Impaired M₃ Muscarinic Acetylcholine Receptor Signal Transduction Through Blockade of Binding of Multiple Proteins to its Third Intracellular Loop

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Third intracellular loop • Muscarinic receptor • Interacting proteins • G-protein coupled receptors • Protein complexes

Abstract
Several motifs found in the third intracellular loop of the M₃ muscarinic receptor are critical for G protein activation and scaffold protein interaction. However, how multiprotein complexes form is not fully understood. A minigene encoding the third intracellular loop of the M₃ muscarinic receptor was constructed to explore whether peptides from this intracellular region could act as inhibitors of the muscarinic multiprotein complex formation and signaling. We found that this construct, when co-expressed with the M₃ receptor, has the ability to act as a competitive antagonist of G protein receptors and receptor-scaffold/accessory proteins. Transient transfection of human embryonic kidney-293 cells with DNA encoding the human M₃ and M₅ receptor subtypes results in a carbachol-dependent increase of inositol phosphate. Co-expression of the M₃ third cytoplasmic loop minigene dramatically reduces both carbachol-mediated G protein activation and inositol phosphate accumulation. Minigene expression also abrogates activation of M₃ and M₅ receptor mitogen-activated protein kinases pathway. Furthermore, minigene expression led to reduced AKT activation. These data, together with results of co-immunoprecipitation of different scaffold and kinase proteins, provide experimental evidence for the role for the third cytoplasmic loop of the human M₃ muscarinic receptor in G-protein activation and multiprotein complex formation.

Introduction
Muscarinic acetylcholine receptors (mACHRs) belong to a class I subfamily of heptahelical, transmembrane G-protein coupled receptors (GPCRs) and are represented by five distinct subtypes, denoted as M₁, M₂, M₃, M₄ and M₅ [1, 2] mACHRs signal through heterotrimeric guanine nucleotide-binding proteins (G-proteins), involving mainly the third intracellular loop (3ILoop). Muscarinic M₁, M₃ and M₅ receptors couple preferentially to the Gq₁₁ subunit type of G-proteins, activating
phospholipase C-β, and inducing a subsequent increase in intracellular calcium concentration [3]. In contrast, M₂ and M₃ couple mainly to G-proteins of the G₁/Q, classes, typically leading to adenylyl cyclase inhibition and activation of inward-rectifier potassium conductance [4, 5]. Among other possible responses, in a suitable cellular context, all mAChR subtypes can regulate a wide network of signaling intermediates, including small GTPase Rho, phospholipase D, phosphoinositide-3 kinase, nonreceptor kinases and mitogen-activated protein kinases [6-9].

Increasing evidence indicates that signaling efficiency/specificity for mACHRs is determined in part by accessory proteins that physically interact or are found in the microenvironment of the receptor [10]. Several proteins have been shown to interact with mACHRs, including other GPCRs, kinases, and scaffold proteins such as β-arrestin [11, 12]. These proteins, along with classical core signaling entities (receptor, G protein and effectors), contribute to form a signalsome complex at the cytoplasmic face of the receptor [13]. Understanding the nature and features of such a complex may be a key step in designing novel strategies to develop next generation drugs. Sequence similarities of different receptor subtypes at the ligand binding sites is the main hurdle for designing and identifying proper subtype-selective ligands [14, 15]. However, important differences in size and sequence homology of the intracellular loops across different muscarinic receptors can be used to specifically identify each subtype. Previous studies using receptor-derived peptides from specific regions of the M₁ and M₃ receptors have shown the C-terminal tail of the 3ILoop is critical for receptor-G protein interaction [16] and the resulting signal transduction mediated by G proteins [17]. More recently, specific motifs in the 3ILoop of M₁ and M₃ receptors have been shown to bind some accessory proteins with high affinity (calmodulin, oncogenic SET protein, and small GTPase Rho) [12, 13, 18]. This experimental evidence points to a specific role of 3ILoop in receptor-G protein coupling, signal transduction and multiprotein complex formation. Thus, we hypothesized that the soluble expressed 3ILoop could act as a 3ILoop receptor analogue, competing with the receptor for its interacting proteins, and affecting the specific G-protein-mediated downstream effects.

To verify this hypothesis, we developed a minigene construct expressing the human M₁ muscarinic acetylcholine receptor 3ILoop (M₁-R-3ILoop minigene), and evaluated its ability to affect downstream mACHRs signaling to G₁/q11 proteins.

### Materials and Methods

#### Materials

Dulbecco’s modified Eagle’s medium, penicillin/streptomycin and fetal bovine serum were purchased from Invitrogen (Carlsbad, CA, USA). [³⁵S]-GTPyS (1,202 Ci/mmol), [³H]-myo-inositol (3.0 Ci/ml) and N-[³H]-methylscopolamine ([³H]-NMS, 81 Ci/mmol), were from Amersham Biosciences (Piscataway, NJ, USA). Restriction enzymes were from New England Biolabs (Beverly, MA, USA). The rabbit anti-hemagglutinin (HA) polyclonal antibody (clone HA.11) was purchased from Covance (Berkeley, CA, USA). Carbamylcholine chloride (carbachol), atropine sulfate and all other reagents were purchased from Sigma-Aldrich (St. Louis, MO, USA). The pcDNA3-Flag-GRK2 encoding the bovine G receptor kinase-2 was kindly provided by Y. Chen (FUMC, Shanghai, China) and the pcDNA3-Flag-CK1-α encoding the human casein kinase 1-α was kindly provided by M. Bini (Palermo University, Italy).

#### Plasmid constructs

The constructs presented here were made using standard techniques employing PCR and fragment replacement strategies. The cDNAs for the human M₁ and M₃ mAChRs (kindly provided by T. Bonner, NIH, USA and D. Bello, ETH, Switzerland, respectively) were subcloned into the mammalian expression vector pcDNA-3.1 (Invitrogen) containing three HA epitopes (gift from P. Calvo, SFU, CA, USA), thus resulting in the 3xHA-M₁-R-pcDNA3.1 and 3xHA-M₃-R-pcDNA3.1 vectors. Briefly, a 1.9 and 1.7 kb fragments encoding the human M₁, R and M₃-R were respectively amplified using sense and antisense primers harboring unique EcoRV and XbaI sites and then subcloned into EcoRV/XbaI sites of the mammalian pcDNA3.1-3xHA vector.

The cDNAs fragment encoding the entire third intracellular loop (3ILoop) of M₁ muscarinic acetylcholine receptor (Thr249-Ser495) was subcloned into the InterPlay® mammalian TAP system, pNTAP-B vector (Stratagene, La Jolla, CA, USA) resulting in the pTAP-M₁-R-3ILoop vectors (M₁-R-3ILoop minigene). The 3ILoop of M₁-R was amplified from the 3xHA-M₁-R-pcDNA vectors using the Expand High Fidelity PCR System (Roche, Basel, Switzerland) and subcloned into BamHI/EcoRI sites of the pNTAP-B vector.

#### Cell culture and transfection

HEK293T cells were maintained in Dulbecco’s modified Eagle’s medium supplemented with 10% fetal calf serum (FCS), 100 U/ml streptomycin, 100 µg/ml penicillin, 2 mM L-glutamine (all from Invitrogen) at 37°C in a humidified 5% CO₂ incubator. For transfections, 2 × 10⁵ cells were seeded into 100-mm dishes. About 24 h later, cells were co-transfected with pTAP-M₁-R-3ILoop vector and the corresponding human muscarinic plasmid, by using the Lipofectamine™ Plus reagent (Invitrogen). Cells were harvested 48 h after transfection and centrifuged at 3000xg for 30 min. Membrane fractions were frozen as aliquots in 5 mM phosphate buffer saline (PBS), pH 7.4, and stored at -80°C until required.

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M₉R-3ILoop minigene transcript analysis. HEK293T cells were washed with PBS 48 h post-transfection, and total RNA was purified using Quick Prep™ total RNA extraction kit (Amersham Biosciences) and subject to RT-PCR (Access RT-PCR system; Promega, USA). Primers were selected according to Gene Bank database (forward: 5'- GCG GAT CCA CTG GAG GAT CTA TAA GG-3'; reverse: 5'- GCC AAT TCG ACC AGG GAC ATC C-3') to amplify a segment of 541 bp. A RT-PCR negative control was performed loading DEPC water instead of cDNAs and a positive control was performed using G3DPH primers.

Membrane preparation and radioligand binding assay

About 48 h after transfection, HEK293T cells were washed twice with cold PBS, harvested and homogenized in binding buffer (25 mM HEPES, pH 7.4, 5 mM MgCl₂, 1 mM EDTA), using a Polytron tissue homogeniser. Cell membranes were collected by centrifugation at 20000xg for 15 min and homogenized as above. After centrifugation at 40000xg for 20 min at 4°C, the final pellet was resuspended in binding buffer, and membranes were either used immediately or frozen in liquid nitrogen and stored at -80°C until needed. Protein concentration was determined by using the Bradford protein assay kit (Bio-Rad, Hercules, CA, USA). To determine the affinity of NMS for each sample, membranes were incubated with different concentrations of [³²P]-NMS (ranging from 12.5 pM to 1.5 nM) in 5 mM sodium phosphate (pH 7.4) containing 5 mM MgCl₂ at 25°C for 60 min. The incubations were stopped by filtration through Whatman (Maidstone, Kent, UK) GF/B filters and washed extensively with ice-cold PBS before scintillation counting. Nonspecific binding was determined in the presence of 10 µM atropine.

Inositol phosphate determination

Transfected HEK293T cells were labeled for 18-24 h with [³²P]-myo-inositol (Amersham Biosciences) in DMEM (with glucose, w/o inositol (Invitrogen). After labeling, cells were washed and preincubated for 5 min in PBS at 37 °C, and subsequently incubated in FCS free medium with different concentrations of carbachol -or without carbachol- in the presence of 10 mM LiCl for 5 min. Reactions were stopped by perchloric acid addition. Inositol phosphates (IPs) were extracted and separated on Dowex AG1-X8 columns (Bio-Rad, Hercules, CA, USA). Total labelled IPs were then counted by liquid scintillation.

[³²P]GTPγS binding assay

HEK293T cell membranes were diluted in an ice-cold buffer containing 10 mM HEPES and 0.1 mM EDTA, 5Mm deoxycholate (pH 7.4). Then pelleted and resuspended in a binding buffer consisting of 10 mM HEPES, 10 mM MgCl₂, and 100 mM NaCl (pH 7.4) at a final protein concentration of 125 µg/ml. Incubations were conducted in a final assay volume of 1 ml (125µg total protein) for 1 h at 30°C in the presence of 1 µM GDP and 0.3 nM [³²P]GTPγS (Amersham Biosciences) and the suitable ligand concentration (carbachol from 1 nM to 1 mM). The reaction was stopped by addition of 5 ml of ice-cold buffer containing HEPES/NaOH (10 mM) (pH 7.4) and MgCl₂ (1 mM), immediately followed by rapid filtration through glass fibre filters GF/C filters (Whatman) presoaked in the same buffer. The filters were washed twice with 5 ml of buffer and the radioactivity was measured by scintillation counting. Nonspecific binding was determined in the presence of 10 µM GTPγS. Assays were performed in triplicate.

Co-immunoprecipitation and western blot

To immunoprecipitate the M₉ mAChR with its associated proteins, plasmids encoding the HA-M₉ receptor as well as the β-arrestin-GFP, Flag-GRK-2 or Flag-CK1-α expressing vectors were transiently co-transfected into HEK293T cells. 48 h later, cells were serum-deprived for 4 h., then incubated with vehicle or carbachol (20 µM) for 10 min and washed once with PBS before being solubilized for immunoprecipitation. Protein immunodetection on membranes was assessed using goat anti-M₉ receptor antibody (1:1000; Santa Cruz Biotechnology, Santa Cruz, CA, USA), rabbit anti-HA polyclonal antibody (1:2000; Covance), mouse anti-flag M2 monoclonal antibody (1:5000; Sigma-Aldrich) and mouse anti-GFP monoclonal antibody (1: 10000; Novus Biological, Spain) as primary antibodies; and then horseradish-peroxidase (HRP) -conjugated goat anti-rabbit IgG (1:60000; Pierce, Rockford, IL, USA) or goat anti-mouse IgG (1:2000; Pierce, Rockford, IL, USA) as secondary antibody and developed using SuperSignal West Pico Chemiluminescent Substrate (Pierce).

MAPK and AKT assay. Co-transfected HEK293T cells expressing M₉ or M₅ receptor subtypes with or without the M₉R-3ILoop minigene were grown to 80% confluence and rendered quiescent by serum starvation overnight before MAPK or AKT phosphorylation assay. Subsequently, additional 2 h incubation in fresh serum-free medium was performed to minimize basal activity. Then, cells were stimulated by adding medium containing the muscarinic agonist carbachol. Rapid rinsing, with ice-cold PBS finished stimulation, and then cell lysis was performed during 10 min by adding 500 µl ice-cold lysis buffer (50 mM Tris-HCL, pH 7.4, 150 mM NaCl, 1% TritonX-100, protease and phosphatase inhibitor cocktail). The cellular debris was removed by centrifugation at 13000xg for 5 min at 4°C, and the total protein content was measured using BCA Protein Assay Reagent (Pierce, Rockford, IL, USA). Aliquots corresponding to 5 µg of protein were mixed with sodium dodecyl sulfate (SDS) loading buffer, applied to 12% SDS-polyacrylamide gel electrophoresis (SDS-PAGE) and analyzed by Western blot. Extracellular signal regulated kinase 1/2 (ERK1/2) and protein kinase B (AKT) activation were assayed by incubating PVDF blots with a mouse antiphospho-ERK1/2 antibody (Sigma-Aldrich) and phospho-AKT antibody (New England Biolabs, UK) respectively. Control blots were also run in parallel and probed with rabbit anti-ERK1/2 antibody (Sigma-Aldrich), and total-AKT antibody that recognized both unphosphorylated and phosphorylated forms. The immunoreactive bands were visualized as described above and then measured by quantitative densitometry.
Forty-eight hours after transfection, HEK293T cells transfected with a constant amount (2 µg) of cDNA of M₃Rluc and increasing amounts of cDNA of E-arrestinGFP₂, were rapidly washed twice in PBS (4% glucose), detached, and resuspended in the same buffer. Cell suspension (40 µg of protein) was distributed in duplicated into 96-well microplates (either clear-bottomed or white opaque plates) for fluorescence and luminescence determinations. The total fluorescence of cell suspensions was quantified and then divided by the background (mock-transfected cells) in a POLARstar Optima plate reader (BMG, Labtechnologies, Offenburg, Germany) equipped with a high-energy xenon flash lamp, using a 10 nm bandwidth excitation filter of 400 nm and an emission filter of 510 nm. Renilla luciferase, total luminescence, was determined on samples incubated 10 min with 5-µM h-coelenterazine as described bellow. The background values for total luminescence were negligible, and they were subtracted from sample values.

For BRET measurement, 40 µg of cell suspensions were distributed in duplicate in 96-well white opaque microplates (Corning, NY) and incubated for 10 min at room temperature in the absence or presence of carbachol. DeepBlueC substrate (Molecular Probes, OR) was added at a final concentration of 5 µM, and readings were performed immediately after, using a POLARstar Optima plate-reader that allows the sequential integration of the signals detected with two filter settings [485 nm (440-500 nm) and 530 nm (510-560 nm)]. The ligand-induced BRET signal is calculated by subtracting the ratio of emission through the acceptor wavelength window over emission through the donor wavelength window for a vehicle-treated cell sample from the same ratio for a second aliquot of the same cells treated with ligand. With this calculation, the vehicle-treated cell sample represents the background, eliminating the requirement for measuring a donor-only control sample. Then, BRET ratio is defined as [GFP₂ emission at 500-530)/(Rluc emission 440-500)] - cf., where cf. corresponds to (emission at 500-530)/(emission at 440-500) for the vehicle-treated cell sample in the same experiment.

**Data analysis**

All binding data were analyzed using the commercial program GraphPad PRISM 4.0 (GraphPad Prism, San Diego, CA, USA). Basal binding was defined as [³⁵S]-GTPγS binding
without agonist in the \(^{35}\text{S}\)-GTP\(_\gamma\)S binding assay. For each agonist concentration, the percentage of binding over basal was calculated to determine the agonist-stimulated \(^{35}\text{S}\)-GTP\(_\gamma\)S binding. Data were fit to a sigmoidal dose-response curve. For statistical evaluation of the biochemical data, unless otherwise specified, one-way analysis of variance (ANOVA) was used followed by Tukey’s Multiple Comparison post-test. \(P\) values less than 0.05 were considered significant.

**Results**

Design, construction and expression of 3ILoop-minigene

To determine whether the expression of the M\(_3\)-3ILoop peptide could function as an M\(_3\) analogue, we generated a minigene vector that encodes the M\(_3\)-3ILoop and a set of peptides (Calmodulin Binding Peptide/Streptavidin Binding Peptide (CBP/SBP)) as described in Materials and Methods (Fig. 1B). HEK293T cells were transiently co-transfected with the M\(_3\)-3ILoop minigene and the M\(_3\) mAChR. Total RNA was isolated 48 hours post-transfection and analyzed by RT-PCR, using a set of primers that spanned the vector (SBP) and inserted sequence (3ILoop), to confirm the transcription of the M\(_3\)-3ILoop minigene. The presence of a single 541-base pair band corresponding to the RT-PCR product confirmed the transfection of cells with the minigene construct (Fig. 1C). To verify the expression of M3R-3ILoop peptide in the transfected HEK293T cells, 48 h post-transfection cells were also harvested and subjected to SDS-PAGE and western blot. A band of approximately 24 kDa molecular mass, corresponding to the expected mass of the designed 3ILoop peptide, was identified in cells that have been transfected with the minigene construct.

Effect of M\(_3\)-3ILoop minigene on receptor expression and ligand binding

The \(K_d\) values, for \([H]\)-NMS binding to membranes, gained from cells expressing the M\(_3\) in the presence and in the absence of the M\(_3\)-3ILoop minigene were not significantly different (Table 1). It is also noticeable that the expression of the M\(_3\)-3ILoop minigene did not alter the receptor density (\(B_{\text{max}}\) value) of the M\(_3\) (Table 1).

HEK293T cells expressing M\(_3\) mAChR were co-transfected with M\(_3\)-3ILoop minigene to determine whether the M\(_3\)-3ILoop protein had any effect on the thermodynamic properties of a related G\(_\gamma\)\(_{11}\)-coupled receptor, the M\(_5\). Interestingly, M\(_5\) showed similar specific \([H]\)-NMS binding properties in the absence or presence of M\(_3\)-3ILoop (Table 1). Overall, these results suggest that M\(_3\)-3ILoop does not affect the conformational state of the orthosteric-binding site of M\(_3\) and M\(_5\).

\(M_{3\text{-}}\text{3ILoop minigene inhibits agonist-mediated stimulation \([35\text{S}]\)-GTP\(_\gamma\)S binding}\)

Since the 3ILoop of M\(_3\) is known to be critical for G-protein coupling and activation [17], the expression of the M\(_3\)-3ILoop minigene might compete for the same pool of G proteins by targeting the receptor-G protein boundary. Therefore, we tested whether the presence of the M\(_3\)-3ILoop minigene, co-transfected in M\(_3\) or M\(_5\) transfected cells, had any effect on G-protein activation which was determined by measuring agonist-induced stimulation of \(^{35}\text{S}\)-GTP\(_\gamma\)S binding to membranes. The ability of carbachol to stimulated \(^{35}\text{S}\)-GTP\(_\gamma\)S binding showed a similar maximal response (\(E_{\text{max}}\)) but different potency (pEC\(_{50}\)) when M3R and M5R were tested (pEC\(_{50}\) = 4.84 ± 0.04 and pEC\(_{50}\) = 5.77 ± 0.05, respectively; means ± S.D., n= 3, \(p<0.05\)) (Fig. 2).

Interestingly, co-expression of M\(_5\) subtype with M\(_3\)-3ILoop minigene resulted in a significant decrement in E\(_{\text{max}}\) (~45%) and 2.2 fold increased of pEC\(_{50}\) of the carbachol-mediated \(^{35}\text{S}\)-GTP\(_\gamma\)S binding, when compared with cells transfected with M\(_3\) alone. We also found that cells expressing the M\(_3\) subtype showed a similar M\(_3\)-3ILoop minigene-mediated change in the agonist-stimulated \(^{35}\text{S}\)-GTP\(_\gamma\)S binding features (maximal response and potency) (Fig. 2).

To further test the specificity of the function of the 3ILoop of the M3 receptor to interact with cellular G proteins, we examined the ability of this peptide to modulate agonist-stimulation of specific \(^{35}\text{S}\)-GTP\(_\gamma\)S

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**Table 1.** Ligand binding properties of M\(_3\) and M\(_3\) mAChR constructs. Radioligand binding studies on HEK293T cell membranes expressing M\(_3\) or M\(_5\) wild type mAChR alone or together with the 3ILoopoop-minigene were carried out as described under Materials and methods. Curves were better fitted by non-linear regression analysis assuming a single binding site. \(K_d\) values were determined by using GraphPad Prism software. Results represent means ± S.E. (n = 5).

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<thead>
<tr>
<th>Receptor construct</th>
<th>(^{[H]})-NMS Binding</th>
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<tr>
<td></td>
<td>(K_d) (pM)</td>
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<tr>
<td>3xHA -M(_3)</td>
<td>85.3 ± 17.7</td>
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<tr>
<td>3xHA -M(_3) + 3ILoop</td>
<td>86.8 ± 27.9</td>
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<tr>
<td>3xHA -M(_5)</td>
<td>164.9 ± 41.4</td>
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<tr>
<td>3xHA -M(_5) + 3ILoop</td>
<td>163.7 ± 55.3</td>
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binding in cells heterologously expressing G<sub>Gi/0</sub>-coupled dopamine D2-long (D<sub>2L</sub>) receptor, as well as cells expressing G<sub>Gs/olf</sub> adenosine A<sub>A2A</sub> receptors. The addition of quinpirole (100nM) in cells expressing D<sub>2L</sub> receptor or CGS 21680 (100nM) in cells expressing A<sub>A2A</sub> receptor, markedly induced an increase in [<sup>35</sup>S]-GTP<sub>JS</sub> binding which was not affected by the expression of the M<sub>R-3ILoop</sub> minigene. These results suggest that M<sub>R-3ILoop</sub> minigene inhibits the G<sub>q/11</sub> protein-coupled mAChRs-mediated [<sup>35</sup>S]-GTP<sub>JS</sub> binding in a specific manner.

**Effect of 3ILoop-minigene Expression on PLC Activation**

We then analyzed agonist-stimulated Ins-(1,4,5)-P3 formation in HEK293T cells expressing each receptor subtype and either the minigene or an empty vector to assess whether the presence of the minigene could compete and recruit G protein-mediated activation of receptors. First, we characterized the time- and concentration-dependence of carbachol-stimulated [<sup>3H</sup>]-myo-InsPs accumulation in transfected M<sub>3</sub> and M<sub>5</sub>-HEK293T cells. [<sup>3H</sup>]-myo-InsPs accumulation reached a peak between 5-10 min after agonist exposure. Complete desensitization occurred within 30 min of high dose stimulation. Nonappreciable carbachol-stimulated [<sup>3H</sup>]-myo-InsPs accumulation was detected in nontransfected cells (Figure not shown).

Fig. 2B shows the concentration-response curves for carbachol-dependent stimulation of [<sup>3H</sup>]-myo-InsPs in M<sub>3</sub> and M<sub>5</sub>-HEK293T cells co-expressing the minigene or the empty vector. Carbachol produced a seven-fold stimulation of inositol phosphate over basal levels, suggesting the presence of endogenous G proteins in HEK293 cells that can effectively activate PLC. In contrast, expression of the minigene vector abolished M<sub>3</sub>-agonist [<sup>3H</sup>]-myo-InsPs accumulation at all agonist concentrations, with a decrease in relative efficacy in about 50%. A similar response was observed in cells expressing M<sub>5</sub> receptors. In this case, carbachol stimulation produced a 6-fold increase in [<sup>3H</sup>]-myo-InsPs accumulation but co-expression of the minigene vector counteracts agonist stimulation (46% reduction in relative efficacy value compared with M5 receptor alone).

Besides the change observed in the maximum PLC response, there was a consistent and statistically significant reduction in the potency of carbachol to stimulate Ins-(1,4,5)-P3 release upon activation of the M3 or M5 receptor ($pEC_{50}$ in the presence of M<sub>R-3ILoop</sub> minigene construct compared with the control, increased $3.1 \pm 0.6$ fold for the M<sub>3</sub> receptor and $3.4 \pm 0.5$ fold for the M<sub>5</sub> receptor).
To ensure that $EC_{50}$ and $E_{\text{max}}$ changes viewed were not due to changes in the receptor density, we plotted the negative logarithm of $EC_{50}$ vs. receptor density for experiments with co-expression of the minigene. Although receptor expression was rather variable, $EC_{50}$ shifts were independent of receptor density (data not shown).

Inhibition of ERK 1/2 and AKT activation is mediated by the expression of minigene construct

Previous experimental evidence supports the idea that at least two distinct mechanisms are involved in activation of ERK1/2 pathway by muscarinic receptors, PCK-dependent and/or receptor phosphorylation dependent [19, 20]. As mentioned above, expression and ligand-binding capacity of $M_3$ and $M_5$ receptor subtypes were unaffected by the presence of minigene vector, whereas coupling to $G_{q/11}$ proteins was altered. Consequently, carbachol failed to stimulate phosphatidylinositol production. We evaluated whether the expression of minigene could further alter ERK1/2 signal transduction after agonist stimulation, and if this vector could be used as a potential inhibitor of this GPCR signaling.

First, a time course for ERK 1/2 activation was performed (Fig. 3A). Stimulation of both $M_3$ receptors co-transfected with empty vector or with minigene, by 20 µM carbachol, caused maximal activation of ERK1/2 at 5 min. However, Fig. 3B shows that ERK1/2 activation by carbachol in cells co-transfected with the minigene was significantly lower than ERK1/2 activation in cells transfected with an empty vector (4-fold in less, P<0.01).

Fig. 3C shows the effects of minigene co-transfection on MAPK activity in cells expressing either $M_3$ or $M_5$ receptor subtypes. A significant decrease in ERK1/2 phosphorylation was observed in both cases when compared with activation in the absence of minigene vector. Preincubation with the MEK specific inhibitor PD-98059 inhibited carbachol-induced ERK1/2 phosphorylation to the same extent as that obtained in cells co-transfected with the minigene construct. This confirmed that $M_3$-3ILoop minigene vector acts as an efficient inhibitory protein of the MAPK signaling pathway after agonist stimulation.

Carbachol binding to the $G_{q/11}$ muscarinic receptors stimulates AKT-mediated cell growth and survival in many cell types [21]. Therefore, the effect of minigene expression on carbachol-induced AKT-activation was also studied.

Activating $M_3$ or $M_5$-expressing cells by carbachol incubation reaches a maximum of AKT phosphorylation...
Fig. 4. 3ILoop effect on AKT phosphorylation mediated upon muscarinic receptor activation. (A) HEK293T cells co-transfected with each receptor subtype and the M3R-3ILoop minigene (lanes 4 and 6 respectively) or empty vector (lanes 3 and 5) were exposed to 1 mM CCh for 10 min after 48 h of transfection. Cell lysates were resolved by SDS-PAGE (12%) and Western blotting performed with an antibody against phosphorylated AKT. Nonstimulated M3-HEK293T cells (lane 1) and CCh-stimulated M3-HEK293T in the presence of a selective PI3K inhibitor, LY294002 (lane 2) were assayed as internal controls. Equal loading was confirmed with rabbit anti-AKT antibody (New England Biolabs, UK). (B) The extent of AKT phosphorylation was quantified by scanning densitometry. Means ± S.E.M is shown; n= 4. **Significantly different compared with M3 (ANOVA; p<0.01). ##Significantly different compared with M5 (ANOVA; p<0.01).

at 10 min after stimulation. No activation occurred in cells pre-incubated with the PI3K antagonist LY294002 as shown in Fig. 4A-B. Western blot analysis revealed that, as previously noted in ERK1/2 signaling, co-transfection of M3 or M5-HEK293T with the minigene vector abolished the phosphorylation of AKT after agonist stimulation (Fig. 4A-B).

Fig. 5. Modulation of scaffold proteins association by co-expression of 3ILoop. (A) HEK293T cells co-transfected with M3 receptor, in the presence or absence of the M3R-3ILoop minigene, plus each of the following proteins, β-arrestin-GFP, Flag-GRK-2 or Flag-CK1-α, were stimulated with CCh before solubilization. Extracts were immunoprecipitated with anti-HA antibody (or nonimmune IgG control) before Western blotting. At the top sections, the immunoprecipitation was probed with an antibody against the M3 muscarinic receptor (goat anti-M3R antibody, 1:1000; Santa Cruz Biotechnology). The middle sections show specific co-immunoprecipitation with β-arrestin-GFP (mouse anti-GFP monoclonal antibody, 1:10000; Novus Biological). A low-level of specific pulldown of β-arrestin-GFP can be seen when cells are co-expressed with the 3ILoop minigene construct. Similar effects were remarked in the bottom sections when immunoprecipitations were probed with anti-FLAG antibody for both kinases (GRK-2 and CK1-α) under the same conditions (mouse anti-flag M2 monoclonal antibody, 1:5000; Sigma-Aldrich). The right panel (positive control) represents the input levels of immunoreactive M3 receptor, β-arrestin-GFP, Flag-GRK-2 or Flag-CK1-α in original extracts. Interacting protein co-immunoprecipitated was normalized for immunoprecipitated 3xHA-M3 receptor. Blots are representative of at least three separate experiments. (B) The extent of scaffold protein immunoprecipitation was quantified by scanning densitometry. Data are the means ± S.E.M from three separate experiments. (**) P<0.01 and (***) P<0.001.
Modulation of receptor functions by sequestration of accessory proteins

Considering the dramatic decrease in ERK1/2 phosphorylation in cells expressing the minigene construct, we decided to analyze whether the minigene expression could modulate receptor function by sequestering receptor scaffold or accessory protein.

At least one of the two mechanisms described for MAPK activation by muscarinic receptor family could involve scaffold proteins, such as ε-arrestin and kinases (GRK-2, GRK-3 and CK1-α), which form a direct complex with the receptor.

Co-immunoprecipitation experiments were carried out to discover if the direct complex formation had been altered or blocked by the presence of the cytoplasmatic 3ILoop. Fig. 5A shows co-immunoprecipitation data from HEK293T cells co-transfected with HA-M₃ receptor in the presence or absence of 3ILoop-M₃ minigene and each of the following vectors: ε-arrestin-GFP2, Flag-GRK-2 and Flag-CK1-α.

Input levels of each protein and the efficiency of M₃ receptor immunoprecipitation were checked to ensure a balance between samples. In addition, we also analyzed the input level of immunoreactive M₃ receptor, ε-arrestin-GFP, Flag-GRK-2 or Flag-CK1-α in original extracts as a positive control of each antibody quality (Fig. 5A, right lane). In all cases, low levels of immunoreactivity were associated with the receptor in basal conditions. However, preincubation of the cells with carbachol caused an increased association of each protein to the M3 receptor, as checked by densitometry of the immunoblots (Fig. 5B). Co-transfection with 3ILoop-M₃ minigene, as well as agonist preincubation, caused a decrease in immunoreactivity for each protein to a level similar to that found under basal conditions. These low levels of co-immunoprecipitation probably resulted from the competition of the M₃R-3ILoop minigene with the same pool of proteins interacting with the receptor, thus blocking their association to the third intracellular loop. Co-immunoprecipitation results could also account for low signal levels gained in ERK ½ phosphorylation assays in the presence of the 3ILoop.

To determine whether sequestration of accessory proteins by the 3ILoop peptide affects their ability to modulate M3 receptor function, we used a BRET assay to determine if the 3ILoop peptide interferes with ε-arrestin recruitment. B-arrestin has been shown to be particularly amenable to BRET analysis, and BRET has been used previously to investigate GPCR-ε-arrestin and ε-arrestin-ubiquitin interactions in parallel [22, 23]. The advantage of BRET compared with a number of other methods is that it is possible to observe interactions in living cells over time, in the presence of agonists and antagonists. We assayed carbachol-induced (1μM) interactions between M₃ receptor and ε-arrestins-1 for 1 min pre-treatment and 120 min post-treatment (different intervals), observing a maximum BRET ratio between 5-20 min (data not shown). In addition, the ability of carbachol to stimulate BRET signal was tested in presence or absence of the 3ILoop. Co-expression of M₃Rluc and ε-arrestinGFP2 in presence of 3ILoop minigene led to a substantial reduction in BRET signal versus M₃Rluc and ε-arrestinGFP2 pair, as seen from the marked reduction of the BRET values (Fig. 6A). These results are consistent with those observed in western blot experiments (Fig. 6B) and phosphorylation level (data not shown) determination; where the presence of 3ILoop produced a dramatic decrease in ε-arrestin binding and receptor interactions.
phosphorylation upon agonist activation.

Discussion

As part of an effort to define the potential role of the 3ILoop in protein-protein interactions and multiprotein complex organization, we report here the co-expression of a minigene construction that encoded the 3ILoop of the M₃ human muscarinic receptor with the intact receptor.

The rationale of this approach was to explore the ability of this structural determinant to interfere with G protein interaction or to compete for other interacting proteins that took part in a putative multiprotein complex formation. Our efforts were specifically focused on the 3ILoop of the M₃ muscarinic receptor subtype, because this region was reported to be involved in direct G protein binding and activation, and the putative site for interaction of a group of scaffold or accessory proteins (arrestin binding, calmodulin and small G protein interactions) [24, 25]. Previous studies of cellular expression of fragments and in vitro G protein activation assays have reported the ability of the intracellular loops-or peptides derived from these loops- to interact with the same molecular partners as the intact receptor [26-28]. These experiments demonstrated that the 3ILoop domain of the M₃ receptor had a recognizable impact on the role of the intact M₃ receptor subtype. In our experiments events occurring at the plasma membrane were dramatically affected in cells co-expressing the 3ILoop minigene construct. Co-expressing the minigene construct with the wild-type receptor decreased G protein activation, phosphatidylinositol production, and subsequent signaling, without affecting normal ligand binding and receptor membrane expression. Our results showed that co-expression of the 3ILoop of the M₃ receptor with the M₃ subtype are similar to previous studies of the 3ILoop of the μ-opioid receptor, which altered the functionality of the intact μ-opioid receptor as well as other classes of GPCR [29].

While we only focused on the study of the effect of the 3ILoop, we cannot exclude the possible involvement of the first and second cytoplasmic loops or the C-terminal tail. Thus, although the 3ILoop is the largest intracellular loop, and is usually proposed to be one of the main sites for intracellular interaction, this does not exclude specific domains in other loops from playing a role in protein-protein interactions such as the formation of a multiprotein complex formation. In fact, the C-terminal tail is recognised to be an important structural determinant with antiapoptotic properties within the muscarinic receptor family [30] and the second intracellular loop, with the DRY motif, has also been described to bind the G protein [31].

In our experimental design, we chose to express the intracellular domain with two-epitope tags (CBP-SBP: TAP-system) at the N-terminal domain to purify the recombinant receptor. This will also allow us to develop further studies using mass spectrometry in order to discover the nature of the interacting proteins. The possibility of conformational and accessibility changes in potentially critical regions close to these tags was unlikely because of their small size (fewer than 15 aa). However, to ensure the proper construct was correctly incorporated into the cell, we used a RT-PCR strategy that involved primers spanning vector and insert sequences that would have been absent in the intact receptor construct.

We found that our minigene-system was able to reduce the mAchR-mediated G protein activation as much as 45%. These observations are consistent with those of other groups using a similar experimental approach with other GPCRs, such as δ-opioid, μ-opioid and α₂-adrenergic receptors [29]. Upon co-expression of the minigene with the M₃ subtype, we also noted low affinity levels with agonist-stimulated [³⁵S]-GTPγS, and a decrease in efficacy. Overall, these experiments suggest that the minigene interferes with the specific function of the G₃q/11-protein pool that couples with M₃ as well as with M₅ subtypes. Supporting this observation, we showed changes in the second messenger levels following expression of the minigene in intact cells. Agonist-mediated activation of Ins-(1,4,5)-P₃ accumulation was blocked in the presence of the minigene, in cells expressing either M₃ or M₅ subtypes. Nevertheless, for M₅ subtype, the inhibitory effect of the 3ILoop was less pronounced. We also saw a decrease, not only in the maximal accumulation of phosphatidylinositol, but also in the potency of carbachol concentration-response curves, indicative of a reversible and competitive process. Our results are consistent with those previously reported for angiotensin receptor, where the presence of the second intracellular loop of angiotensin AT1a receptor resulted in a rise in the angiotensin II concentration needed for 50% stimulation for Ins-(1,4,5)-P₃ release [32].

The low sequence homology within these receptors (at the level of 3ILoop) cannot account for the significant effect detected for the minigene impairment of receptor-G protein coupling process, suggesting that structural G-protein interaction sites are similar in both receptors [33]. This idea is consistent with the substantial reduction of Ins-(1,4,5)-P₃ production and G protein activation, which,
however, did not decrease more than 50% with regard to the wild type. This indicates that multiple distinct structural domains could be involved in this interaction, as seen for dopamine and M₃ muscarinic receptors [16]. An alternative explanation is that the relevant domain that interacts with the G protein is exposed in the natural receptor, while the adapted minigene may not present or contain this structure, and therefore does not effectively compete for the binding with its homologous partner.

Several lines of evidence show that muscarinic receptors can activate the MAPK pathway in various ways, dependent on or independent of the classical G-protein activation pathway [20]. One of these mechanisms is PKC-dependent and is essential in ERK activation; it can involve βγ-subunits and Gₛα₁₁-subunits. Other mechanisms may depend more on the phosphorylation state, and are mediated by a group of different kinases, like GRK2/3 or CK1α, recruiting β-arrestin scaffold protein or other adapter proteins, respectively. Our experiments showed that cells co-expressing the minigene construct failed to phosphorylate ERK1/2 upon receptor activation, suggesting that PKC-dependent mechanism, as well as other mechanisms, which depend upon a group of kinases and accessory proteins, have been blocked.

In addition, using immunoprecipitation and bioluminescent assays, we analyzed whether the intact receptor, once activated, loses its ability to recruit and bind CK1α or β-arrestin in the presence of the minigene. This may help us understand whether the dramatic blockade of ERK1/2 phosphorylation observed involves the inhibitory effects of the minigene construct at the level of the G protein. Our immunoprecipitation and bioluminescent experiments showed a loss of CK1α and β-arrestin respectively, in the presence of the minigene in comparison with the wild type receptor. Taking this observation into consideration, one possible scenario is that the minigene construct not only disrupts the G subunit association, but also acts as an inhibitory or competitor subunit that potentially disrupts the multiprotein complex formation at the third intracellular loop, thus resulting in a markedly reduced ERK1/2 phosphorylation upon M3 receptor activation. Thus, the specific recruitment of accessory/scaffold proteins by the 3ILoop may be important for their ability to modulate M3 receptor function.

The M3 3ILoop is 256 residues long, containing multiple motifs of basic and acidic residues and some currently recognized functional sequence motifs. This primary sequence pattern is not conserved throughout the G protein-coupled receptor superfamily, and not even within the class A rhodopsin-like receptor subfamily that represents the closest structurally related class of GPCRs. This may eventually form a conformational structural motif that will be more broadly representative and determine part of the specificity for each receptor in these families. However, no structural data are available to confirm or refute this hypothesis.

Interacting proteins for such motifs appearing in a modular form can be identified by affinity purification approaches, such as yeast two-hybrid screening and immunoprecipitation. Our current work, in which the minigene construct representing the 3ILoop of the M₃ muscarinic receptor specifically influences the intracellular signaling of the intact receptor, probably blocking putative multiprotein complex formation, supports the notion that important motifs could be present in this region. The fact that this loop is long could make it an ideal tool for exploring potential molecular partners that might mediate the observed effect under different cellular conditions. The use of a TAP-system strategy that complements the co-expression experiments of the construct with the intact receptor under different physiological conditions would be a productive approach to study protein-protein interactions, not only for muscarinic receptors but also for other types of GPCR.

In summary, our results suggest that the presence of the M₃-R-3ILoop minigene construct not only prevents G-protein coupling to M₃ receptors, but also impairs coupling of other GPCRs that selectively interact with the same G protein population (Gₛα₁₁). We found that the minigene construct can inhibit the M₃ muscarinic acetylcholine signal transduction pathway and its functionality, probably by recruiting M₃-R interacting proteins as confirmed by co-immunoprecipitation and bioluminescent experiments. These results highlight the functional relevance of the interplay among GPCRs and selective G-protein pools, a key process for cell regulation. Furthermore, muscarinic receptor-derived peptides could be used as selective inhibitors for protein-protein interaction, this may be useful for developing a novel pharmacological strategies.

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References


