Characterization of central inhibitory muscarinic autoreceptors by the use of muscarinic acetylcholine receptor knock-out mice

Weilie Zhang, National Institute of Diabetes and Digestive and Kidney Diseases
Anthony S. Basile, National Institute of Diabetes and Digestive and Kidney Diseases
Jesus Gomeza, National Institute of Diabetes and Digestive and Kidney Diseases
Laura A. Volpicelli, Emory University
Allan I Levey, Emory University
Jurgen Wess, National Institute of Diabetes and Digestive and Kidney Diseases

Journal Title: Journal of Neuroscience Nursing
Volume: Volume 22, Number 5
Publisher: Lippincott, Williams & Wilkins | 2002-03-01, Pages 1709-1717
Type of Work: Article | Final Publisher PDF
Publisher DOI: 10.1523/JNEUROSCI.22-05-01709.2002
Permanent URL: https://pid.emory.edu/ark:/25593/vdvdx

Final published version:
http://dx.doi.org/10.1523/JNEUROSCI.22-05-01709.2002

Copyright information:
Copyright © 2002 Society for Neuroscience
This is an Open Access work distributed under the terms of the Creative Commons Attribution 4.0 International License (https://creativecommons.org/licenses/by/4.0/).

Accessed August 9, 2020 12:02 AM EDT
Characterization of Central Inhibitory Muscarinic Autoreceptors by the Use of Muscarinic Acetylcholine Receptor Knock-Out Mice

Weile Zhang,1 Anthony S. Basile,1 Jesus Gomez,1 Laura A. Volpicelli,2 Allan I. Levey,2 and Jürgen Wess1

1Laboratory of Bioorganic Chemistry, National Institute of Diabetes and Digestive and Kidney Diseases, Bethesda, Maryland 20892, and 2Department of Neurology, Emory University School of Medicine, Atlanta, Georgia 30322

Forebrain muscarinic acetylcholine (ACh) receptors (mAChRs; M1–M5) are predicted to play important roles in many fundamental central functions, including higher cognitive processes and modulation of extrapyramidal motor activity. Synaptic ACh levels are known to be regulated by the activity of presynaptic muscarinic autoreceptors mediating inhibition of ACh release. Primarily because of the lack of ligands with pronounced subtype selectivity, classical pharmacological studies have led to conflicting results regarding the identity of the mAChR subtypes mediating this activity in different areas of the brain. To investigate the molecular identity of hippocampal, cortical, and striatal inhibitory muscarinic autoreceptors in a more direct manner, we used genetically altered mice lacking functional M2 and/or M4 mAChRs [knock-out (KO) mice]. After labeling of cellular ACh pools with [3H]choline, potassium-stimulated [3H]ACh release was measured in superfused brain slices, either in the absence or the presence of muscarinic drugs. The nonsubtype-selective muscarinic agonist, oxotremorine (0.1–10 μM), inhibited potassium-stimulated [3H]ACh release in hippocampal, cortical, and striatal slices prepared from wild-type mice by up to 80%. This activity was totally abolished in tissues prepared from M2–M4 receptor double KO mice. Strikingly, release studies with brain slices from M2 and M4 receptor single KO mice indicated that autoinhibition of ACh release is mediated primarily by the M4 receptor in hippocampus and cerebral cortex, but predominantly by the M2 receptor in the striatum. These results, together with additional receptor localization studies, support the novel concept that autoinhibition of ACh release involves different mAChRs in different regions of the brain.

Key words: acetylcholine release; autoreceptors; knock-out mice; muscarinic receptors; oxotremorine; presynaptic receptors

Many of the important central actions of acetylcholine (ACh) are mediated by members of the muscarinic ACh receptor family (M1–M5) that belong to the superfamily of G-protein-coupled receptors (Huile et al., 1990; Wess, 1996). Central mAChRs are involved in a large number of important physiological functions, including the control of extrapyramidal locomotor activity (Di Chiara et al., 1994) and higher cognitive processes such as learning and memory (Coyle et al., 1983; Mash et al., 1985; Quirion et al., 1984). Identification of the mAChR subtypes involved in these diverse central muscarinic functions has proven a difficult task, primarily because of the lack of ligands with pronounced subtype selectivity (Buckley et al., 1989; Dörje et al., 1991; Caulfield, 1993; Wess, 1996) and the fact that most central tissues express multiple mAChRs (Levey, 1993; Vilaro et al., 1993).

As is the case with many other neurotransmitter systems, synaptic levels of ACh are known to be regulated by the activity of presynaptic mAChRs mediating feedback inhibition of ACh release from cholinergic nerve terminals (for review, see Kilbinger, 1984; Starke et al., 1989). In situ mRNA hybridization studies have shown that all five mAChRs are expressed in areas of the CNS known to contain cholinergic cell bodies, raising the possibility that multiple mAChRs participate in presynaptic modulation of ACh release (Vilaro et al., 1994). Functional studies using classical pharmacological tools (muscarinic agonists and antagonists) have led to conflicting results regarding the molecular nature of the mAChRs mediating autoinhibition of ACh release. For example, inhibitory muscarinic autoreceptors in the hippocampus have been classified as either M2 (Pohorecky et al., 1988; Lapchak et al., 1989), M3 (Raiteri et al., 1989), or M4 (McKinney et al., 1993) receptors, or as mixtures of M1 and M4 mAChRs (Vannucchi and Pepeu, 1995). Similarly, it has been proposed that striatal muscarinic autoreceptors predominantly consist of either M1 (Kawashima et al., 1991), M2 (James and Cubeddu, 1987; Lapchak et al., 1989; Weiler, 1989; Billard et al., 1995), M3 (De Boer et al., 1990; Büyüküysal et al., 1998), or M4 (Dolezal and Tucek, 1998) receptors. It is likely that these discrepant results are primarily caused by the limited subtype selectivity of the pharmacological agents used in these studies (Buckley et al., 1989; Dörje et al., 1991; Caulfield, 1993).

To circumvent these difficulties, we and others recently developed mutant mouse lines in which specific mAChR genes had been inactivated via gene-targeting techniques (Hamilton et al., 1997; Gomez et al., 1999a,b; Yamada et al., 2001). Using these mice, we performed systematic studies of ACh release from brain slices, to examine the molecular nature of central muscarinic autoreceptors in a more direct manner. Specifically, we studied muscarinic agonist-mediated inhibition of potassium-stimulated [3H]ACh re-
lease using superfused hippocampal, cortical, and striatal tissues from M<sub>2</sub> and M<sub>4</sub> receptor single knock-out (KO) mice (Gomez et al., 1999a,b) and M<sub>2</sub>–M<sub>4</sub> receptor double KO mice (A. Duttaroy, J. Gomeza, and J. Wess, unpublished observations).

MATERIALS AND METHODS

Animals. The generation of homozygous M<sub>2</sub> receptor KO (genetic background: 129J1/H11003) and M<sub>4</sub> receptor KO (129J1 × 129Ev × EV) mice has been described previously (Gomez et al., 1999a,b). M<sub>2</sub>–M<sub>4</sub> receptor double KO mice (129J1 × 129Ev × EV) were generated by intermixing homozygous M<sub>2</sub> and M<sub>4</sub> receptor mutant mice (A. Duttaroy, J. Gomeza, and J. Wess, unpublished observations). In all experiments, aged-matched wild-type (WT) mice of the matching genetic background were used as control animals. Mouse genotyping was performed by PCR analysis of mouse tail DNA.

Acetylcholine release studies. Mice (males; 2–4 months old) were killed by decapitation, and brains were removed. Hippocampal, cortices, and striata were dissected and chopped into 250 μm prisms using a Sorvall tissue chopper (Newton, CT). Hippocampal, cortical, and striatal slices prepared from one mouse were dispersed in 25 ml of oxygenated (95% O<sub>2</sub> and 5% CO<sub>2</sub>) Krebs’–Ringer’s solution buffer, pH 7.4 (in mM: 11.5 glucose, 25 NaHCO<sub>3</sub>, 1.2 MgCl<sub>2</sub>, 1.2 NaH<sub>2</sub>PO<sub>4</sub>, 118 NaCl, 4.8 KCl, 2.5 CaCl<sub>2</sub>, and 0.004 Na<sub>2</sub>-EDTA) at 33°C for 20 min. Slices were then incubated with [3H]choline (75 Ci/mmol; NEN Life Science Products, Boston, MA) at a final concentration of 0.1 μM for 30 min. This low concentration of [3H]choline favors the selective uptake of choline only into cholinergic neurons through their high-affinity uptake system (Pittel et al., 1990). After rinsing, slices were transferred to a superfusion system (SF-12; Brandel, Gaithersburg, MD) and superfused at 33°C at a constant rate of 0.4 ml/min with Krebs’–Ringer’s solution buffer containing 10 μM hemicholinium to prevent uptake of [3H]choline during the release experiments. Fractions were collected every 4 min beginning after a 60 min superfusion. Two 2 min periods of 25, 30, or 20 mM KCl (hippocampus, cerebral cortex, and striatum, respectively) were applied after 72 (S1) and 104 (S2) min of superfusion. Tetrodotoxin (600 nM) (Sigma, St. Louis, MO) was added at the beginning of the superfusion period where indicated. Drugs were added to the superfusion buffer 20 min before S2. The efflux of tritium collected was calculated as a percentage of the total tritium present in the slices at the start of the fraction considered. The net efflux of tritium was calculated by subtracting the average of three fractions (expected basal value) before KCl stimulation. The results were expressed as the S2:S1 ratio of release.

Separation of [3H]choline and [3H]ACh by reverse-phase HPLC. Hippocampal, cortical, and striatal slices prepared from M<sub>2</sub> receptor WT mice were incubated with [3H]choline (75 Ci/mmol), superfused, and stimulated with KCl as described in the previous paragraph. To prevent enzymatic degradation of released ACh, phostigmine (100 μM) was added to the superfusion medium. Fractions were collected every 4 min (rate of superfusion: 0.4 ml/min), as described above. [3H]Choline and [3H]ACh were separated by reverse-phase HPLC followed by liquid scintillation spectrometry, essentially as described (Wessler and Wetard, 1990). Aliquots of each fraction (100 μl) were injected onto the HPLC (ESA Inc., Chelmsford, MA; S82; reverse phase C18 5 μm, 250 × 3.2 mm). The elution solvent consisted of 0.1 mM phosphate buffer, pH 4.7, that contained methanol (8% vol) and 0.2 mM tetramethylammonium. The flow rate was 0.5 ml/min, and the effluent was collected in 1 min fractions. The retention times of choline and ACh were determined using the radioactive standards [3H]choline and [3H]ACh (20 Ci/mmol; ARC, St. Louis, MO) (Fig. 1A).

Muscarinic receptor–vesicular acetylcholine transporter colocalization studies. Mice were injected with heparin, anesthetized with sodium pentobarbital, and perfused transcardially with 15–20 ml 0.9% NaCl and 0.005% sodium nitrite, followed by 80–120 ml of 4% paraformaldehyde in 0.1 M phosphate buffer. The brain was removed and placed in 4% paraformaldehyde overnight at 4°C. Sections (∼50-μm-thick) of the striatum were cut on a vibratome and rinsed several times in PBS. All solutions were diluted in PBS, and all incubations were performed at room temperature with gentle agitation. The sections were treated with 3% hydrogen peroxide for 10 min, followed by three rinses in PBS. The sections were blocked in 5% normal goat serum, 5% normal horse serum, and 10 μg/ml avidin for 30 min. Incubations with primary antibodies were performed in buffer containing 1% normal goat serum, 1% normal horse serum, and 50 μg/ml biotin. The following antibodies were used (Levey et al., 1991; Hersch et al., 1994; Gilmore et al., 1996; Bernard et al., 1998, 1999): M<sub>3</sub> rat monoclonal antibody (1:100), M<sub>4</sub> mouse monoclonal antibody (1:1000), and vesicular acetylcholine transporter (VACHT) rabbit polyclonal antibody (1 μg/ml). The specificity of these antibodies has been described in detail previously (Levey et al., 1991; Hersch et al., 1994; Gilmore et al., 1996; Bernard et al., 1998, 1999). For muscarinic receptor–VACHT double-labeling studies, striatal slices were coincubated with the two primary antibodies. The sections were then rinsed and incubated for 60 min with donkey anti-rabbit rhodamine red-X (1:100; Jackson ImmunoResearch, West Grove, PA) in secondary buffer (1% normal goat serum and 1% normal horse serum). Subsequently, sections were rinsed and incubated with biotinylated goat anti-mouse or anti-rat antibody (1:100; Jackson ImmunoResearch) in secondary buffer for 60 min. After several rinses, the sections were incubated with avidin–biotin complex (ABC elite; Vector Laboratories, Burlingame, CA) for 60 min, rinsed, and incubated in tyramide-fluorescein diluted in amplification diluent (1:100; NEN) for 10 min. The sections were rinsed and incubated for 30 min in 10 mM cupric sulfate in 50 mM ammonium acetate, pH 5.0, to eliminate autofluorescence (Schell et al., 1999). After rinsing, sections were mounted using Vectorshield mounting media for fluorescence (Vector Laboratories). Control incubations included omission of the primary antibodies to test for nonspecific binding of the secondary antibodies and incubation with one primary but both secondary antibodies to demonstrate the absence of bleedthrough and cross-labeling.

Sections were scanned using a Zeiss (Oberkochen, Germany) LSM 510 laser-scanning confocal microscope coupled to a Zeiss 100M axiovert and a 63× Plan-Apochromat oil-immersion lens. Quantitation of colocalization between M<sub>2</sub> or M<sub>4</sub> muscarinic receptors and VACHT was analyzed using Metamorph Imaging System Software (Universal Imaging Corporation, West Chester, PA). To define background, sections incubated with both secondary but no primary antibodies were used, and the average grayscale pixel intensity + 1 SD was measured in the FITC and rhodamine channels. To subtract background from double-labeled images, the threshold of each channel was set at the value obtained for background. The average pixel intensity + 1 SD was measured, and the threshold was set to this new value. The percentage of the area of overlap between M<sub>2</sub> or M<sub>4</sub> pixels over VAChT pixels was measured. Colocalization was assessed on two randomly chosen fields of the striatum from three different animals per genotype. Data are presented as means ± SEM.

Figures. All statistics were analyzed by one-way repeated measures ANOVA followed by a Student–Newman–Keuls test. Data are given as means ± SEM.
RESULTS

Muscarinic agonist-mediated inhibition of stimulated ACh release is abolished in brain slices from M₂–M₄ receptor double KO mice

The M₂ and M₄ mAChRs are selectively coupled to G-proteins of the G₁/G₅ family, whereas the M₁, M₃, and M₅ mAChRs are preferentially linked to G-proteins of the G₉ subclass (Hulme et al., 1990; Caulfield, 1993; Wess, 1996). Because mAChR-activated G₁/G₅ proteins mediate the inhibition of voltage-sensitive Ca²⁺ channels (Caulfield, 1993; Shapiro et al., 1999) that are known to be intimately involved in the regulation of neurotransmitter release, we speculated that the M₂ and/or M₄ receptor subtypes represent the major inhibitory muscarinic autoreceptors. To test this concept in a direct and unambiguous manner, we initially analyzed ACh release using mutant mice lacking both M₄ and M₄ mAChRs (M₂–M₄ double KO mice) (Duttaroy, Gomez and Wess, unpublished observations). The M₂–M₄ receptor double KO mice showed no obvious morphological abnormalities and did not differ from their WT littermates in overall health, fertility, and longevity. Moreover, immunoprecipitation studies with receptor subtype-selective antisera indicated that the lack of M₂ and M₄ receptors did not lead to compensatory changes in the levels of the remaining mAChR subtypes (Duttaroy, Gomez, and Wess, unpublished observations).

In vitro ACh release studies were performed with superfused hippocampal, cortical, and striatal slices prepared from WT and mAChR mutant mice. Initially, cellular ACh pools were radioactively labeled by incubating brain tissues with [³H]choline. Subsequently, potassium-stimulated [³H] release was measured either in the absence (S1 phase) or the presence of drugs (S2 phase), as described in Materials and Methods.

To verify that the potassium-dependent [³H] outflow in the three different tissues primarily consisted of [³H]ACh, we used a reverse-phase HPLC method that can efficiently separate [³H]choline and [³H]ACh with a recovery rate of ~100% (Wessler and Werhand, 1990). To prevent enzymatic degradation of [³H]ACh, physostigmine (100 μM) was added to the perfusion medium in this set of experiments. Control experiments with [³H]choline and [¹⁴C]ACh standards indicated that both amines could be clearly separated with virtually no overlap (Fig. 1A). In hippocampal, cortical, and striatal preparations from WT mice (M₄ receptor WT mice), the basal (spontaneous) [³H] efflux consisted predominantly of [³H]ACh in all three tissues investigated (88–96% of total net [³H] outflow) (Table 1). The [³H] recovery rate was ~95% in all three tissues investigated (data not shown). When calcium was omitted from the superfusion medium, the KCl-induced increase in [³H]ACh release was virtually abolished in all three tissues studied (<10% of [³H]ACh release observed in the presence of calcium; n = 4). Consistent with previous studies (Pedata et al., 1986; Marchi et al., 1990), these data indicate that the potassium-stimulated [³H] outflow in slices from different central tissues preincubated with [³H]choline predominantly represents authentic [³H]ACh (shown for striatal slices in Fig. 1B). Throughout the text, we therefore refer to potassium-stimulated [³H] outflow simply as [³H]ACh release.

Incubation of hippocampal, cortical, and striatal slices from WT mice with oxotremorine (0.1–10 μM), a nonsubtype-selective muscarinic agonist, led to a dose-dependent inhibition of stimulated [³H]ACh release (Figs. 2A, 3A, 4A, top panels). At the highest oxotremorine concentration used (10 μM), the average inhibition of [³H]ACh release amounted to 73 ± 2% in hippocampal, 74 ± 4% in cortical, and 56 ± 2% in striatal preparations, respectively. The oxotremorine (10 μM)-mediated inhibition of transmitter release was completely abolished in the presence of atropine (2 μM), confirming the involvement of mAChRs (Figs. 2A, 3A, 4A). Incubation of tissue slices with atropine (2 μM) alone had no significant effect on potassium-evoked [³H]ACh release (Figs. 2A, 3A, 4A), suggesting that the

Table 1. Basal and potassium-evoked release of [³H]choline and [³H]ACh in hippocampal, cortical, and striatal slices from WT mice prelabeled with [³H]choline, as determined by reverse-phase HPLC

<table>
<thead>
<tr>
<th>Tissue</th>
<th>[³H]Choline (Disintegrations per minute/preparation)</th>
<th>[³H]ACh (Disintegrations per minute/preparation)</th>
<th>% [³H]ACh (of total [³H])</th>
</tr>
</thead>
<tbody>
<tr>
<td>Hippocampus</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Basal release</td>
<td>957 ± 60</td>
<td>1119 ± 48</td>
<td>52.9 ± 2.1</td>
</tr>
<tr>
<td>Release after K⁺ stimulation</td>
<td>1092 ± 56</td>
<td>2516 ± 109</td>
<td>69.7 ± 1.1</td>
</tr>
<tr>
<td>Net increase in release after K⁺ stimulation</td>
<td>180 ± 27</td>
<td>1442 ± 104</td>
<td>88.9 ± 1.5</td>
</tr>
<tr>
<td>Cortex</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Basal release</td>
<td>959 ± 10</td>
<td>944 ± 64</td>
<td>49.5 ± 1.5</td>
</tr>
<tr>
<td>Release after K⁺ stimulation</td>
<td>1166 ± 30</td>
<td>2285 ± 135</td>
<td>66.1 ± 1.6</td>
</tr>
<tr>
<td>Net increase in release after K⁺ stimulation</td>
<td>208 ± 37</td>
<td>1491 ± 159</td>
<td>87.7 ± 2.0</td>
</tr>
<tr>
<td>Striatum</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Basal release</td>
<td>1087 ± 140</td>
<td>1940 ± 182</td>
<td>64.3 ± 1.4</td>
</tr>
<tr>
<td>Release after K⁺ stimulation</td>
<td>1254 ± 164</td>
<td>5421 ± 383</td>
<td>81.3 ± 1.7</td>
</tr>
<tr>
<td>Net increase in release after K⁺ stimulation</td>
<td>167 ± 33</td>
<td>3481 ± 298</td>
<td>95.5 ± 0.7</td>
</tr>
</tbody>
</table>

*Sum of disintegrations per minute of aliquots (100 μl) taken from the three fractions collected immediately before K⁺ stimulation (fractions 2–4).

Table 1. Basal and potassium-evoked release of [³H]choline and [³H]ACh in hippocampal, cortical, and striatal slices from WT mice prelabeled with [³H]choline, as determined by reverse-phase HPLC

- Basal release* 957 ± 60 1119 ± 48 52.9 ± 2.1
- Release after K⁺ stimulation 1092 ± 56 2516 ± 109 69.7 ± 1.1
- Net increase in release after K⁺ stimulation 180 ± 27 1442 ± 104 88.9 ± 1.5
- Basal release 959 ± 10 944 ± 64 49.5 ± 1.5
- Release after K⁺ stimulation 1166 ± 30 2285 ± 135 66.1 ± 1.6
- Net increase in release after K⁺ stimulation 208 ± 37 1491 ± 159 87.7 ± 2.0
- Basal release 1087 ± 140 1940 ± 182 64.3 ± 1.4
- Release after K⁺ stimulation 1254 ± 164 5421 ± 383 81.3 ± 1.7
- Net increase in release after K⁺ stimulation 167 ± 33 3481 ± 298 95.5 ± 0.7

*Sum of disintegrations per minute of aliquots (100 μl) taken from the three fractions collected immediately after K⁺ stimulation (fractions 5–7).

ACh release was virtually abolished in all three tissues studied (0–10% of [³H]ACh in all three tissues investigated (88–96% of total net [³H] outflow) (Table 1). The [³H] recovery rate was ~95% in all three tissues investigated (data not shown). When calcium was omitted from the superfusion medium, the KCl-induced increase in [³H]ACh release was virtually abolished in all three tissues studied (10% of [³H]ACh release observed in the presence of calcium; n = 4). Consistent with previous studies (Pedata et al., 1986; Marchi et al., 1990), these data indicate that the potassium-stimulated [³H] outflow in slices from different central tissues preincubated with [³H]choline predominantly represents authentic [³H]ACh (shown for striatal slices in Fig. 1B). Throughout the text, we therefore refer to potassium-stimulated [³H] outflow simply as [³H]ACh release.

Incubation of hippocampal, cortical, and striatal slices from WT mice with oxotremorine (0.1–10 μM), a nonsubtype-selective muscarinic agonist, led to a dose-dependent inhibition of stimulated [³H]ACh release (Figs. 2A, 3A, 4A, top panels). At the highest oxotremorine concentration used (10 μM), the average inhibition of [³H]ACh release amounted to 73 ± 2% in hippocampal, 74 ± 4% in cortical, and 56 ± 2% in striatal preparations, respectively. The oxotremorine (10 μM)-mediated inhibition of transmitter release was completely abolished in the presence of atropine (2 μM), confirming the involvement of mAChRs (Figs. 2A, 3A, 4A). Incubation of tissue slices with atropine (2 μM) alone had no significant effect on potassium-evoked [³H]ACh release (Figs. 2A, 3A, 4A), suggesting that the
The biophase concentration of ACh did not reach levels that were sufficiently high to produce autoinhibition of ACh release. It is likely that degradation of released ACh by cholinesterases (all experiments were carried out in the absence of cholinesterase inhibitors) and removal of ACh by the superfusion stream were the major factors leading to the rapid reduction of synaptic ACh levels (for review, see Starke et al., 1989).

In all three tissues examined, K⁺-evoked [³H]ACh release and oxotremorine-induced inhibition of stimulated [³H] outflow remained essentially unaffected by incubation with tetrodotoxin (600 nm) (data not shown), suggesting that agonist-dependent inhibition of [³H]ACh release does not require the propagation of nerve impulses. As reviewed by Starke et al. (1989), it is therefore likely that autoinhibition of [³H]ACh release in central tissues is primarily mediated by mAChRs located directly on cholinergic nerve terminals. Strikingly, in hippocampal, cortical, and striatal slices prepared from M2–M4 receptor double KO mice, oxotremorine (0.1–10 μM) completely lost its ability to mediate inhibition of stimulated [³H]ACh release (Figs. 2A, 3A, 4A, bottom panels). This ob-

**Hippocampus**

Figure 2. Effect of oxotremorine on potassium-stimulated [³H]ACh release in hippocampal slices from M₂–M₄ receptor double KO, M₂ receptor single KO, and M₄ receptor single KO mice (A–C, bottom panels) and their corresponding WT control mice (A–C, top panels). Each bar represents the mean ± SEM of S2/S1 values from 6–11 independent experiments (mice). Concentrations shown are micromolar. Asterisks indicate significant differences from the control group (no drug) (*p < 0.05; **p < 0.01).

**Cortex**

Figure 3. Effect of oxotremorine on potassium-stimulated [³H]ACh release in cortical slices from M₂–M₄ receptor double KO, M₂ receptor single KO, and M₄ receptor single KO mice (A–C, bottom panels) and their corresponding WT control mice (A–C, top panels). Each bar represents the mean ± SEM of S2/S1 values from 6–11 independent experiments (mice). Concentrations shown are micromolar. Asterisks indicate significant differences from the control group (no drug) (*p < 0.05; **p < 0.01).
servation demonstrates in a very convincing manner that M2 and/or M4 receptors mediate autoinhibition of ACh release in these brain tissues.

The M2 subtype represents the predominant muscarinic autoreceptor in hippocampus and cerebral cortex

To examine the relative contributions of M2 and M4 receptors to autoinhibition of stimulated ACh release, we next performed analogous studies with hippocampal and cortical tissues from M2 receptor single KO mice and their corresponding WT controls. Interestingly, oxotremorine (0.1–10 μM) failed to exert a significant effect on stimulated [3H]ACh release in hippocampal preparations from M2 receptor KO mice (Fig. 2B). On the other hand, oxotremorine displayed comparable inhibitory effects on neurotransmitter release in hippocampal preparations from M4 receptor KO and their WT control mice (Fig. 2C). Similar observations were made when analogous studies were performed with cortical tissues (Fig. 3). As observed with hippocampal preparations, oxotremorine failed to inhibit stimulated [3H]ACh release in cortical slices from M2 receptor KO mice (Fig. 3B) but showed similar activities in cortical preparations from M4 receptor KO and their WT control mice (Fig. 3C). These findings strongly suggest that the M2 subtype represents the predominant muscarinic autoreceptor in mouse hippocampus and cerebral cortex.

The M4 subtype functions as the major muscarinic autoreceptor in striatum

In striking contrast to the findings obtained with hippocampal and cortical slices, oxotremorine (0.1–10 μM) retained the ability to mediate inhibition of stimulated [3H]ACh release in striatal preparations from mice lacking M4 receptors (Fig. 4B). In contrast, oxotremorine virtually lacked the ability to inhibit stimulated [3H]ACh release in striatal preparations from mice lacking M4 receptors (Fig. 4C). Although there was a trend to slightly reduced [3H]ACh levels at the two highest oxotremorine concentrations used (1 and 10 μM) (Fig. 4C), this effect did not reach statistical significance. These data strongly suggest that the M4 subtype represents the major muscarinic autoreceptor in mouse striatum.

Colocalization of M2 and M4 muscarinic receptors with VACHT in the striatum

It can be argued that the lack of muscarinic autoinhibition of ACh release observed with striatal preparations from M4 receptor KO mice may be caused by alterations in the distribution of M2 receptors at striatal cholinergic terminals. For example, M2 receptors may perhaps act as primary autoreceptors in M4 receptor WT mice but may no longer localize to cholinergic terminals in M4 receptor KO mice, thus providing a possible explanation for the lack of autoinhibition of ACh release observed with striatal preparations from M4 receptor KO mice. To test this hypothesis, we determined to which extent M2 receptors colocalized with the VACHT, a marker of cholinergic terminals (Gilmor et al., 1996), in the striatum of M4 receptor WT and M4 receptor KO mice. As shown in Figure 5, the M2 receptor was colocalized with VACHT in striatal cholinergic terminals in the striatum of both M4 receptor WT and KO mice. Quantitation of confocal images (see Materials and Methods for details) revealed ~ 8.7% (± 0.4) and 7.1% (± 0.5) colocalization of M2 receptors with VACHT in cholinergic terminals in M4 receptor WT and M4 receptor KO mice, respectively. These values represent the minimum degree of colocalization as either marker may be present but may not detectable with the method used here. Because the majority of M2 receptors does not localize to VACHT-positive cholinergic terminals (in either M4 receptor WT or KO mice),
most of the M₂ receptor immunoreactivity is likely to represent M₂ receptors present on aspiny dendrites of cholinergic interneurons or M₂ receptors localized to terminals of noncholinergic, excitatory synapses (Hersch et al., 1994). In any case, our data clearly show that the lack of M₄ receptors does not lead to a redistribution of M₂ receptors on cholinergic terminals in the striatum.

Because the neurochemical studies suggested that M₄ receptors function as muscarinic autoreceptors in the striatum, we also determined to which extent M₄ receptors colocalized with VACHT in striatal cholinergic terminals. Quantification of confocal microscopic images indicated that ~6.5% (±0.3) of M₄ receptors colocalized with VACHT in cholinergic terminals (Fig. 6). Similar to M₂ receptors, most of the striatal M₄ receptors do not localize to cholinergic terminals (M₄ receptors are predominately found on dendritic spines of striatal projection neurons and also on terminals of asymmetric, excitatory synapses (Hersch et al., 1994; Bernard et al., 1999). It is possible that the level of M₄ receptors on striatal cholinergic terminals is lower than on dendrites which could limit the sensitivity of detection of these receptors by immunofluorescence. Taken together, the muscarinic receptor–VACHT colocalization data are consistent with our observation that M₄ receptors can act as release-inhibiting autoreceptors in the striatum.

**DISCUSSION**

The present study was designed to identify the mAChR subtypes that mediate autoinhibition of ACh release in various brain regions. Based on the results of functional studies using muscarinic agonists and antagonists of limited mAChR subtype selectivity, considerable controversy exists regarding which specific mAChRs represent the primary inhibitory autoreceptors in different regions of the CNS (see introductory remarks for more details). To avoid the pitfalls associated with the use of classical pharmacological tools (ligands), we decided to study [³H]ACh release using brain slices from different mAChR KO strains. Specifically, we examined the nature of the inhibitory muscarinic autoreceptors in hippocampus, cerebral cortex, and striatum, using tissues prepared from M₂–M₄ receptor double KO and M₂ and M₄ receptor single KO mice (and their corresponding WT controls). The M₂ and M₄ mAChRs (but not the M₁, M₃, and M₅ mAChRs) are efficiently coupled to G-proteins of the Gₛ/G₁ family (Hulme et al., 1990; Caulfield, 1993; Wess, 1996) that are predicted to play a role in the inhibition of neurotransmitter release via inhibition of voltage-sensitive Ca²⁺ channels (Caulfield, 1993; Shapiro et al., 1999). It seemed therefore reasonable to assume that M₂ and/or M₄ receptors are the most likely candidates involved in autoinhibition of ACh release.

By using a strategy similar to that described here (use of mutant mice lacking specific α₂-adrenergic receptor subtypes), recent studies have identified specific adrenergic receptor subtypes mediating autoinhibition of electrically stimulated norepinephrine release in several central and peripheral tissues (Trendelenburg et al., 1999, 2001).

**The M₂ receptor subtype represents the predominant inhibitory muscarinic autoreceptor in hippocampus and cerebral cortex**

We initially performed [³H]ACh release studies using superfused brain slices derived from M₂/M₄ receptor double KO and their WT control mice. These experiments showed that oxotremorine-induced inhibition of potassium-stimulated [³H]ACh release was totally abolished in hippocampal, cortical, and striatal preparations from M₂–M₄ receptor double KO mice (Figs. 2A, 3A, 4A).

To examine whether autoinhibition of ACh release was mediated by M₂ or M₄ receptors (or by a mixture of the two receptors), we next performed analogous studies with tissues from M₂ and M₄ receptor single KO mice. In hippocampal and cortical preparations, the lack of M₄ receptors had no significant effect on agonist-dependent inhibition of stimulated [³H]ACh release (Figs. 2C, 3C). In contrast, in hippocampal and cortical preparations from M₂ receptor KO mice, this activity was completely abolished (Figs. 2B, 3B). These observations demonstrate in a direct and unambiguous manner that the M₂ subtype represents the predominant muscarinic autoreceptor in hippocampus and cerebral cortex. This observation is consistent with high-resolution microscopic studies indicating that the M₂ receptor is abundantly expressed on cholinergic nerve terminals in the hippocampus (Rouse et al., 2000). Moreover, a recent in vivo microdialysis study using antisense oligodeoxynucleotides to reduce the expression levels of M₂ or M₄ receptors also arrived at the conclusion that hippocampal inhibitory muscarinic autoreceptors primarily consist of M₂ receptors (Kitaiuchi et al., 1999).

Reduced levels of ACh are consistently found in cerebral cortex and hippocampus of patients suffering from Alzheimer’s disease, and considerable evidence suggests that this neurochemical deficit makes a major contribution to the cognitive impair-
The M₄ receptor subtype represents the primary inhibitory muscarinic autoreceptor in the striatum

Properly regulated muscarinic neurotransmission in the striatum is known to play an important role in the regulation of extrapyramidal locomotor activity (Di Chiara et al., 1994). Most notably, a misbalance between muscarinic and dopaminergic neurotransmission in the striatum is considered a hallmark of Parkinson’s disease (Hornykiewicz, 1981; Fahn et al., 1990). Identification of the mAChR subtype that mediates autoinhibition of ACh release in the striatum is therefore of considerable therapeutic interest. In contrast to what we observed with hippocampal and cortical tissues, oxotremorine-induced inhibition of stimulated [³H]ACh release remained largely intact in striatal slices from M₄ receptor KO mice (Fig. 4B). On the other hand, oxotremorine virtually lost its ability to mediate inhibition of stimulated [³H]ACh release in striatal slices from M₄ receptor KO mice (Fig. 4C). These data clearly demonstrate that the M₄ subtype represents the predominant inhibitory muscarinic autoreceptor in mouse striatum. Because centrally active muscarinic antagonists are widely used in the treatment of Parkinson’s disease (Fahn et al., 1990; Standaert and Young, 1996), our findings should be of considerable therapeutic relevance.

Previous studies have shown that M₂ muscarinic receptors are abundantly expressed in striatal cholinergic interneurons (Hersch et al., 1994; Bernard et al., 1998). These neurons are known to provide the source of striatal ACh and innervate virtually all striatal projection neurons (Di Chiara et al., 1994). It has therefore been proposed that muscarinic autoinhibition of striatal ACh release may be mediated predominantly by M₂ receptors (Hersch et al., 1994; Bernard et al., 1998). However, as discussed in the previous paragraph, our neurochemical data strongly suggest that the M₄ receptor subtype represents the primary inhibitory muscarinic autoreceptor in mouse striatum. Immunoprecipitation studies showed that the lack of M₄ receptors in M₄ receptor KO mice had no significant effect on M₂ receptor expression levels in the striatum (Gomeza et al., 1999b). Similarly, disruption of the M₂ receptor gene had no noticeable effect on overall striatal M₄ receptor densities (Gomeza et al., 1999a). In addition, immunofluorescence studies demonstrated that both M₂ and M₄ receptors are colocalized with VACHT, a marker of cholinergic terminals (Gilmor et al., 1996), in the striatum (Figs. 5, 6). This observation is consistent with previous findings indicating that both M₂ and M₄ receptors are expressed by most striatal cholinergic interneurons (Bernard et al., 1992, 1998, 1999; Hersch et al., 1994; Yan and Surmeier, 1996). However, to the best of our knowledge, this is the first study demonstrating the presence of M₄ receptors on cholinergic terminals in the striatum. Moreover, M₂ receptor/VACHT double labeling studies showed that the lack of M₄ receptors had not significant effect on the pattern of M₂ receptor expression on cholinergic terminals in the striatum (Fig. 5). These data therefore exclude the possibility that the lack of muscarinic autoinhibition observed with striatal preparations from M₂ receptor KO mice is an artifact caused by altered M₂ receptor expression levels or altered M₂ receptor distribution.

To verify that potassium-stimulated [³H]outflow in mouse striatal slices primarily represented radiolabeled ACh, we used a reverse HPLC method to separate [³H]ACh from [³H]choline (Fig. 1A). This analysis showed that >90% of the potassium-stimulated [³H]overflow in the striatum consisted of authentic ACh (Fig. 1B, Table 1), thus excluding the possibility that the preferential release of radiolabeled compounds other than [³H]ACh may have affected the outcome of the neurochemical studies in the striatum.

Taken together, both the neurochemical and receptor localization data support the concept that M₄ muscarinic receptors act as inhibitory muscarinic autoreceptors in the mouse striatum. Because M₂ receptors are also present on the terminals of cholinergic striatal interneurons (see Discussion), it remains unclear at present why muscarinic autoinhibition of striatal ACh release was not significantly affected by the absence of M₂ receptors. One possibility is that a potential contribution of M₂ receptors to muscarinic autoinhibition of striatal ACh release may have been masked by the presence of the functionally predominant M₄ receptors in the M₂ receptor KO mice. In contrast to M₂ receptors, M₄ receptors are abundantly expressed by a subpopulation of striatal projection neurons (Hersch et al., 1994; Bernard et al., 1999). At present, we cannot completely rule out the possibility that the absence of these receptors may have contributed, through an indirect mechanism, to the loss of muscarinic autoinhibition in striatal slices from M₂ receptor KO mice, e.g., by altering the release of other neurotransmitters such as GABA or glutamate.

In conclusion, our results demonstrate that mAChR KO mice...
represent highly useful tools to assess the molecular identity of the mACHRs that mediate autoinhibition of ACh release in different regions of the brain. We provide direct evidence that the M$_3$ subtype represents the predominant inhibitory muscarinic autoreceptor in hippocampus and cerebral cortex, whereas the M$_4$ receptor subtype functions as the primary inhibitory muscarinic autoreceptor in striatum. These results provide a rational basis for the development of novel muscarinic drugs for a variety of pathophysiological conditions including Alzheimer’s and Parkinson’s disease. Moreover, our findings suggest that it should be possible to design therapeutic strategies aimed at selectively modulating ACh release in distinct regions of the brain.

REFERENCES


Hamilton SE, Loose MD, Qi M, Levey AI, Hille B, McKnight GS, Di Chiara G, Morelli M, Consolo S (1994) Modulatory functions of acetylcholine in rat hippocampus and cerebral cortex, whereas the M$_4$ receptor subtype functions as the primary inhibitory muscarinic autoreceptor in striatum. These results provide a rational basis for the development of novel muscarinic drugs for a variety of pathophysiological conditions including Alzheimer’s and Parkinson’s disease. Moreover, our findings suggest that it should be possible to design therapeutic strategies aimed at selectively modulating ACh release in distinct regions of the brain.


