Research Article

Role of M2 Muscarinic Receptor in the Airway Response to Methacholine of Mice Selected for Minimal or Maximal Acute Inflammatory Response

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Received 3 December 2012; Revised 16 February 2013; Accepted 18 February 2013

Academic Editor: Alexandre de Paula Rogerio

Airway smooth muscle constriction induced by cholinergic agonists such as methacholine (MCh), which is typically increased in asthmatic patients, is regulated mainly by muscle muscarinic M3 receptors and negatively by vagal muscarinic M2 receptors. Here we evaluated basal (intrinsic) and allergen-induced (extrinsic) airway responses to MCh. We used two mouse lines selected to respond maximally (AIRmax) or minimally (AIRmin) to innate inflammatory stimuli. We found that in basal condition AIRmin mice responded more vigorously to MCh than AIRmax. Treatment with a specific M2 antagonist increased airway response of AIRmax but not of AIRmin mice. The expression of M2 receptors in the lung was significantly lower in AIRmin compared to AIRmax animals. AIRmax mice developed a more intense allergic inflammation than AIRmin, and both allergic mouse lines increased airway responses to MCh. However, gallamine treatment of allergic groups did not affect the responses to MCh. Our results confirm that low or dysfunctional M2 receptor activity is associated with increased airway responsiveness to MCh and that this trait was inherited during the selective breeding of AIRmin mice and was acquired by AIRmax mice during allergic lung inflammation.

1. Introduction

The cholinergic system plays a role in the regulation of many pathophysiological mechanisms. In the lung, the vagal parasympathetic nervous system via muscarinic receptors represents the dominant autonomic control of airway smooth muscle tone. Acetylcholine released at neuromuscular junctions binds to M3 muscarinic receptors in the smooth muscle and promotes airway contraction through a number of well-defined intracellular signaling mechanisms [1, 2]. An important negative feedback mechanism is given by the neuronal M2 muscarinic receptor that inhibits acetylcholine release [3, 4]. Airway hyperresponsiveness (AHR) is an exaggerated smooth muscle constriction observed in certain individuals among a population, which can occur in response to a variety of stimuli such as histamine, exercise, cold air, and methacholine (MCh). AHR is a cardinal feature of asthma [5]. Although there are exceptions [6], studies in asthmatic patients indicate a positive correlation between AHR and allergic eosinophilic inflammation [7]. Accordingly, susceptibility to develop asthma is associated with IL-5, a key cytokine for eosinophil differentiation, activation, and survival [8, 9].
One of the proposed mechanisms for allergic AHR to MCh is loss of function of vagal M2 receptors caused by eosinophilic inflammation [10–14]. Indeed, blockage of M2 receptors results in an increased acetylcholine release from vagal nerve endings that ultimately enhances bronchoconstriction [4, 15]. In this sense, AHR can be viewed as an integrated immune-neural circuit. Another well-established immune-mediated neural circuit is the cholinergic anti-inflammatory pathway [16–18]. This efferent pathway is triggered by proinflammatory cytokines via afferent sensory neurons and culminates in the release of acetylcholine by the vagus nerve. In turn, acetylcholine signals thorough nicotine acetylcholine receptor subunit α7 (α7nAChR), expressed on immune cells, inhibiting proinflammatory cytokine production [19]. Because the cholinergic system is involved in bronchoconstriction as well as regulation of inflammation, the present study aimed to evaluate the airway responses to MCh and airway allergic responses in two mouse lines selected to respond maximally (AIRmax) or minimally (AIRmin) to an innate inflammatory stimuli [20], focusing mainly on the role of M2 receptors. These mouse lines were produced by bidirectional selective breeding from a genetically heterogeneous population and behave differently in several experimental models such as arthritis [21], lung cancer [22], and bacterial infection [23]. Here we evaluated whether an innate response that selected the gene combinations determining AIRmax and AIRmin phenotypes would affect basal (intrinsic) or allergic (extrinsic) airway responses to MCh as well as an immune-adaptive T-helper-type-2- (Th2-) mediated stimuli in an experimental model of lung eosinophilic inflammation.

2. Results

2.1. Respiratory Pattern and Expression of M2 and M3 Muscarinic Receptors. It was previously reported that the basal airway responsiveness varies markedly among inbred strains of mice [24, 25]. Indeed, under basal noninflammatory condition multiple quantitative trait loci linked to AHR were identified in inbred mice strains [26, 27]. Therefore, we first determined the basal respiratory pattern of AIRmin and AIRmax in response to increasing doses of MCh in conscious unrestrained mice using a noninvasive whole-body barometric plethysmography. AIRmin mice showed a marked increase in Penh values when compared to AIRmax mice (Figure 1(a)). The dose of MCh required to elicit a 200% increase over baseline values (PC200) was 2.5-fold higher in AIRmax than in AIR min (Figure 1(b)). Our results indicate that the genetic selective process markedly influenced airway response to MCh. In this context, AIRmin animals can be considered as hyperresponsive whereas AIRmax animals are hyporresponsible to MCh.

The contractile response of airway smooth muscles to muscarinic agonists is induced mainly by activation of muscle M3 receptor subtype and inhibited by activation of vagal M2 receptor subtype. Thus, we determined the expression of M3 and M2 muscarinic receptors in lung tissue. We found that M3 receptor expression was lower in AIRmin than in AIRmax animals, as revealed by real-time PCR and Western blot analysis (Figures 1(c)-1(d)), thus, it could not explain the hyperresponsiveness of AIRmin mice to MCh. Therefore, we evaluated M2 muscarinic receptor expression by PCR and Western blot analyses. M2 receptor expression was lower in AIRmin than in AIRmax mice (Figures 1(e)-1(f)). Altogether, these results indicate that low expression of muscarinic M2 receptor in AIRmin mice might be associated with the increased AHR to MCh observed in these animals.

2.2. In Vivo M2 Muscarinic Receptor Activity. To evaluate the role of M2 receptor subtype on the airway responses in vivo, we treated the animals with gallamine, a selective M2 receptor antagonist. Gallamine treatment increased significantly Penh values in AIRmax but not in AIRmin mice when compared with untreated mice (Figures 2(a)-2(b)). However, gallamine treatment increased the responses to MCh in AIRmin animals indicating some activity of M2 receptor, but this activity was not sufficient to generate statistically significant data when compared to untreated group. In our study, we only used one dose of gallamine based on a previous report [28]. Taken together, these results indicate that the negative modulatory effect exerted by vagal M2 receptors was present in AIRmax mice but minimally in AIRmin animals. Therefore, the robust response to MCh observed in AIRmin mice and the lack of significant effect of gallamine indicate dysfunctional vagal M2 activity in this mouse line.

2.3. Ex Vivo M2 Receptor Activity. The results obtained in vivo with the noninvasive method indicated a major role of vagal M2 receptors in inhibiting airway constriction. In order to determine the role of muscle M2 receptors, we evaluated ex vivo the effect of gallamine on smooth muscle constriction to MCh in isolated trachea. In this scenario, vagal receptors are excluded, and only muscarinic receptors of airway smooth muscle responded to MCh. We found that tracheal smooth muscle constriction induced by MCh was more intense in AIRmax when compared with AIRmin animals (Figure 3(a)). Gallamine treatment did not affect tracheal smooth muscle constriction induced by MCh (Figures 3(b)-3(c)), indicating that muscle M2 receptors have no role in smooth muscle constriction induced by MCh.

2.4. Allergic Lung Inflammation and Expression of Muscarinic Receptors. Loss of M2 receptor function due to allergic inflammation has been well documented [29, 30]. It has been shown that eosinophilic major basic protein plays a key role in M2 receptor dysfunction (for review see [10]). We first determined the magnitude of allergic inflammation developed in AIRmin and AIRmax mice referred as OVA group. Figure 4 shows that allergic lung inflammation, as revealed by total number of inflammatory cells (Figure 4(a)) and eosinophils (Figure 4(b)) in BAL, was more intense in AIRmax than in AIRmin animals. Accordingly, the levels of Th2 cytokines (IL-4 and IL-13) and mucus production were higher in AIRmax than in AIRmin mice (Figures 4(c)-4(e)). Histological analysis confirmed the more intense allergic inflammation in AIRmax when compared with AIRmin.
Figure 1: Respiratory pattern and expression of muscarinic receptors in AIRmin and AIRmax mice. Respiratory pattern was determined in awake, unrestrained mice by noninvasive whole-body barometric plethysmography. (a) Penh values were used as an index of bronchoconstriction induced after sequential delivery of increasing concentrations of MCh (3, 6, 12, and 25 mg/mL) and (b) provocative concentration of aerosol MCh at a 200% increase (PC200) over baseline values. Gene (c and e) or protein (d and f) expression of M2 and M3 muscarinic receptors was evaluated by real-time PCR or Western blot analysis in lungs from AIRmin and AIRmax mice. Real-time PCR was carried out using β-actin gene expression as internal control for normalization of M3R (c) and M2R (e) mRNA transcription levels. All PCR reactions were quantitative reactions made by real-time PCR. In Western blot analysis the density of M3R (d) and M2R (f) protein expression was nor aerosol at a malized to actin expression in each sample. Western blots were quantified by densitometry using the ImageJ software (NIH). Real-time PCR and Western-blot analyses were performed using pooled lungs from 5 mice. Data are expressed as mean ± SEM of five mice per group and are representative of three experiments; *P < 0.05.
Figure 2: Respiratory pattern and airway resistance of AIRmin and AIRmax mice in the presence or absence of gallamine, a specific M2 receptor antagonist. Respiratory pattern to inhaled MCh was measured using noninvasive method in AIRmin (a) and AIRmax (b) mice. Airway resistance was measured by invasive method in AIRmin (c) and AIRmax (b) mice. Measurements were performed in the presence (black symbols) or absence (open symbols) of gallamine, in response to increasing concentrations of MCh. Results are reported as Penh values (a-b). Penh data represent the means ± SEM of five mice per group and are representative of three experiments (unpaired t-test, * P < 0.05 as compared to PBS group).

3. Discussion

Our work shows that AIRmin and AIRmax mouse lines represent a model to study lung muscarinic receptor functions and its relation with the development of an allergic lung disease. The main cholinergic pathways are depicted in Figure 6. We found that AIRmin and AIRmax animals develop different patterns of airway reactivity under basal or inflammatory conditions. Notably, respiratory responses to increasing doses of MCh observed in AIRmin mice in basal condition were significantly higher than that obtained in AIRmax animals. Previous studies identified multiple quantitative trait loci (QTL) linked to basal, noninflammatory airway hyperreactivity in specific inbred mouse strains [26, 27, 31]. It was found in A/J mice that the enhanced baseline airway reactivity was independent of lymphocytes or bone marrow-derived cells [32]. Airway responses to MCh are regulated by muscarinic M3 receptor present on muscle cells, which induces muscle constriction, and negatively regulated by the activation of muscarinic M2 receptors on vagal nerve endings, which reduce acetylcholine release at the neuromuscular junction [2, 3, 33, 34]. Here we present evidence that enhanced basal (intrinsic) airway reactivity of AIRmin mice is associated with the low expression and activity of M2 receptors. This assumption is based on the following findings: firstly, mRNA and protein expression of M2 receptors were decreased in AIRmin mice compared to AIRmax animals; secondly, although mRNA and protein expression of M3 receptors, the major muscarinic pathway for airway smooth muscle constriction, were increased in AIRmax mice, these animals did not show enhanced basal airway reactivity; thirdly, the use of gallamine, a selective M2 receptor antagonist significantly increased the airway resistance of AIRmin mice as shown by peribronchovascular cellular infiltrate and increased mucus (Figure 4(f)).

We next determined the respiratory pattern of allergic AIRmin and AIRmax animals in the absence or presence of gallamine. Firstly, we found that allergic AIRmin and AIRmax mice showed higher Penh values when compared with control mice (Figures 5(a)-5(b)). Statistical analysis performed on data, represented by area under the curve, confirmed that the responses to MCh were significantly increased in allergic mice when compared with control non-allergic mice (Figures 5(c) and 5(d)). However, no significant differences were observed in Penh values of allergic mice treated or not with gallamine (Figures 5(a)-5(d)).

Because allergic mice showed an enhanced response to MCh when compared with control mice and gallamine treatment was ineffective in increasing Penh values, we asked whether the expression of muscarinic receptors has changed during allergic inflammation. We found that allergic inflammation did not change the expression of M2 or M3 receptors (Figures 5(e)-5(h)). Altogether, these data suggest that M2 muscarinic receptors dysfunction rather than altered muscarinic expression might be responsible for the increased response to MCh observed in allergic AIRmax or AIRmin mice.
responses of AIRmax mice, but affected minimally the responses of AIRmin mice. These results indicated that gallamine treatment impaired M2 function in AIRmax but not in AIRmin mice. Importantly, in conditions where the participation of vagal M2 receptors were precluded such as in *ex vivo* experiments with isolated tracheal rings, the tracheal responsiveness of AIRmin mice to MCh was not affected by gallamine treatment, reinforcing the predominant
Figure 4: Allergic lung inflammation in AIRmin and AIRmax mice. AIRmin or AIRmax mice were sensitized twice (s.c.) with ovalbumin adsorbed on alum and challenged twice with ovalbumin (i.n.) (OVA group). Control group consisted of nonmanipulated animals. The experiments were performed 24 h after the last OVA challenge. The total inflammatory cells numbers (a), the number of eosinophils (b), and the levels of cytokines IL-5 (c) and IL-13 (d) were determined in bronchoalveolar lavage (BAL) fluid. The mucus index was determined by histocytometry of lung sections stained with periodic acid-Schiff (e) representative lung micrographs stained with HE of control and allergic AIRmin or AIRmax mice (original magnification ×200) (f). Data are expressed as mean ± SEM of four mice per group and are representative of two experiments; Statistical analyses of Student's t-test for (a), (b), (c), and (d). Statistical analyses of Mann Whitney-test for (e). *P < 0.05 relative to control group. #P < 0.05 relative to AIRmin mice.
Figure 5: Continued.
role of M3 receptors in smooth muscle constriction. Our ex vivo experiments with tracheal rings are in line with previous studies in M2 receptor KO mice, which showed that M2 receptors do not exert an inhibitory effect on tracheal constriction induced by cholinergic stimuli [35, 36]. Several studies have shown that the loss of M2 receptor function increases acetylcholine release and potentiates vagally mediated bronchoconstriction (reviewed in [15]). Similar to our findings, experiments performed with estrogen receptor-α KO mice showed enhanced airway responsiveness to inhaled MCh and serotonin under basal conditions and this was associated with reduced M2 receptor expression and function [28, 36]. In addition, our results are in line with the work of Fisher et al. showing that M2-deficient (KO) mice display a significantly enhanced in vivo bronchoconstrictor response to vagal stimulation [37].

In allergic conditions, AIRmin and AIRmax mice showed a significant increase over baseline response (control group) in Penh value. The change in respiratory pattern of AIRmax animals after allergic inflammation could not be attributed to increased expression of M3 receptors. Because gallamine treatment had no effect, we favor the notion that the major mechanism for the increased respiratory responses of allergic AIRmax mice was due to decreased expression and/or dysfunction M2 receptor. This finding is in accordance with previous studies in allergic mice or asthmatic patients indicating that M2 receptor dysfunction rather than M3 receptor overexpression is associated with airway hyperreactivity (reviewed in [3, 4, 33]). It was shown by Ricci et al. (2002, 2008) that patients with asthma or rhinitis present different expression M2 and M3 receptors on lymphocytes, but the role of these receptors in these cells is still elusive [38, 39].

A striking difference between AIRmin and AIRmax animals was related to the intensity of T-helper-type-2-(Th2-) mediated allergic responses such as eosinophilic inflammation, type 2 cytokines production, and mucus production that were all significantly higher in AIRmax when compared with AIRmin animals. It is known that inflammatory mediators play a role in enhancing bronchoconstriction by several pathways. The major basic protein from eosinophils can bind directly to M2 receptors, blocking the interaction with acetylcholine that leads to a higher release of acetylcholine in airway smooth muscles [10]. In addition, type 2 cytokines play a critical role in the development of bronchial responsiveness. IL-13, for instance, may increase bronchoconstriction by directly modulating smooth muscle contractibility and promoting airway hyperreactivity through multiple molecular mechanisms [40, 41]. The participation of these signaling pathways in our asthma murine model in AIRmin and AIRmax animals remains to be determined.

Interestingly, AIRmin and AIRmax mice that were selected by an innate stimuli maintained the polarized inflammatory profile with a stimulus derived from T-cell-dependent adaptive inflammation. We speculate that the enhanced Th2 responses of AIRmax animals was due to an enhanced production of inflammatory cells in bone marrow, as shown previously for neutrophils [42].
FIGURE 6: A model illustrating cholinergic receptors on pulmonary parasympathetic nerves and airway smooth muscle. Acetylcholine (ACh) released at parasympathetic ganglion binds to nicotinic (NI) and muscarinic (M1) receptors and activates postganglionic cholinergic nerve to release ACh at neuromuscular junction. ACh at neuromuscular junctions binds to M3 receptors in the airway smooth muscle, inducing muscle contraction, and to the autoinhibitory M2 receptors located in the vagus nerve, that limits ACh release at the neuromuscular junction and decreasing smooth muscle constriction.

How M2 receptors could be associated with two contrasting situations? On one side, M2 dysfunction was associated with AIRmin animals that have been selected for low inflammatory response and on the other side with AIRmax animals that develop an intense allergic inflammation. These apparently conflicting findings can be reconciled by considering that acetylcholine can also regulate inflammation via the cholinergic anti-inflammatory pathway [16, 17, 43, 44]. It follows that cholinergic anti-inflammatory pathway would favor the selective process of AIRmin animals. The a7nAChR is coded by Chrm7 gene, which is mapped closed to inflammatory response modulator 1 (Irm1) locus on chromosome 7 [45]. The Irm1 locus region contains about 230 known genes that are involved in inflammatory response regulation [45]. Interestingly, AIRmin mice present the same Irm1 locus haplotype as A/J MCh hyperresponsive mice. The genotype of AIRmin at the rs32017050 (SNP at the peak of the Irm1 locus) is “TT” similar to A/J mice, which is one of the 8 parental strains used in F0 population of those selected mice. On the other hand, AIRmax mice are “CC”, showing identical genotypes of C57BL/6J, CBA/J, DBA/2J, P/J, and SWR/J mice (mouse genomic informatics) that are hyporesponsive to MCh. Thus, M2 receptor dysfunction in AIRmin animals can be viewed as an additional mechanism inherited during the selective process that contributed for the low inflammatory phenotype. In the same vein, the loss of M2 receptor function observed in AIRmax animals during allergic inflammation could be interpreted as a neural homeostatic mechanism of the organism to control allergic inflammation [46]. Overall, the neuronal circuits that operate in inflammation or allergic asthma can be viewed as homeostatic mechanisms to counterbalance the inflammatory process.

Since asthma is a complex syndrome, the AIRmin/AIRmax model might mirror clinical cases where airway reactivity is more prominent than airway inflammation as described in fatal asthma and in clinical conditions where airway inflammation is more robust than airway reactivity [47].

4. Material and Methods

4.1. Mice. The AIRmax and AIRmin original mice were obtained by bidirectional selective breeding starting from a high polymorphic population produced by intercross between eight inbred mouse strains (A, DBA-2, P, SWR, CBA, SJL, BALB/c, and C57Bl/6) [20]. The selective breeding of these animals for maximal or minimal acute inflammatory response was based on both cellular infiltrate and protein contents in the inflammatory exudate 24 h after subcutaneous implant of polyacrylamide beads (Biogel). The main interline difference was characterized by the strong differential neutrophil PMN count in inflammatory exudate. Several quantitative trait loci (QTL) were recently detected in these mouse lines [46, 48]. Animals were obtained and maintained at the animal facilities of the Immunogenetics Laboratory at Butantan Institute on standard pellet food and water ad libitum. All experiments were performed following the guidelines for animal use approved by the Ethics Committee in Animal Experimentation of the University of São Paulo, Brazil, which is in accordance with the Ethical Principals of the Brazilian College of Animal Experimentation (COBEA), ethics protocol number 26, page 30, Book 2, ICB, USP.

4.2. Ovalbumin (OVA) Sensitization and Challenge. Mice were sensitized and boosted subcutaneously (s.c.) with 4 μg of chicken OVA (Sigma-Aldrich, St. Louis, MO, USA) and 1.6 mg of aluminum hydroxide gel in 0.2 mL of sterile PBS on days 0 and 7. Removal of lipopolysaccharides from OVA was performed as previously described [49]. After LPS removal, the endotoxin level of OVA was below the limit of detection in the LAL (Limulus amoebocyte lysate) assay (QCL-1000 kit, BioWhittaker, Walkersville, MD, USA). Airway inflammation was induced by two intranasal (i.n.) challenges with 10 μg OVA on days 14 and 21, animals were anaesthetized with ketamine (50 mg/kg body weight) and xilazine (20 mg/kg body weight) intraperitoneally in 200 μL of PBS for intranasal instilations. Experiments were performed 24 h after the last i.n. OVA challenge, on day 22.

4.3. Airway Responsiveness to Methacholine by Noninvasive Barometric Plethysmography. Respiratory parameters were determined before and after administration of increasing doses of inhaled MCh (3, 6, 12, and 25 mg/mL) by noninvasive method in conscious unrestrained mice using whole-body plethysmograph (Buxco Electronics Inc. Wilmington, North
4.4. Ex Vivo Trachea Responsiveness to Methacholine. To test M2 muscarinic receptor function, mice were exposed or not to M2 muscarinic receptor antagonist gallamine (30 μM/animal) [28], nebulized 30 minutes before recording of respiratory parameters.

4.5. Bronchoalveolar Lavage Fluid. Mice were deeply anaesthetized by intraperitoneal injection of xilazine (50 mg/kg body weight) and ketamine (100 mg/kg body weight), followed by cervical dislocation. Blood samples from the retro-orbital plexus were collected for serum antibody determinations. The trachea was cannulated, and lungs were washed twice with 0.5 and 1.0 mL PBS, respectively. Total and differential cell counts of bronchoalveolar lavage (BAL) fluid were determined by haemocytometer and cytospin preparation stained with Instant-Prov, hematoxylin-eosin (Newprov, Pinhais, PR, Brazil).

4.6. Cytokines Measurements. The levels of cytokines (IL-5 and IL-13) in the BAL fluid were assayed by sandwich kit ELISA according to the manufacturer’s instruction (PharMingen, San Diego, CA, USA), as previously described [53]. For IL-13 determinations, the pair used was 38 213.11 and biotinylated goat polyclonal anti-IL-13 from R&D Systems (Minneapolis, MN, USA). Values were expressed as pg/mL deducted from standards that run in parallel with recombinant cytokines. The limit of detection was 10 pg/mL for IL-5 and 31 pg/mL for IL-13.

4.7. Histological Analyses. After BAL collection, lungs were perfused via the right ventricle with 10 mL of PBS to remove residual blood, then immersed in 10% phosphate-buffered formalin for 24 h, and in 70% ethanol until embedded in paraffin. Lung sections with 5 μm were stained with hematoxylin/eosin for evaluation of peribronchial and perivascular lung inflammation or with periodic acid-Schiff (PAS)/haematoxylin for the evaluation of mucus production, as previously described [51]. Briefly, a quantitative digital morphometric analysis was performed using the application program Metamorph 6.0 (Universal Images Corporation, Downingtown, PA, USA). The circumference area of bronchi and the PAS-stained area were electronically measured, and the mucus index was determined by the following formula: (PAS-stained area/bronchial circumference area) × 100.

4.8. Western-Blot Assays. Lungs from mice were homogenized in ice-cold lysis buffer containing protease inhibitors, 10 mM TrisHCl (pH 7.5), 8 M urea, 20 mM EDTA, 150 mM NaCl, and 1% Triton X-100. Membrane proteins (40 μg) were fractionated by differential centrifugation, separated by SDS-PAGE and transferred to nitrocellulose membranes. The incubation procedure was performed using primary and secondary antibodies, as previously described [54–57]. The following polyclonal rabbit antisera were used: anti-M1 (1:200), anti-M2 (1:100), anti-M3 (1:100), and anti-β-actin (1:400) (Santa Cruz Biotechnology, Santa Cruz, CA, USA). β-actin protein expression was used as an internal standard for relative quantification of muscarinic receptors expression levels. Western blots were quantified by densitometry using the ImageJ software (NIH, Washington DC, USA).

4.9. RNA Isolation, Reverse Transcription, Real-Time PCR, and Conventional-PCR. Total RNA was isolated using Trizol (Invitrogen, Carlsbad, CA, USA) from lungs of nonmanipulated Airmax and Airmin mice. Integrity of the isolated RNA was verified by separation of an aliquot of the extracted RNA on a 2% ethidium bromide-stained agarose gel. DNA was removed from RNA samples by incubation with DNase I (Ambion Inc., Austin, TX, USA).

Total RNA (3 μg) was reversely transcribed to cDNA with 200U of RevertAid H MInus M-MuLV-reverse transcriptase (Fermentas Inc., Hanover, MD, USA). DNA templates were amplified by real-time PCR on the 7000 Sequence Detection System (ABI Prism, Applied Biosystems, Foster City, CA, USA) using the Sybr green method [54, 56] or were amplified by PCR and analyzed as described previously [54, 56]. Variations in cDNA concentrations were normalized with β-actin as an internal control, which is a constitutively expressed gene. Experiments were performed in triplicate for each data point. Primer sequences for reverse transcription and quantitative PCR amplification (qRT-PCR) of mRNA used in this study were β-actin forward CTT CTC CCC TGC TTA TGC GTC TT, reverse CCA GCA GTC GCT GGC GTC TT, M2AChR forward GCT GGT GGT GCC TCT CCA TCC, reverse CCA GCC AGA TGG TCT TCC, and M3AChR forward CCA GCC AGA TGG TCT TCC, reverse CCA GCC AGA TGG TCT TCC. Negative controls were conducted on water and on total
RNA. All PCR reactions were quantitative reactions, made by real-time PCR.

4.10. Statistical Analysis. Statistical analyses were performed employing JMP 9.0 (JMP SAS Institute Inc). Results are expressed as mean ± SEM. Continuous variables were log-transformed for the analyses when the normality of the distribution was rejected by the Shapiro-Wilk test. For comparisons between two groups, normality was tested and then data were submitted to Student’s t-test. For comparisons amongst 3 or more groups it was used ANOVA following In the case of absence of normality, even in log-transformed variables, Mann-Whitney was used for comparisons between 2 groups and or Kruskal-Wallis tests for three or more groups.

Acknowledgments

The authors thank Thiago Andrade Patente for the help in statistical analyses and Katia Haipek for the illustration of Figure 6.

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