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Endocytosis of activated muscarinic m2 receptor (m2R) in live mouse hippocampal neurons occurs via a clathrin-dependent pathway

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Running title: Muscarinic m2 receptor clathrin-dependent endocytosis in live neurons
ABSTRACT

Our aim was to examine the dynamics of the muscarinic m2 receptor (m2R), a G-protein coupled receptor (GPCR), after agonist activation in living hippocampal neurons, and especially clathrin dependency endocytosis. We have previously shown that the m2R undergoes agonist-induced internalization in vivo. However, the dynamics and the nature of the endocytotic pathway used by m2R after activation are still unknown in living neurons. Using live cell imaging and quantitative analyses, we have monitored the effect of stimulation on the fate of the membrane-bound m2R and on its redistribution in intraneuronal compartments. Shortly (6 min) after activation, m2R is internalized in preexisting clathrin-coated pits and not in newly-formed pits. Furthermore, after clathrin-dependent endocytosis, m2R associates with early and late endosomes and with subcellular organelles involved in degradation. Together, these results provide, for the first time, a description of m2R dynamics and trafficking in living neurons and prove unambiguously that m2R undergoes clathrin-dependent endocytosis before being degraded.

KEYWORDS

Internalization ; G protein-coupled receptor ; Mouse ; Trafficking ; Time lapse confocal microscopy.

ABBREVIATIONS

GPCR, G protein-coupled receptor; m2R, m2 receptor; ACh, acetylcholine; LM, light microscopy; CCh : carbamylcholine ; CCP : clathrin coated pits, Tf, Alexa Fluor® 594 Conjugated Tf, GFP : green fluorescent protein ; ICC : immunocytochemistry ; NHS, normal horse serum ; SEP : super ecliptic pHluorin ; PBS : phosphate-buffered sodium.
INTRODUCTION

Most neurotransmitter and neurotransmitter-related drugs modulate neuronal activity through G-protein-coupled receptors (GPCRs). Mechanisms that control GPCR compartmentalization, including membrane availability, enable a neuron to adapt its response to local changes in neurotransmitter environment.

Ligand-induced endocytosis is characterized by the internalization of membrane molecules, including GPCRs, from the cell surface into internal membrane compartments. Endocytosis is a complex process that involves different steps. First, endocytosis of GPCRs, classically involves recruitment of agonist-occupied receptor into vesicles for entry into the endocytic pathway. This early vesicular trafficking can be divided into two main pathways: the classic, clathrin-mediated endocytic pathway and the atypical, clathrin-independent, that may be caveolin-1 or flotillin-1-enriched lipid-raft-dependent (Hansen & Nichols, 2009). Second, the cell may lead receptor containing vesicles to further endosomal processing through different subcellular compartments and may either recycle the GPCR back to the plasma membrane and/or degrade them. These early and late trafficking events mediate important functions for the neuron, tuning its responsiveness to ligands over both short-term and long-term periods and regulating receptor coupling to signal transduction pathways.

The molecular mechanisms underlying the endocytotic processing are still not clearly defined but are receptor-specific and may vary between cell types. For example, the highly related dopamine D1 or D2 receptors may have different internalization pathways (Vickery & von Zastrow, 1999). Intracellular signaling pathway may also be dependent of the cell type as shown for ErbB2 or 5-HT1A receptor (Carrel et al., 2006; Hashizume et al., 2008).

GPCR endocytosis studies have mostly been performed in cell lines and rarely in neurons. Yet, as polarized and arborized cells, neurons may display endocytosis features that serve their specific physiological functions, including receptor targeting to distinct subcellular compartments (McDonald et al., 2007a).

Our work focuses on the muscarinic receptor m2R, a metabotropic acetylcholine receptor involved in autoregulation of ACh release especially in the hippocampus and cortex (Zhang et al., 2002). In the present study, we have investigated the dynamics of the early endocytosis steps of the acetylcholine muscarinic m2 receptor (m2R) in live neurons. Indeed, the subcellular events after the stimulation of m2R may play a key role in the function of cholinergic neurons, especially in the regulation of their neuronal activity and/or in the inhibition of ACh release. We have previously shown that m2R displays endocytosis in vivo in striatal cholinergic neurons after acute stimulation (Bernard et al., 1998; Liste et al., 2002; Decossas et al., 2003; Bernard et al., 2006). However, the precise endocytotic pathways used by m2R in living neurons are still unknown. One of the aims of our work was to determine whether m2R internalization occurs via clathrin-coated pits.

The m2R dynamics was investigated in hippocampal neurons after agonist activation using new fluorescent m2R fusion proteins N-terminally tagged with green fluorescent protein (GFP) or super-ecliptic pHluorin (SEP), a pH-sensitive chimera which facilitate the detection of surface receptor expression in live cells (McDonald et al., 2007b). Live-cell confocal imaging was used to visualize, analyse and quantify m2R dynamics. Real-time early trafficking events of the m2R were especially examined with regard to clathrin, a key protein of the endocytic pathway, and to other intraneuronal post-endocytic compartments.

MATERIALS AND METHODS
All relevant experimental procedures followed the guidelines of the European Communities Council Directive (86/809/EEC) regarding the care and use of animals for experimental procedures, and the Ministère de l'Agriculture et de la Forêt, Service Vétérinaire de la Santé et de la Protection Animale (permission no. A 94-028-21), and were approved by the Regional Ethics Committee no. 3 of Ile-de-France region on Animal Experiments.

**DNA constructs**

Two plasmids, mM2-pcDPS and pRK-ssGFP-NK3 encoding the m2R and GFP-neurokinin 3 receptor, were used to generate the GFP-m2R construct. The m2R fragment was amplified from the mM2-pcDPS plasmid by PCR and introduced as a SalI/XbaI fragment in the pRK-ssGFP-NK3 plasmid to replace NK3 and to generate a pRK-ssGFP-m2R plasmid and obtain the N-terminal labeled version of the receptor. The GFP-m2R fragment was flanked upstream of GFP by an optimized artificial signal sequence derived from the human growth hormone (hGH1; a signal sequence (ss)) (McDonald et al., 2007b). This plasmid is designated as GFP-m2R throughout this paper. Another plasmid encoding for m2R tagged with the super-ecliptic pHluorin, a pH dependent fluorochrome, was generated (SEP-m2R). The SEP fragment was amplified from the SEP-TOPO plasmid by PCR and introduced as a BglII/SalI fragment in the GFP-m2R plasmid to replace GFP and to generate a pRK-ssSEP-m2R plasmid. This plasmid is designated as SEP-m2R throughout this paper. The mM2-pcDPS, pRK-ssGFP-NK3 and SEP-TOPO were generous gifts from J. Wess (NIH, Bethesda, USA), A. Irving (University of Dundee, UK) and J. Henley (University of Bristol, UK), respectively. Alternatively, we have removed GFP from the pRK-ssGFP-m2R plasmid to produce a pRK-ss-m2R plasmid that coded for the wild-type m2R that was used to check the absence of negative effects of GFP in the endocytotic processes. This plasmid is designated as WT-m2R throughout this paper. The integrity of the constructs was confirmed by sequencing. CAV1-mCherry was a gift from Ari Helenius (Addgene plasmid # 27705, Hayer et al, 2010).

**Neuronal cultures and transfections**

Post-natal day 0 C57BL/6 mice were euthanized by decapitation. Hippocampi were dissected from mouse brains and dissociated in Hanks' Balanced Salt Solution (HBSS) with papaine (Worthington Biochemical Corp. Lakewood, NJ, USA; 9001-73-4; 25U/ml). Hippocampal neurons were plated on glass coverslips previously coated with poly-L-Lysine 0.01% (Sigma-Aldrich, St. Louis, MO, USA). Neurons were grown in Neurobasal A medium supplemented with 2% B27, 1% glutamax and 0.5% penicillin-streptomycin (Life technologies; 10888022; 35050038; 17504044; 15140122; respectively) and maintained in an incubator with 5% CO2. Hippocampal neurons were transfected at day in vitro (DIV) 7 with the appropriate cDNA (WT-m2R, GFP-m2R, SEP-m2R or DsRed-clathrin) using Lipofectamine 2000 (Life technologies; 11668019) in OptiMEM medium (Life technologies; 31985-062). All experiments were performed the day after transfection (DIV8). The m2R is not constitutively expressed by hippocampal neurons in culture.

**Pharmacological treatments.**
The effect of a muscarinic receptor agonist carbamylcholine, further referred to as “carbachol” (CCh) (Sigma, St. Louis, MO, USA), on m2R trafficking was observed in hippocampal neurons. For real-time experiments, the imaging chamber was perfused 1–120 min with 30 or 100µM CCh diluted in the isotonic medium. In some experiments, neurons were perfused with 10nM of the muscarinic receptor antagonist atropine (Sigma-Aldrich, St. Louis, MO, USA), 10 min prior to 100µM CCh. CCh was added then together with atropine. In order to reveal receptors associated with acidic intraneuronal organelles after SEP-m2R transfection, NH4Cl (50mM) was added in the perfusion bath.

For other experiments, neurons were incubated with 30 or 100µM CCh in Neurobasal medium for 3, 6, 20min, 1 or 2 hrs and fixed with 2% paraformaldehyde for 5 min. In some experiments, neurons were perfused with 10nM of the muscarinic receptor antagonist atropine (Sigma, St. Louis, MO, USA) 10 min prior to CCh and then during endocytosis 15 min together with 30µM CCh.

Clathrin-dependent endocytosis blockade

The clathrin-dependence endocytosis of was investigated by blocking this pathway using different biochemical and molecular means.

Molecular manipulation of a selected clathrin-dependent endocytosis pathway protein. To block the clathrin-dependent route in the endocytic pathway, we blocked the function of a key protein in the endocytic pathway, Eps15, by expressing dominant-negative proteins, fused to GFP. DIII and EH29 mutants were generated by deleting distinct parts of the DNA coding for Eps15 (Benmerah et al., 1999). Plasmid constructs of dominant negative Eps15 (DIII and EH29) and control (D3Δ2) were kindly provided by A. Benmerah (Hôpital Necker-Enfants Malades, Paris, France). Neurons were transiently co-transfected with dominant negative plasmids and the WT-m2R plasmid using Lipofectamine 2000 (Life Technologies, Saint Aubin, France). Neurons expressing simultaneously Eps15 mutants or control (identified by GFP staining) and m2R (identified by m2R ICC) were analysed.

Al594-Tf uptake used as a marker of clathrin-mediated endocytosis in hippocampal neurons.

Transferrin uptake occurs through a clathrin-mediated endocytosis (Benmerah et al., 1999). In order to know wether m2R is internalized with the same pathway, we have studied the colocalization of fluorescence for m2R and Al594-Tf in untreated neurons and after CCh stimulation. For that, the neurons were incubated with Al594-Tf alone for 10 min, then with or without CCh for 15 min. After fixation, neurons were observed under the confocal microscope and the colocalization of fluorescent m2R ICC signal and Al594-Tf was analyzed using the Jacop ImageJ Plugin (see below).

Antibodies and immunocytochemistry

Antibodies. The m2R expressed after transfection with the WT-m2R plasmid was immunolocalized using a monoclonal anti-m2R antibody raised in rat against an intracellular epitope of the receptor (rat, Chemicon, Cat# MAB367, Lot# RRID: AB_94952). The antibody recognized a single band on Western blots corresponding to the m2i3-GST fusion
In immunohistochemistry, it exhibited a pattern identical to that seen previously with polyclonal antibodies against the same antigen (Levey et al., 1991, 1995). No immunoreactivity was seen when the antibody was used on tissue from m2 receptor knockout mice (Duttaroy et al., 2002).

In some experiments, GFP or SEP expressed after transfection with the GFP-m2R or SEP-m2R plasmids was detected using a anti-GFP antibody (mouse, Roche Applied Science Cat# 11814460001, Lot# RRID:AB_390913). To identify subcellular organelles associated with m2R after stimulation with CCh for 3, 6 or 20min, 1 or 2hrs, the following antisera were used: anti-Clathrin heavy chain (CHC; mouse; BD Biosciences Cat# 610499 Lot# RRID:AB_397865); anti-Golgi matrix protein of 130kDa (GM130; mouse; BD Biosciences Cat# 610822 Lot# RRID:AB_398141); anti-Rab5 (mouse; BD Biosciences Cat# 610281 Lot# RRID:AB_397676); anti-Rab9 (mouse; Thermo Fisher Scientific Cat# MA3-067 Lot# RRID:AB_2175599); anti-protein disulphide isomerase (PDI; mouse; Thermo Fisher Scientific Cat# MA3-019 Lot# RRID:AB_2163120); anti-cathepsin D (CathD; mouse; Santa Cruz Biotechnology Cat# sc-6494 Lot# RRID:AB_2087097). Secondary antibodies used were donkey anti-rat Alexa568-conjugated or goat anti-rat Alexa488 (m2R, Thermo Fisher Scientific Cat# A-11077, RRID:AB_2534121 or Molecular Probes Cat# A-11006, RRID:AB_141373; respectively) and goat anti-mouse Alexa488-conjugated (GFP, (Thermo Fisher Scientific Cat# A32723, RRID:AB_2633275) or goat anti-mouse Alexa688-conjugated (GFP, CHC, GM130, Rab5, Rab9, PDI, Molecular Probes Cat# A-11004, RRID:AB_141371).

Immunocytochemistry. Neurons were fixed with 2% paraformaldehyde for 5 min at room temperature. The cells were washed in PBS and incubated 30min with 4% normal donkey serum (Sigma, St. Louis, MO, USA). Primary antibodies were diluted in PBS with 1% normal donkey serum and 0.075% saponin and incubated overnight at 4°C. Neurons were washed in PBS and subsequently incubated with fluorescence-coupled secondary antibodies diluted in PBS with 0.075% saponin for 1 hr at room temperature. Finally, cells were washed in PBS and mounted in Prolong gold (ThermoFisher Scientific).

Time-lapse imaging of cultured hippocampal neurons

Time-lapse imaging was used to analyse 1) the pH-dependence of the SEP-m2R construct, 2) the effect of CCh on the membrane associated m2R and 3) clathrin-dependence of m2R endocytosis and its dynamics.

Time-lapse sequences from cultured hippocampal neurons transfected with selected plasmids were collected using a Leica DMi6000B inverted microscope (Leica Microsystems, Deerfield, IL; USA) equipped with a Yokogawa CSU-X1 spinning disc confocal head (Roper Scientific, Lisses, France) and a 100 mW 491 and 561 nm laser controlled by MetaMorph (Molecular Devices, St. Grégoire, France). The setup was enclosed in a thermal incubator set to 37°C under 5% CO2. Images were collected through a 63x/1.4 numerical aperture oil-immersion objective and an additional 2x lens on a QuantEM:512SC EMCCD (Photometrics, Tucson, AZ).

For validation of SEP pH-dependence, hippocampal neurons transfected with the SEP-m2R were observed under the spinning disk microscope for 30min and the pH of the medium was changed (7.4 to acidic pH (around 6.0)) time to time with or without NH4Cl (50mM). Stacks of images were collected every 30s for 30min. In order to analyse the effect
of CCh on the membrane associated m2R, hippocampal neurons transfected with the SEP-
m2R were observed under the spinning disk microscope for 120 min and stacks of images
were collected. The clathrin-dependence of m2R endocytosis was analysed from double
transfected neurons with and GFP-m2R plasmids. Stacks of images were acquired every 30 s
for 30 min. Images were treated using Fiji (Schindelin et al., 2012) and Adobe Photoshop
softwares.

**Imaging of fixed cultures by confocal microscopy**

Images were acquired on a Leica SP5 confocal system (Leica Microsystems, Deerfield, IL;
USA). z-Series stacks of confocal images were acquired at 1024 × 1024 pixel resolution,
with a pinhole setting of one Airy unit and optimal settings for gain and offset. For double
immunolabeling quantifications, images were taken with a 63×/1.4 numerical aperture
(N.A.) Plan-Apochromat, an argon laser at an excitation wavelength of 488 nm, and a diode
561 nm or two diodes at 561 and 633 nm. Images were treated using Fiji (Schindelin et al.,
2012) and Adobe Photoshop softwares.

**Quantification and statistical analyses**

*Quantification of colocalization of fluorescence*

The quantification of colocalization of m2R with GFP or SEP (for validation of the
constructs) and m2R with clathrin or organelle markers was analyzed with the ‘Just Another
Colocalization Program’ (JACoP) Plugin (ImageJ, National Institutes of Health), and
statistical data are reported from the Costes’s randomization-based colocalization module
(Bolte and Cordelieres 2006). Costes’s randomization method for measurement of
colocalization was used to confirm, with > 95% certainty, that the colocalization observed
between the m2R and clathrin or organelle immunofluorescent signals was not caused by
chance coincidence (Costes et al. 2004). A Pearson’s coefficient (pc) was calculated.
Costes’s randomization was applied on five neurons from four mice of each genotype using
at least 150 iterations per image. For validation of the constructs, analyses were performed
on somatic areas and on the neuropile. For the colocalization of m2R with clathrin and
organelle markers, analyses have been restricted to the somatic area. Just individual images
(and not stacks of images) were analyzed. The quantification of colocalization was
performed from the labeling on images observed with the 63x objective (surface of the field:
655 µm²).
The pc calculated in colocalizations analyses in WT and stimulated neurons were compared
using a Mann-Whitney U test or the Kruskal-Wallis test followed by Dunn’s Multiple
Comparison Test when more than two groups had to be compared. All data are shown as the
means ± SEM; NS : not significant; ***: p<0.0001.

*Quantification of the density of m2R clusters in mutants of Eps 15*

Hippocampal neurons were observed using the 63x objective and acquisitions were
performed under the confocal microscope. Intracellular immunofluorescent clusters,
representing m2R in endosomes, were segmented and counted using the FIJI/ImageJ
software. Results are expressed as intracellular immunofluorescent clusters per µm²
cytoplasmic surface in Eps15 dominant negative-treated and control neurons.

*Quantification of variation of fluorescence in time lapse experiments.* The quantification
of variation of fluorescence levels with time was automatically performed using the Fiji
software. Mean intensity measurements following background subtraction from the whole
neuron were pooled for each cell. Control values corresponded to the mean fluorescence
intensity immediately at the beginning of the experiment or before addition of the
muscarinic agonist. Data were compared using the repeated ANOVA test followed by the
Dunnet post-hoc test comparing each value to the value at the beginning of CCh treatment.
The $p$ values values are: * $p < 0.05$; **: $p<0.001$; ***: $p<0.0001$. The quantification of the
number of clusters of internalized m2R per surface of neuron after CCh treatment was
performed on projections of the stacks images. Data were compared using the repeated
ANOVA test followed by the Dunnet post-hoc test comparing each value to the value at
0 min or the Mann-Whitney U test when the data were unpaired. The $p$ values values are: *
$p < 0.05$; **: $p<0.001$; ***: $p<0.0001$.

All the experiments have been replicated at least three times. For quantitative studies, 15 to
25 neurons per group were analyzed.
RESULTS

Validation of the GFP-, SEP- and WT-m2R constructs

Expression of GFP-m2R and SEP-m2R in living and fixed neurons. To analyse the dynamics of GPCRs with high resolution in fixed and living neurons, the m2R was tagged at the N terminus with GFP or SEP. By live or fixed-cell confocal microscopy, we have detected a predominant plasma membrane distribution of GFP-m2R or SEP-m2R fluorescence (Fig. 1A,A’,A’’,C,C’’,D,D’’,E,G) in the cell body and the proximal and distal dendrites of hippocampal neurons. This distribution was similar to that detected for the m2R using a third intracellular loop directed antibody (Fig. 1A’,A’’,D,D’’) or to the endogenous receptor in hippocampal neurons (Bernard et al., 2003). Cytoplasmic fluorescence signal was low. Similar labelings were observed after transfection with the WT-m2R plasmid and immunocytochemistry with an m2R antibody (Fig. 1H). The analysis of the colocalization of GFP with m2R-ICC and SEP with m2R-ICC was performed on fixed neurons using the Jacop Plugin of ImageJ are reported from the Costes’ randomization-based colocalization module (see Materials and methods). The high Pearson’s coefficient at or higher than 0.8 (0.8707+/−0.0171, n=22; 0.8489+/−0.0199, n=25; respectively) confirmed the validation of the GFP-, SEP- and WT-m2R constructs.

GFP and SEP are correctly addressed to the external plasma membrane. In order to check that the m2R was correctly folded and addressed as expected to the plasma membrane with its N terminus at its extracellular side, we have co-detected GFP and SEP by their native fluorescence and by immunocytochemistry using an anti-GFP antibody. GFP immunofluorescence on fixed and unpermeabilized neurons, was restricted to the plasma membrane in perikarya and dendrites (Fig. 1C’,C’’). The analysis of the colocalization of GFP with GFP-ICC and SEP with GFP-ICC using the Jacop Plugin of ImageJ gave a high Pearson’s coefficient (0.8006±0.0266, n=14 and 0.7696±0.0612, n=7; respectively) confirmed the validation of the GFP-, SEP- and WT-m2R constructs.

Surface expression of GFP-m2R and SEP-m2R is dynamically regulated by agonist exposure. As for other GPCRs, the m2R is internalized from the cell surface following agonist binding (Bernard et al., 1997; Bernard et al., 1998; Bernard et al., 2003; Bernard et al., 2006). We especially checked that GFP or SEP did not disturb m2R internalization. The GFP-m2R and SEP-m2R underwent a time-dependent loss of cell surface receptors following CCh (100 µM) exposure in agreement with data for endocytosis of wild-type m2R expressed in neurons (Fig. 1B,B’,B’,F) (Bernard et al., 1998; Bernard et al., 2003; Bernard et al., 2006). Internalization was also observed when neurons were transfected with the WT-m2R plasmid and the m2R detected by immunocytochemistry with an m2R antibody (Fig. 1I).

Validation of pH-dependence of SEP-m2R. Although GFP is useful to report receptor localization, it is not possible to distinguish between surface and intracellular receptors in live cells using fusion proteins incorporating GFP. However, SEP, a pH-sensitive variant of GFP can be used to report surface expression, when expressed at an extracellular site (Miesenbock et al., 1998; Ashby et al., 2006; McDonald et al., 2007b). Genetically encoding SEP into the extracellular domain of a membrane protein of interest positions the fluorophore on the luminal side of the endoplasmic reticulum (ER) and in the extracellular region of the cell. SEP is fluorescent when the pH is greater than 6, but remains in an off state at lower pH values. Therefore, receptors tagged with SEP fluoresce when residing in
the ER or upon insertion in the plasma membrane (PM) but not when confined to a trafficking vesicle. We therefore generated a SEP-m2R chimera by switching GFP for SEP in the original GFP-m2R construct. The pH dependence of SEP-m2R fluorescence was characterized in transfected hippocampal neurons. At physiological pH 7.4, SEP exhibits similar fluorescence to wild-type GFP (Fig. 2A). A strong membrane-associated fluorescence was observed in the whole dendritic arborization and in the cell body. A faint staining was seen in the cytoplasm. When the pH is decreased to 6, the dendritic and cell body membrane labeling strongly decreases as well as the intracytoplasmic staining (-81% and -98%, respectively) (Fig. 2B,E). Changes in fluorescence levels are reversible when the pH is back to 7.4 (Fig. 2C,E). Also, SEP-m2R fluorescence is visible again at the plasma membrane of dendrites and cell body (Fig. 2C). The quantification shows that SEP-m2R fluorescence is 10% under control (pH 7.4) values in the whole neuron (Fig. 2E).

Administration of NH4Cl (50mM), a compound that equilibrates luminal pH of acidic intracellular vesicles to the extracellular neutral pH, reveals labeling in some intracytoplasmic organelles when the pH is back around 6.0 to 7.4, and the SEP-m2R fluorescence is 59% above control values (Fig. 2C,E). In contrast, when the extracellular medium, including intravesicular pH (due to NH4Cl presence) is set to a more acidic pH (around 6.0), SEP-m2R labeling disappears at membranes and in the soma (-87% and -84% (Fig.2D,E)).

**Specificity of activation of muscarinic receptors by carbachol.** Receptor internalization induced by CCh was totally blocked in the presence of the muscarinic receptor antagonist atropine (10nM) (Fig. 1J).

All validation experiments were performed on three independent cultures, and in each culture, at least ten neurons were analysed.

**Dynamics of m2R internalization**

To investigate the dynamic properties of the m2R after agonist activation, we have developed a combination of experiments using individual live hippocampal neurons transfected with the GFP-m2R or SEP-m2R plasmids and exposed to 100µM CCh for 100-120 min and monitored by time-lapse confocal microscopy. SEP-m2R staining allowed us to analyse and quantify the dynamics of membrane m2R disappearance only. GFP-m2R labeling experiments were useful to analyse the dynamics of m2R internalization.

SEP-m2R fluorescence slowly drops to reach a minimum of 62% of initial levels 54min after the application of the drug (Fig. 3B,D). Then SEP-m2R labeling rises back to reach control values at 84min after exposure to CCh. The statistical analysis shows a significant decrease of the m2R, 39, 54 and 69min after the initiation of the treatment (Fig.3D). However, m2R fluorescence levels are not different from 39 to 69min, revealing a steady state of the m2R intensity levels. Addition of NH4Cl reveals that m2R is associated with acidic compartments, by inducing the appearance of a strong punctiform labelling in the cytoplasm at the level of the soma and dendrites (Fig. 3C,D).

While SEP-m2R fluorescence decreases at the plasma membrane, a punctiform GFP-m2R labeling appears close to the membrane and in the cytoplasm as early as 6 min after administration of CCh (Fig. 4C). The statistical analysis demonstrates that the number of m2R clusters significantly increases as early as 10 min after the beginning of CCh treatment (Fig. 4G). Time-lapse analysis shows that the number of internalized clusters increases with
agonist exposure during the first 30 min and stabilizes afterwards.

m2R endocytosis in clathrin-coated pits (CCP)

Plasma membrane proteins, especially GPCRs, internalize following agonist stimulation through a variety of distinct endocytic pathways (Doherty & McMahon, 2009). The best-characterized pathway is the well-known clathrin-dependent endocytosis, although other clathrin-independent pathways also exist (Roseberry & Hosey, 2001). The nature of the m2R endocytotic pathway, especially in neurons, is still under debate. We have tested here the m2R clathrin-dependent endocytosis hypothesis.

In order to know if m2R clusters after stimulation in native CCPs, we have identified these compartments in fixed neurons using an anti-clathrin heavy chain antibody in cells transfected with WT-m2R. In control neurons, no obvious colocalization of fluorescent m2R and clathrin was detected (Fig. 5A-A’). In contrast, m2R is highly colocalized with clathrin in neurons treated with CCh (Fig.5B-B’; arrows). The statistical analysis confirmed that Pearson’s coefficients are significantly different in control compared to treated neurons (Mann-Whitney U test : p<0.0001; Fig. 5C).

Live imaging allowed us to determine that activated m2R clusterized to preexisting CCPs in neuronal cells. For that, live hippocampal neurons were co-transfected with GFP-m2R and DsRed-Clathrin and observed by spinning disk confocal microscopy before and during a 30-min-long CCh (30µM) treatment. Our observations revealed that agonist treatment clearly induced appearance of green fluorescence clusters in both cell bodies and dendrites (Fig. 6D-E’) as early as 6 min after administration of CCh. These clusters co-localized in CCPs identified by DsRed-Clathrin expression (arrows in Fig. 6D-E’). These results suggest that membrane-associated receptors move to preexisting CCPs shortly (6 min) after activation.

Al594-Tf uptake as a marker of m2R clathrin-mediated endocytosis

In order to know if m2R internalization involves clathrin-mediated endocytosis (CME), we analyzed uptake of a protein that is well known to display constitutive CME (Benmerah et al., 1999) in control neurons and in neurons treated with CCh. For that, the same neurons were incubated with Al594-Tf all along the experiment (see Materials and methods). In a control neuron, Al594-Tf is detected in the cytoplasm (Fig. 7A’,A’’), whereas m2R is present at the plasma membrane (Fig. 7A,A’’). In contrast, in a neuron treated by CCh, m2R and Al594-Tf often colocalized (Fig. 7B’’). The quantitative analyses of Pearson’s coefficients, demonstrated that colocalization of m2R immunofluorescence and Al594-Tf significantly increased after CCh stimulation (Kruskal-Wallis test, followed by Dunn’s Multiple Comparison Test: ***,p<0.0001; Fig.7C). Atropine prevented the increased colocalization of m2R and Al594-Tf (Fig.7C).

Blockade of m2R clathrin-mediated endocytosis in hippocampal neurons

In order to determine if m2R internalization is strictly clathrin-dependent or whether it may involve other endocytic pathways, we have used dominant-interfering mutant proteins (Eps15).

Expression of Eps15 mutants disrupts m2R trafficking. Eps15 is a constitutive component of plasma membrane CCP (Benmerah et al., 1999). To determine if disruption of
Eps15 function influences m2R trafficking, we co-expressed in living hippocampal neurons two different Eps15 mutants (GFP-EH29 or GFP-DIII) or a control mutant, GFP-D3Δ2, with WT-m2R. We then analysed m2R post-endocytic trafficking by confocal microscopy after a 15-min-long CCh (30µM) treatment. The expression of the mutants was checked by the detection of GFP staining in neurons (Fig. 8A’, B’ and C’). Expression of GFP-EH29 or GFP-DIII mutants completely prevents the punctate staining characteristic of m2R endocytosis induced by CCh. As shown in Fig. 8A and 8B, m2R immunostaining is located mostly at the plasma membrane of soma and dendrites 15 min after the initiation of CCh treatment. In contrast, m2R is still internalized when the control mutant, GFP-D3Δ2, is expressed (Fig. 8C). The statistical analysis (Fig. 8D) confirmed that 1) GFP-EH29 or GFP-DIII mutants block m2R endocytosis (Mann-Whitney U test : NS; Fig. 8D) and 2) the control mutant has no effect on m2R internalization (Mann-Whitney U test :***: p<0.0001; Fig. 8D).

Role of caveolin 1 in m2R endocytosis

Proteins may also internalize through clathrin-independent pathways (Doherty & McMahon, 2009). One of these pathways involves caveolae. To address the question of a role of caveolae-dependent pathway in m2R internalization, we have co-transfected the Cav1-mCherry plasmid with WT-m2R, and we have quantified colocalization of Cav1 and m2R fluorescent signals without or after 6 min, 12 min or 15 min of treatment with CCh (Fig. 9). We did not find any difference in Pearson’s coefficients between control and treated neurons (Mann-Whitney U test : NS; Fig. 9C).

Post-endocytic fate of m2R

Shortly after activation (6 min), m2R immunoreactivity is detected in numerous vesicles positive for CHC, a marker of CCP (CHC), EEA1 markers of early (EEA1) and late endosomes (Rab9) (Fig. 10A-C’’’) and cathepsin D, a marker of lysosomes (Fig. 10F-F’’’). After 20 min of CCh exposure, m2R immunoreactivity is identified in some vesicles positive for PDI, a marker of endoplasmic reticulum and in GM130, a marker of Golgi apparatus (Fig. 10D-E’’’).
DISCUSSION

In the present study, we have developed a live-cell imaging approach to gain insights into the dynamics of a GPCR in living neurons, the muscarinic m2R. We have produced and validated different DNA constructs to allow expression of m2R in hippocampal neurons in vitro. We have especially studied the early steps of m2R endocytosis triggered by the stimulation. We have demonstrated, for the first time, that m2R is internalized in live neurons, after stimulation by an agonist, through a clathrin-mediated endocytic pathway.

Methodological aspects

Structural validation of constructions: Consideration of receptor structure and function are important factors in the generation of fluorescent tag/GPCR chimeras. The construction has to preserve the native ability to address the m2R to the plasma membrane, to bind its ligands and not to modify intracellular receptor signaling. We have thus chosen to attach GFP at the extracellular N-terminus of the m2R, since this site is commonly used as a tagging site for many other GPCRs (McDonald et al., 2007b; Lelouvier et al., 2008). Several complementary experiments argue for the fact that our m2R-GFP and m2R-SEP constructs are valuable tools for such dynamic studies. Indeed, the m2R-GFP and m2R-SEP displayed the same subcellular localization as the wild type or the endogenous m2R (Bernard et al., 1998); i.e., is homogeneously distributed at the plasma membrane of the somatodendritic domain. With or without tag, most of m2Rs, revealed by GFP, or SEP or ICC, are detected at the plasma membrane, suggesting that the m2R is correctly addressed to the plasma membrane of soma and dendrites. Moreover, the easy detection of GFP and SEP at the plasma membrane using an anti-GFP antibody, in a non-permeabilized condition, demonstrates that the tag is correctly fused to the extracellular N-terminus of m2R protein. The colocalization of GFP or SEP fluorescence with anti-m2R ICC shows that GFP and SEP are faithful markers of m2R.

Validation of pH-dependence of SEP-m2R: We used the SEP-m2R constructs to monitor and quantify variations of m2R at the plasma membrane upon agonist stimulation. Since SEP-m2R is tagged at the extracellular N-terminus, SEP will be present in the lumen of intracellular organelles during receptor endocytosis. Since these organelles have acidic pH (Demaurex, 2002), the fluorescence of endocytosed SEP-tagged receptors will be obscured. In agreement with these data, we have indeed demonstrated that 1) SEP-m2R fluorescence is quenched at acidic pH and 2) neutralization of intraneuronal vesicles medium by NH4Cl reveals their content in m2R. The SEP-m2R construct is therefore well suited to studying dynamic changes in surface receptor expression in live cells.

Muscarinic receptor stimulation induces m2R internalization in hippocampal neurons

We have shown here that agonist stimulation of m2R induces internalization of this receptor in neurons in vitro. This is in agreement with previous data observed in vivo for the native receptor (Bernard et al., 1998; Liste et al., 2002; Decossas et al., 2003; Decossas et al., 2005). Our results demonstrate that the incorporation of GFP into the m2R protein does not modify its ability to bind its ligands and internalize upon agonist stimulation. Indeed, m2R internalization induced by CCh stimulation was observed with the same timing (6 min after the initiation of activation) when GFP- and WT-m2R constructions are transfected. Moreover, we have checked that m2R internalization was actually due to specific activation of muscarinic receptors since it was blocked by atropine.
The study of receptor internalization phenomena requires the monitoring of two critical parameters: 1) the variation in receptor availability at the plasma membrane and 2) appearance of these receptors in intraneuronal compartments. The use of the SEP-m2R construct allowed the identification of three steps in the dynamics of membrane m2R density changes induced by agonist stimulation. First, m2R membrane density regularly decreases during the first forty minutes. In the same time, GFP-m2R experiments demonstrate m2R clusters appearance in the somatic and dendritic cytoplasm. The use of NH4Cl on SEP-m2R expressing neurons reveals that the compartments containing m2R are acidic and thus probably correspond to endosomes. Taken together, our results suggest that this first step is mainly operated through endocytosis of membrane m2R into endosomes. Second, surface and internalized m2R densities stabilize (as measured by both SEP- and GFP-m2R). This suggests that the bulk internalization process is over. This may reveal a saturation of the endocytosis machinery, especially saturation of binding to protein involved in endocytosis. Lou et al. (2008) have shown that saturation of the endocytosis process occurs in absence of dynamin 1, a predominant component of the endocytic response. We may also assume that clathrin-dependent endocytosis has limited capacity and is saturated when m2R are saturated themselves as demonstrated for the EGF receptor (Schmidt-Glenewinkel et al., 2008). The third step is characterized by the recovery of m2R at the plasma membrane. At the same time, m2R is still accumulated in acidic compartments in the cytoplasm as demonstrated by NH4Cl application on SEP-m2R neurons and by the appearance of cytoplasmic GFP-m2R clusters. Recycling and/or neosynthesis of m2R may contribute to restore a normal receptor density at membranes. Recycling has been well studied for some GPCRs (Hanyaloglu & von Zastrow, 2008; Lelouvier et al., 2008; Zenko & Hislop, 2017) and was shown to be a key phenomenon in the recovery of cell function. Interference with each process through exposure to monensin (recycling) or cycloheximide (neosynthesis) should help to determine what is the mechanism leading to the normalization of surface m2R density.

The m2R endocytosis is clathrin-dependent

We have demonstrated here unambiguously that the m2R was endocytosed in a clathrin-dependent way in neurons. First, we have shown that, shortly after stimulation, the m2R colocalized with clathrin-coated pits. Second, we have shown that m2R is partly internalized together with Al594-Tf, a molecule known to be internalized through a CME pathway. Third, disruption of CCP using over-expression of Eps15 negative dominants abolished m2R endocytosis. The clathrin-dependence of m2R endocytosis is still under debate (Zenko & Hislop, 2017). Some studies demonstrated that m2R internalization pathway involves CCP only (Pals-Ryliaardsdam et al., 1997; Jones et al., 2006; Yamanushi et al., 2007). Other claimed that m2R is internalized through a clathrin-independent process (Vogler et al., 1999; Delaney et al., 2002; Wan et al., 2015). Ockenga and Tikkanen (2015) propose that m2R endocytosis takes place by means of an atypical clathrin-mediated pathway that may involve a specific subset of CCP. Finally, some authors showed that the internalization of the m2R utilizes neither clathrin-coated pits nor caveolae (Roseberry & Hosey, 2001). These discrepancies may be explained in different ways. The signalling and trafficking properties of GPCRs may depend on the cell and cellular context (Ritter & Hall, 2009). Non-neuronal cells may not natively produce all the proteins involved in the clathrin mediated endocytic machinery. In contrast, we have demonstrated that in neurons, m2R endocytosis is clathrin-
dependent, even without overexpression of any endocytotic complex proteins. Indeed, we
have shown agonist-induced m2R internalization in native CCP, i.e. detected by clathrin
immunocytochemistry.

Many studies have demonstrated the essential role of CCP in endocytosis and
cellular signalling processes at the plasma membrane. CCP have also been shown to play a
role in the transport of hydrolases from the Golgi complex to the lysosome and for polarity
of the basolateral plasma membrane proteins in the epithelial cell line MDCK, and from the
somato-dendritic membrane to axonal membrane in neurons (Deborde et al., 2008). We may
hypothesize that CCP may play a role in the transport of endocytosed m2R from one
subcellular compartment to another. Alternatively, clathrin was also shown to participate in
rapid recycling after cargo accesses early endosomes (Zhao & Keen, 2008). CCP containing
m2Rs may thus contribute to recycling of m2R at the membrane, as we have suggested
above.

The m2R is endocytosed at pre-existing CCP

One important issue in cell biology of CCP is to know whether the agonist stimulation
initiates the formation of clathrin-coated domains that are specialized for endocytosis of
activated m2R or whether they are simply mobilized to pre-existing CCP. Time lapse
experiments on living neurons allowed us to answer that muscarinic receptor activation
induces m2R accumulation in already pre-existing CCP. Indeed, when m2R and clathrin
were co-expressed in the same living neuron, we were able to show that m2R clusters co-
localized with Ds-Red-clathrin spots that were already present in dendrites or in cell bodies
before stimulation. We hypothesize that, upon activation, phosphorylated m2Rs may shift in
mobility and be trapped at preformed CCP. Similar evidence, i.e. activated receptor clathrin-
mediated endocytosis in preexisting CCP, have been obtained previously in non-neuronal
cell cultures (Santini et al., 2002; Scott et al., 2002). Our results with those obtained for
somatostatin type 2A receptors are the only ones to our knowledge to reveal this behaviour
in neurons (Lelouvier et al., 2008). Further experiments are needed in order to determine
whether this feature can be generalized to all GPCRs in neurons.

The study of arrestin translocation and redistribution of receptor-arrestin complexes in CCPs
like β-arrestin 2 and chimera between Eps15, a constitutive component of CCPs, during the
early stages of ligand-mediated endocytosis as shown by Scott et al. (2002) may help to
confirm our data. However, due to the fact that current microscopy techniques cannot
distinguish between single pre-formed clathrin structures at the plasma membrane and
clusters of dynamic clathrin-coated pits, some of which are formed in close proximity to
other clathrin structures, we cannot exclude that part of m2R endocytosis occurs through de
novo CCPs in response to the m2R agonist binding.

The m2R does not involve caveolae-mediated endocytosis.

When m2R and CAV1-mCherry were co-expressed in the same neuron, we did not find that
m2R clusters colocalized with CAV1-mCherry. This suggests that m2R endocytosis does
not use the clathrin-independent pathway involving caveolae. Further experiments are
required to determine whether other m2R undergoes other clathrin-independent endocytotic
pathways.

Post-endocytotic fate of m2R
We have identified to which subcellular organelles m2R is targeted in order to identify the post-endocytotic pathway where the activated receptors are sorted (Fig. 10). We have detected m2R in vesicles expressing Clathrin heavy chain (CHC), EEA1 and Rab9, as soon as 6 min after stimulation (Fig. 10A-C’’). This suggests that, after endocytosis in clathrin-coated pits (identified by CHC immunohistochemistry, Fig. 10A-A’’), m2R is sorted to early, then late endosomes (Fig. 10B-C’’). Early endosomes are considered as the first sites where internalized proteins, including GPCRs, are targeted before being either recycled, or degraded (Lakadamyali et al. 2006).

The colocalization of m2R with cathepsin D, a marker of lysosomes (Fig. 10G-G’’), confirms the hypothesis of m2R degradation after activation and endocytosis. This is in agreement with earlier data showing the accumulation of m2R into multivesicular bodies, which are organelles resulting of the fusion of lysosomes (Bernard et al. 1998; Tsuga et al. 1998).

The fact that m2R content is not increased in endoplasmic reticulum, revealed by PDI ICC in fixed neurons suggests that m2R is not targeted to compartments involved in m2R neosynthesis (Fig. 10C-C’’). This is in agreement with the absence of m2R-SEP labeling in live experiments. Indeed, if m2R-SEP is present in the endoplasmic reticulum, a neutral compartment, SEP should emit light. This is in agreement with earlier studies (Bernard et al. 1998).

**CONCLUSION**

We have demonstrated here for the first time that m2R is endocytosed into living neurons via a clathrin-dependent pathway and through pre-existing clathrin coated pits. The role of clathrin-mediated endocytosis in signal transduction has yet to be fully understood. It is known that m2R, as an autoreceptor, modulates acetylcholine release in hippocampus and cortex (Zhang et al., 2002). How does m2R endocytosis alters acetylcholine release and is clathrin-mediated endocytosis involved in this alteration are still open questions. It is likely that clathrin-mediated endocytosis plays a key role in the regulation of signal transduction by physically removing activated m2R from the cell surface, that would have as a consequence to terminate the signal. Unless m2Rs recycle from endosomes. Another role of clathrin-mediated endocytosis may be to produce transport vesicle to convey m2R to axonal varicosities where it is involved in the regulation of acetylcholine release. This phenomenon called transcytosis has been reported for Trk receptors (Ascano et al. 2009) but never for a muscarinic receptor. Analysis of m2R redistribution at the axonal levels may help to consider this hypothesis. If our work demonstrates that m2R is internalized through clathrin-mediated endocytosis, we cannot exclude that another endocytotic pathway contributes to m2R internalization.

The regulation of m2R membrane availability may thus contribute to regulate neuronal sensitivity to acetylcholine and relative drugs in physiological or pathological conditions displaying abnormalities in acetylcholine transmission such as Alzheimer’s disease or schizophrenia (Wess et al., 2007).
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AUTHOR CONTRIBUTIONS STATEMENT

LL has performed neuronal cultures, immunocytochemistry and imaging. DD participated in immunocytochemistry experiments and analysis. AF participated to neuronal cultures. ZC and PD contributed to data interpretations. VB conceptualized the research, designed the project, participated in the analysis and data interpretation and drafted the work.

CONFLICT OF INTEREST STATEMENT

The submitted work was not carried out in the presence of any personal, professional or financial relationships that could potentially be construed as a conflict of interest.
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FIGURE LEGENDS

Figure 1
Validation of GFP-m2R, SEP-m2R and WT-m2R expressing vectors and localization of transfected m2R in hippocampal neurons. Hippocampal neurons were transfected with a plasmid encoding the GFP-tagged receptor (A-C’’), SEP-tagged receptor (D-G) or wild-type receptor (WT-SS-m2R: H-J), fixed, and processed for visualization of the receptor by confocal microscopy. Equatorial images of neurons (0.5µM in depth) were selected and illustrated on this panel. The m2R localization was identified by GFP (A,A’’,B,B’’,C,C’’), or SEP (D,D’’E,F,G) native fluorescence or by fluorescent ICC using an anti-m2R (A’,B’,D’,H’-J) or anti-GFP antibody (C’,G). Whatever the construction, the fluorescent signal is localized at the membrane of the soma and the dendrites. A faint signal is detected in the cytoplasm. B,F,I. The stimulation with a muscarinic receptor agonist (Carbachol (CCh), 100µM) induces a huge decrease of the signal at the somatic and dendritic membranes and a the appearence of a punctiform labeling in the cytoplasm when using a GFP (B) or SEP-tagged (F) or WT (I) construction. J: A neuron, that has been pre-incubated with Atropine (10nM), a muscarinic receptor antagonist, display a membrane labeling at the soma and dendrites similar to a control staining. A-B’’,D-D’’: Fluorescent signals detected by direct visualisation of GFP (A,B) or SEP (D) and by m2R ICC (A’,B’,D’) in a same neuron perfectly colocalize (A’’,B’’,D’’). C-C’’: GFP or SEP detection by ICC (C’,G) display a membrane labeling in a non-permeabilized neuron that colocalizes with the direct GFP or SEP fluorescence (C,C’’,G).

Figure 2
Validation of pH-dependence of SEP-m2R. A living hippocampal neuron transfected with SEP-m2R was observed by spinning disk confocal microscopy for 30min. A stack of 20 images (0.5µM in depth) were collected at 30s intervals. A projection of the stack images was performed and an equatorial image was extracted (insert) and illustrated on this panel (A-D). The effect of pH was observed on the fluorescence level with or without NH4Cl. A: At pH7.4, the SEP-m2R is detected at the membrane of cell body and proximal dendrites. B: At acidic pH, SEP-m2R labeling strongly decreases. C: The SEP-m2R labeling is seen again at the plasma membrane when the medium is back to pH7.4. When NH4Cl (50mM) that is known to reveal receptors associated with acidic intraneuronal organelles is added in the medium, punctiform m2R labeling was also seen in the cytoplasm. D: At acidic pH with NH4Cl, the SEP-m2R labeling is very weak again. E: At pH7.4, the SEP-m2R is detected again. F: Quantification of the fluorescence level. Fluorescence was measured at the level of the whole neuron and in soma using the Fiji software. Data are expressed as normalized values compared to the fluorescence level at pH7.4 at 0min. SEP-m2R labeling strongly decreases at acidic pH at plasma membranes and in the cytoplasm. NH4Cl at pH7.4 induces an increase of the staining close to the control values at the plasma membranes and much higher in soma. The acidic pH with NH4Cl strongly decreases fluorescence at membranes and in soma. The recovery of SEP-m2R fluorescence is shown in both compartments when the medium is back to pH7.4.

Figure 3
Time lapse imaging and quantification of SEP-m2R membrane labeling in a living neuron after stimulation by CCh, a muscarinic receptor agonist. A living hippocampal neuron transfected with SEP-m2R was observed by spinning disk confocal microscopy for 75min. A stack of 20 images were collected at 30s intervals. An equatorial image (0.5µM in depth) was selected and illustrated on this panel (A-C). A: In control condition, the SEP-
m2R staining is detected at the membrane of the cell body and proximal dendrites. A faint signal is also shown in the cytoplasm. B : CCh (100µM) induces a decrease of SEP-m2R labeling at cell body and dendrites levels. C : Application of NH4Cl (50mM) that reveals receptors associated with acidic intraneuronal organelles induces an abundant and intense punctiform staining in the cytoplasm. D : Quantification of the effect of CCh on the fluorescence level +/- SEM in 4 neurons using the Fiji software. Fluorescence was measured on three different neurons on a projection of the stack images at the level of the whole neuron using the Fiji software. Data are expressed as normalized values compared to the fluorescence level at 9 min before CCh application. The quantification shows a significant difference of m2R fluorescence with time. The statistical analysis (Repeated measures ANOVA test followed by the Dunnett post-hoc test), performed on raw data, shows that CCh induces a significative decrease of fluorescence 39, 54 and 69 min after the beginning of the treatment. Post hoc anlayses were performed on two segments of the slope to analyse 1) the effect of CCh (from T=0min until 84 min) and 2) the effect of NH4Cl (from T=84 min until 120 min) on fluorescence levels. The values are compared to the values at T=9 min, the initiation point of CCh application for the CCh effect and at T=84 min, the initiation of NH4Cl application, for NH4Cl effect. Results show a significant decrease of the fluorescent level from 39 to 54 min after CCh stimulation. From 54 min, fluorescence slowly returns to normal values. In contrast, NH4Cl, which reveals m2R attached to acidic vesicles, induces a significant increase of SEP-m2R fluorescence levels. NS : not significant, * : p<0.05 ; ** : p<0.001 ; ***: p<0.0001.

**Figure 4**

**Time lapse imaging and quantification of internalization of m2R in a living neuron after stimulation by CCh, a muscarinic receptor agonist.** A living hippocampal neuron transfected with GFP-m2R was observed by spinning disk confocal microscopy before and during a 30-min-long carbachol (100µM) treatment. A stack of 20 consecutive confocal images (0.5µM in depth) were acquired every 30 s. A projection of the stack images was performed and illustrated on this panel (A-F). An equatorial image was selected and an enlargement of a dendritic shaft is shown at the bottom of each image (insert). Here is shown the GFP-m2R labeling in this neuron 3 min before CCh and every 3 min for 30 min. Before CCh addition (A) and 3 min after the beginning of agonist treatment (B), m2R was detected mainly at the plasma membrane of soma and dendrites. Agonist induces internalization of membrane-associated m2R and clusterization 6 min after treatment initiation in the cytoplasm of soma and dendrites (arrows in C). G : Quantification of the effect of CCh on the density of fluorescent clusters +/- SEM in the cytoplasm of 3 neurons. Fluorescence was measured on a projection of the stack images using the Fiji software. The clusters density increases during the first 30 min and stabilized afterwards. The statistical analysis (Repeated measures ANOVA test followed by the Dunnett post-hoc test) shows a significant increase of the number of clusters as early as 10 min after the beginning of CCh stimulation.

**Figure 5**

**Internalization of m2R in native clathrin-coated pits in neurons after stimulation by CCh.** Hippocampal neurons were transfected with a plasmid encoding the WT-m2R, stimulated with CCh (30µM) for 9 min and fixed. A stack of 10 images were collected and an equatorial image was selected and an enlargement of a cell body is shown at the bottom of each image (insert). (A-B’’). (A-B’’) : Native CCP where detected by immunocytochemistry using an anti-clathrin heavy chain antibody (A’,B’). A-A’’ : A
control neuron displays no m2R and clathrin colocalization. Nine minutes after the initiation of CCh treatment, some m2R clusters colocalize with native CCP (arrows). C : The analysis of the colocalization of clathrin and m2R-ICC was performed on fixed neurons using the Jacop Plugin of ImageJ are reported from the Costes’s randomization-based colocalization module (see Materials and methods). The quantification of m2R and clathrin colocalization and the statistical analysis demonstrates a significant increase of the Pearson’s coefficient in treated neurons (n=20) compared to cells treated with CCh (n=20) (Mann Whitney U test; * : p<0.0001).). Control neurons : n=20; CCh-treated neurons : n=21.

Figure 6 :
Internalization of m2R in clathrin-coated pits in a living neuron after stimulation by CCh. A living hippocampal neuron transfected with GFP-m2R and Ds-Red clathrin was observed by spinning disk confocal microscopy before and during a 30-min-long carbachol (30µM) treatment. A stack of 10 images was collected and selected images at the indicated times show a representative dendrite and a selected area in a cell body (insert) of a neuron. Six minutes after the beginning of CCh treatment, m2R clusters appear at loci of clathrin-coated-pit (CCP) spots (arrows) in the dendrite and the cell body (A’’-E’’). Note that the m2R clusters form in preexisting CCP.

Figure 7
Amount of Tf uptake and m2R clathrin-dependent endocytosis in hippocampal neurons. A- D’ : Hippocampal neurons were transfected with m2R-WT, pre-incubated with Tf-Al594, 10 min before CCh treatment. Cells were fixed after 15min treatment. In control neurons, Al594-Tf is detected in the cytoplasm as a punctiform labelling (A’-A’’). CCh treatment induces a strong decrease of membrane m2R labeling and the appearance of m2R punctiform staining (B’,B’’). Al594-Tf and m2R-ICC signal often colocalize (B,B’, arrows). The quantitative analysis of the colocalization of m2R and Al594-Tf in neurons was performed using the Jacop Plugin of ImageJ and statistical data are reported from the Costes’s randomization-based colocalization module (see methods). Data are expressed as a Pearson’s coefficient (pc) and pc were compared using the Kruskal-Wallis test followed by the Dunn’s Multiple Comparison Test. Our analysis shows that the colocalization observed between the m2R immunofluorescent signal and Al594-Tf is higher after treatment with CCh compared to untreated neurons (***(p<0.0001)). Atropine prevents the increase of m2R and Al594-Tf colocalization (Atropine treatment vs Control : NS : not significant). Control neurons : n=25; CCh-treated neurons : n=19; CCh-treated + atropine neurons: n=9.

Figure 8
Blockade of m2R clathrin-dependent endocytosis in hippocampal neurons with negative dominant of Eps15. Hippocampal living neurons were co-transfected with WT-m2R plasmid and GFP-tagged EH29 and DIII mutants or their control (D3Δ2). The day after transfection, neurons were treated with CCh (30µM) for 15min and fixed. The m2R was detected by ICC. When the mutants are expressed (labeling in A’, B’), m2R labeling is seen at the plasma membrane of neurons (A,B). The expression of the control plasmid (labeling in C’) does not block m2R internalization (C). D : Intracellular immunofluorescent clusters, representing m2R in endosomes, were segmented and counted using the FIJI/ImageJ software. Results are expressed as intracellular immunofluorescent clusters per μm² cytoplasmic surface in Eps15 dominant negative-treated and control neurons. The statistical analysis shows that the expression of the EH29 and DIII mutants blocks m2R clusterization (Mann-Whitney U test: NS : Not significant; EH29 : Control neurons : n=19; CCh-treated
neurons: n=19; DIII: Control neurons: n=19; CCh-treated neurons: n=30). In contrast, the control mutant (D3Δ2) does not inhibit m2R clusterization (Mann-Whitney U test: p < 0.0001; Control neurons: n=19; CCh-treated neurons: n=15).

Figure 9
Absence of internalization of m2R in caveole in fixed neuron after stimulation by CCh.

Hippocampal neurons were co-transfected with a plasmid encoding the wild-type receptor (WT-SS-m2R: A,B) and CAV1-mCherry (A’,B’) fixed, and processed for visualization by confocal microscopy. In control and treated neurons, CAV1-mCherry is detected in the cytoplasm as a punctiform labelling (A’,A’’, B,B’’). Some m2R and CAV1-mCherry clusters colocalize (arrows) in both treated and untreated neurons. The quantitative analysis of the colocalization of m2R and CAV1-mCherry in neurons was performed using the Jacop Plugin of ImageJ and statistical data are reported from the Costes’s randomization-based colocalization module (see methods). Data are expressed as a Pearson’s coefficient (pc) and pc were compared using the Kruskal-Wallis test followed by the Dunn’s Multiple Comparison Test. Our analysis shows that pc values do not significantly differ in control neurons and neurons treated with CCh for 6, 12 and 15 min (NS: not significant).

Figure 10
Immunohistochemical localization of m2R in neuronal compartments involved in endocytosis, synthesis, maturation and degradation in fixed hippocampal neurons.

Hippocampal neurons were transfected with m2R-WT. Neurons were stimulated with CCh at 30µM for 6, 20 min and 1hr fixed, and processed for visualization of m2R together with markers of intraneuronal compartments and observed by confocal microscopy. A-C’’: 6min after CCh stimulation (30µM), some m2R immunopositive puncta colocalize with CHC in clathrin-coated pits, EEA1 in early endosomes and Rab9 in late endosomes (arrow heads).

D-F’’: 20min after CCh stimulation (30µM), we failed to detect no colocalization of m2R with PDI, a marker of endoplasmic reticulum and GM130 and TGN38, markers of Golgi apparatus. G-G’’: 1hr after CCh stimulation (30µM), some m2R immunopositive puncta colocalize with CathD, a marker of lysosomes (arrow heads). The quantitative analysis of the colocalization of m2R and markers of subcellular compartment in neurons was performed using the Jacop Plugin of ImageJ and statistical data are reported from the Costes’s randomization-based colocalization module (see methods). Data are expressed as a Pearson’s coefficient (pc) and pc were compared using the Mann-Whitney U test. Our analysis shows that the colocalization of the immunofluorescent signals for m2R with CHC, EEA1, Rab9 and CathD is higher after treatment with CCh compared to untreated neurons (CHC, Rab9 and CathD: ***: p<0.0001; EEA1: **: p<0.01). In contrast, the colocalization of the immunofluorescent signals for m2R with PDI and GM130 do not significantly differ in CCh-treated neurons compared to untreated Control neurons: CHC n=20, EEA1 n=16, Rab9 n=15, PDI n=21, GM130 n=17, CathD n=17; CCh-treated neurons: CHC n=12, EEA1 n=18, Rab9 n=15, PDI n=20, GM130 n=18, CathD n=15.