<tr>
<td>McN-A-343</td>
<td>4.69 ± 0.10</td>
<td>1.19 ± 0.12</td>
</tr>
<tr>
<td>AC-42</td>
<td>5.86 ± 0.11</td>
<td>0.92 ± 0.13</td>
</tr>
<tr>
<td>77-LH-28-1</td>
<td>5.95 ± 0.15</td>
<td>1.00 ± 0.16</td>
</tr>
</tbody>
</table>

$^a$ Negative logarithm of the equilibrium dissociation constant, determined with the Schild slope constrained to equal one.

*Significantly different (p < 0.05) from the corresponding control value at the same receptor

N/P – Not performed
Figure 1
Figure 5
Figure 6
Figure 8
Figure 9
Figure 10