**M₁ and M₃ muscarinic receptors may play a role in the neurotoxicity of anhydroecgonine methyl ester, a cocaine pyrolysis product**

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The smoke of crack cocaine contains cocaine and its pyrolysis product, anhydroecgonine methyl ester (AEME). AEME possesses greater neurotoxic potential than cocaine and an additive effect when they are combined. Since atropine prevented AEME-induced neurotoxicity, it has been suggested that its toxic effects may involve the muscarinic cholinergic receptors (mAChRs). Our aim is to understand the interaction between AEME and mAChRs and how it can lead to neuronal death. Using a rat primary hippocampal cell culture, AEME was shown to cause a concentration-dependent increase on both total [³H]inositol phosphate and intracellular calcium, and to induce DNA fragmentation after 24 hours of exposure, in line with the activation of caspase-3 previously shown. Additionally, we assessed AEME activity at rat mAChR subtypes 1–5 heterologously expressed in Chinese Hamster Ovary cells. I-[N-methyl-³H]scopolamine competition binding showed a preference of AEME for the M₂ subtype; calcium mobilization tests revealed partial agonist effects at M₁ and M₃ and antagonist activity at the remaining subtypes. The selective M₁ and M₃ antagonists and the phospholipase C inhibitor, were able to prevent AEME-induced neurotoxicity, suggesting that the toxicity is due to the partial agonist effect at M₁ and M₃ mAChRs, leading to DNA fragmentation and neuronal death by apoptosis.

Cocaine, a tropane alkaloid found in *Erythroxylum coca* leaves, is a recreational drug used worldwide. Its abuse is a public health problem, affecting 17 million people (range 14–21 million) during 2012, which corresponds to a 0.4% rate of annual prevalence. Cocaine usage occurs with high incidence in North and...
South America (1.8 and 1.2% annual prevalence rates, respectively), Oceania (1.5%), and Western and Central Europe (1%)\(^1\). It blocks the uptake of serotonin, norepinephrine and dopamine in presynaptic nerve terminals, as well as voltage-specific sodium channels, responsible for the local anesthetic effect\(^7\). Cocaine may cause intense vasoconstriction, endothelial cell dysfunction, oxidative stress and platelet aggregation\(^2\)–\(^5\), which are responsible for the main systemic adverse effects of its abuse, including stroke, myocardial infarction, arterial dissection, vascular thrombosis, rhabdomyolysis and renal complications\(^6\)–\(^7\).

There are two distinct chemical forms of cocaine: hydrochloride (‘street’ cocaine, ‘coke’), a water-soluble powder which can be taken orally, intranasally or intravenously; and ‘freebase’ or ‘crack’ cocaine, which is cocaine without the hydrochloride moiety\(^8\)–\(^9\). Crack cocaine is the smoked form of cocaine and has greater addictive potential than other routes of cocaine administration\(^10\). As crack cocaine has a low melting point (96–98°C), the heating process quickly volatilizes the cocaine, which is rapidly absorbed by the lungs and reaches the brain faster than any other route. Along with cocaine, anhydroecgonine methyl ester (AEME), a cocaine pyrolysis product, is also absorbed by the lungs\(^11\). Up to 80% of cocaine can be converted to AEME, depending on the temperature, the purity of the crack cocaine and the smoking devices\(^12\).

Little is known about AEME effects. To the best of our knowledge, the data available about this substance thus far includes only studies in the peripheral system, i.e., reductions in blood pressure and heart rate in rabbits with an increase in the respiratory rate\(^13\); a negative inotropic effect in vitro possibly mediated by muscarinic cholinergic receptors (mACHRs), since it was reversed by atropine, a nonspecific muscarinic receptor antagonist\(^14\); cardiovascular effects, e.g., hypotension and tachycardia in sheep, which are also antagonized by intravenous administration of atropine, again consistent with a muscarinic cholinergic effect\(^15\). However, the mechanisms of AEME in the central nervous system are poorly investigated and not well understood.

Our group first described the neurotoxicity of AEME and also the involvement of mACHRs in AEME-induced neuronal death in rat primary hippocampal cell cultures\(^16\). AEME seems to be a neurotoxic agent with greater neurotoxic potential than cocaine, showing an additive effect when combined. Caspase-3 activity in the hippocampal neurons was increased after 6 hours of exposure and seems to be one of the main mechanisms of AEME-induced neurotoxicity. Also, atropine prevented AEME-induced neurotoxicity, reinforcing that mACHRs are involved in AEME’s effects. In addition, binding experiments with hippocampi membrane preparations have confirmed the affinity of AEME for muscarinic cholinergic receptors\(^16\).

The mACHRs belong to the G-protein coupled receptors and include five distinct subtypes, denoted as M\(_1\)–M\(_5\) mACHRs, which are widely distributed throughout the body. The odd-numbered mACHRs subtypes are coupled to G\(_i\)/G\(_\alpha11\) proteins and induce the hydrolysis of phosphoinositide lipids by phospholipases, while M\(_2\) and M\(_3\) mACHRs subtypes, coupled to G\(_i\)/G\(_\alpha\) proteins, inhibit adenylyl cyclase activity. Thus, the activation of M\(_1\), M\(_2\) and M\(_3\) mACHRs subtypes produces inositol trisphosphate (IP\(_3\)) and intracellular calcium, while M\(_2\) and M\(_3\) mACHRs subtypes cause a downstream decrease in cAMP levels\(^17\)–\(^18\). It is important to emphasize that all five mACHRs are expressed in the hippocampus\(^19\)–\(^20\) and they modulate hippocampal function through the inhibition of synaptic activity and/or the increase of neuronal excitability\(^21\).

To better understand the interaction between AEME and mACHRs, the present study investigated: 1) the effects of AEME on the production of total inositol phosphate and intracellular calcium release in rat primary hippocampal cell cultures, as well as DNA fragmentation as a result of caspase-3 activation, which was previously observed by our group\(^16\); 2) the affinity and the action of AEME at mACHRs using Chinese hamster ovary (CHO) cells expressing all five individual rat mAChR subtypes; 3) the effects of M\(_1\) and M\(_3\) selective antagonists on AEME-induced neurotoxicity, as well as the inhibition of phospholipase C (PLC). The comprehension of the mechanisms underlying AEME-induced neurotoxicity could contribute to explain why crack cocaine smoke is more devastating than other routes of cocaine administration.

**Results**

**Experiments with rat primary hippocampal cell culture.** Effect of AEME on total \[^3H\]inositol phosphates accumulation and on intracellular calcium release. AEME and the cholinergic agonist carbachol (control) (10\(^{-8}\) to 10\(^{-3}\) M) caused a concentration-dependent increase of total \[^3H\]inositol phosphate in the primary hippocampal cell culture. Maximum inositol phosphate accumulation was obtained with 10\(^{-3}\) M (10\(^{-3}\) M) AEME and carbachol (Fig. 1A). The basal level of the total \[^3H\]inositol phosphate was 30626 ± 4250 dpm/10\(^6\) cells.

There was a concentration-dependent increase in intracellular calcium release after AEME exposure (Fig. 1B), with the lowest effect observed with 10\(^{-4}\) M AEME and highest with 10\(^{-3}\) M AEME (F\(_{3,36}\) = 13.76; \(p < 0.001\)).

Effect of AEME on DNA fragmentation. Hippocampal cells exposed either to 10\(^{-4}\) or 10\(^{-3}\) M AEME for 12 hours (Fig. 2A) showed the same pattern of DNA fragmentation as the control group (F\(_{2,4}\) = 4.73; \(p = 0.059\)). However, after 24 hours of exposure, 10\(^{-3}\) M AEME increased the number of cells with
fragmented DNA (44.1%; $F_{2.8} = 9.77; p = 0.007$) compared to control (28.6%) and the lower AEME concentration (32.1%; Fig. 2B).

**Experiments with CHO cells.** $B_{\text{max}}$ and $K_d$ determination for each mACHR subtype expressed in CHO cells. The saturation binding of $[^3H]$NMS for each mACHR subtype expressed in CHO cells was specific and saturable and the specific binding fitted best a one-site model (see Supplementary Figure S1). An analysis of three experiments, each one performed in triplicate yielded dissociation constant ($K_d$) and binding capacity ($B_{\text{max}}$), summarized in Table 1.

**Radioligand competition binding.** $[^3H]$NMS displacement curves for AEME are shown in Fig. 3. The AEME $K_d$ values (μM) were: 25.7 for rat M1, 19.5 for rat M2, 33.9 for rat M3, 24.5 for rat M4, and 29.5 for rat M5. The one-way ANOVA showed that the affinity for rat M2 was greater than rat M4; rat M4 greater than rat M3; and rat M1 greater than M3 ($p < 0.05$). Other than this observation, there was no difference among rat M1, rat M2, and rat M3, and no difference between rat M4 and rat M5. The averages of absolute values of controls are: $2274 \pm 187$, $888 \pm 25$, $3698 \pm 303$, $2831 \pm 231$, and $4059 \pm 415$ dpm/25 μg of protein for rat M1, rat M2, rat M3, rat M4, and rat M5, respectively.
Calcium mobilization assays (Figs 4 and 5) were performed to determine AEME’s mechanism of action. AEME exhibited partial agonist-activity at the M₁ and M₃ mAChRs subtypes as observed by an increase in intracellular calcium fluorescence. The effective concentration that increased the response to 50% of maximum (EC₅₀) was greater than 100μM for both rat M₁ and rat M₃, reaching 38.3% (rat M₁) and 27.2% (rat M₃) of the maximum acetylcholine response (Fig. 4). There was

| mAChR subtype | Bₘₐₓ (fmol/mg of protein) | Kᵦ (×10⁻³ μM) |
|----------------|---------------------------|---------------|
| Rat M₁         | 1087 ± 165                | 0.085 ± 0.004 |
| Rat M₂         | 1470 ± 053                | 0.159 ± 0.022 |
| Rat M₃         | 1132 ± 033                | 0.063 ± 0.008 |
| Rat M₄         | 1835 ± 127                | 0.038 ± 0.004 |
| Rat M₅         | 1564 ± 027                | 0.224 ± 0.016 |

Table 1. Saturation binding experiment of [³H]NMS in separate CHO cells expressing one of the five subtypes of mAChRs. Bₘₐₓ and Kᵦ values were determined for radioligand competition binding using GraphPad Prism 5 (data represent the mean of three independent experiments performed in triplicate). Data presented as mean ± SEM.

Figure 3. AEME competition binding curves obtained in CHO cells expressing individual subtypes of muscarinic receptors (data represent the mean of three independent experiments performed in triplicate). The Kᵦ values for AEME (μM) were plotted alongside with the competition binding curves. The affinity for rat M₂ was greater than rat M₄ (p < 0.05), rat M₄ was greater than rat M₁ (p < 0.05), and rat M₁ greater than rat M₃ (p < 0.05). There was no difference among rat M₄, rat M₁, and rat M₃. The averages of absolute values of controls are: 2274 ± 187, 888 ± 25, 3698 ± 303, 2831 ± 231, and 4059 ± 415 dpm/25 μg of protein for rat M₁, rat M₂, rat M₃, rat M₄, and rat M₅, respectively. Data presented as mean ± SEM.

Figure 4. AEME concentration-response curve normalized to the maximum acetylcholine response. CHO cells expressing rat M₁, rat M₃, and rat M₅ were studied (data represent the mean of three independent experiments, each one performed in triplicate). The AEME concentration effect was greater than 100μM for both rat M₁ and rat M₃. The averages of absolute values of control are: 21950 ± 133, 23107 ± 77, and 20117 ± 1323 arbitrary units/4 × 10⁴ cells for rat M₁, rat M₃, and rat M₅, respectively. Data presented as mean ± SEM.

**AEME functional assays.** Calcium mobilization assays (Figs 4 and 5) were performed to determine AEME’s mechanism of action. AEME exhibited partial agonist-activity at the M₁ and M₃ mAChRs subtypes as observed by an increase in intracellular calcium fluorescence. The effective concentration that increased the response to 50% of maximum (EC₅₀) was greater than 100μM for both rat M₁ and rat M₃, reaching 38.3% (rat M₁) and 27.2% (rat M₃) of the maximum acetylcholine response (Fig. 4). There was...
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The effects of 100 μM AEME on acetylcholine concentration-response curves are shown in Fig. 5.

Weak partial agonist activity at rat M₁ and rat M₃ was confirmed by the presence of an increase in calcium fluorescence at low concentrations of acetylcholine (Fig. 5A,C). AEME antagonized the effects of ACh at rat M₂, rat M₄, and rat M₅ (Fig. 5B,D,E; EC₅₀s are summarized in Table 2). In light of the ability of AEME to displace NMS, this suggested potential antagonist activity of AEME at rat M₅. The averages of absolute values of control are: 20307 ± 149, 18233 ± 86, and 23438 ± 205 arbitrary units/4 × 10⁴ cells for rat M₂ and rat M₄, respectively.

As the AEME antagonist effect appeared most robust at the rat M₅ mAChR subtype, Schild analyses were performed to determine antagonist affinity (pA₂). Using this technique, we estimated the pA₂ and Schild slopes, which were 4.87 ± 0.10 and 0.95 ± 0.07, respectively (Fig. 6). These results suggest that AEME is a competitive orthosteric antagonist at rat M₅. The average of absolute values of control is 21255 ± 124 arbitrary units/4 × 10⁴ cells.

Table 2. The EC₅₀ for acetylcholine concentration-response curve in the absence and in the presence of 100 μM AEME (data represent the mean of three independent experiments, each one performed in duplicate). *p < 0.05, **p < 0.01 and ***p < 0.001, compared with ACh concentration-response curve alone for each mAChR subtype. mAChR: muscarinic acetylcholine receptor, DMSO: dimethyl sulfoxide, ACh: acetylcholine. Results are presented as mean ± SEM in nM.
Effect of M1 and M3 selective mAChR antagonists and the PLC inhibitor on AEME-induced neurotoxicity.

MTT viability assay (Fig. 7) was performed to determine the involvement of either M1 or M3 mAChRs, as well as the PLC inhibitor (U73122) on AEME-induced neurotoxicity. After 24 hours of exposure, $10^{-3}$ M AEME presented a significant decrease in neuronal viability, presenting 67.2% of viable cells ($F_{9,220} = 20.87; p < 0.001$). The incubation of AEME in the presence of the M1 selective antagonist, pirenzepine, prevented its neurotoxicity (98.4% of viable cells; $p < 0.001$ compared with AEME), as well as with the M3 selective antagonist, p-fluorohexahydro-sila-difenidol hydrochloride (p-F-HHSiD) (85.4% of viable cells; $p < 0.05$ compared with AEME) and iPLC (U73122) (102.6% of viable cells; $p < 0.01$ compared with AEME).

Discussion

This manuscript comprises the first description of the mechanisms of AEME at mAChRs. In saturation binding experiments with $[^3H]NMS$, we observed monophasic curves, allowing us to determine $B_{max}$ and $K_d$ values for individual mAChR subtypes. To characterize the effect of AEME at these receptors, different concentrations of AEME were examined for their ability to compete with $[^3H]NMS$ for its binding site. The binding curves in competition experiments with AEME were indicative of a single binding site for $[^3H]NMS$. The range of $pK_i$ values obtained for AEME in CHO cells from rat M1 to rat M5 was comparable with $pK_i$ values for AEME from our previous studies with $[^3H]quinuclidinyl benzilate ($[^3H]$QNB; 4.15 ± 0.15) in rat hippocampus, indicating that mAChRs may be a target of AEME. Although the $pK_i$ values were very similar, we observed a slight preference for the M2 mAChR.

In the hippocampus of male rats, immunoprecipitation studies indicated a predominance of the M1 subtype (55%) and low expression of M3 (10%) and M4 (15%)[23]. Their order of abundance has been ranked as $M_1 \gg M_2 > M_3 = M_4 > M_5$[19,20]. In hippocampus of female rats in proestrus, immunoprecipitation studies also confirmed that, although hippocampus expresses all mAChR subtypes, the population of M1 receptors is predominant. In addition, the amount of M2 in the hippocampus is higher than the

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**Figure 6.** Acetylcholine concentration-response curves in the absence or presence of different concentrations of AEME ($10^{-6}$ to $10^{-3.5}$ M) in a calcium mobilization assay using CHO-K1 cells stably expressing rat M5. Schild regression of the dose ratios (DR) derived from the AEME antagonism of acetylcholine was presented and the slope was 0.95 ± 0.07. The average of absolute values of control is 21255 ± 124 arbitrary units/4 × 10^4 cells. Data presented as mean ± SEM. ACh, acetylcholine.

**Figure 7.** Effect of M1 selective antagonist pirenzepine (10 nM), M3 selective antagonist p-fluorohexahydro-sila-difenidol (p-F-HHSiD, 10 nM) and PLC inhibitor (U73122, 10 nM) after 24 hours of exposure to $10^{-3}$ M AEME (n = 4). Potassium chloride (250 mM) was used as a positive control of neuronal death. ***$p < 0.001$, compared with neurobasal medium (control group), **$p < 0.05$, **$p < 0.01$, ###$p < 0.001$compared with $10^{-3}$ M AEME (ANOVA and Newman-Keuls multiple comparison).
amount of M1, M4 and M5 mACHR subtypes in rat hippocampus\textsuperscript{19,20,22}. In the present study we showed that this substance is a partial agonist at M1 and M4 mACHR subtypes through the increase in the intracellular calcium, even in the presence of low concentrations of acetylcholine. Also, 100 μM AEME slightly shifted the acetylcholine concentration-response curves to the right for the M4 mACHR subtypes, characterizing a weak antagonist effect. The same antagonist effect was also observed for M3 mACHR; however, the effect of AEME seems to be relatively higher for mACHRs with lower EC\textsubscript{50} for acetylcholine. It is important to note that AEME increased, in a concentration-dependent manner, total ['H]inositol phosphate, indicating that AEME may act as an agonist at the M1 and/or M4 mACHR subtypes.

The intracellular cascade involved in the activation of the M1 mACHRs increases the intracellular calcium\textsuperscript{23}, which in turn might be involved in the activation of the caspase signaling, leading to neuronal death. In fact, Shih et al.\textsuperscript{24} showed that arecoline, a mACHR agonist, induced neuronal death by apoptosis at concentrations from 50–200 μM. The generation of reactive oxygen species, the decrease in antioxidant defenses and the activation of caspase-3 were some of the mechanisms studied. Besides the chemical similarity among AEME and arecoline, with similar pK\textsubscript{a} values in binding studies on cloned M1-M5 mACHRs\textsuperscript{25}, it seems that these substances share the same neuronal death pathways, culminating in the activation of caspase-3\textsuperscript{16,24}. This executioner enzyme can mediate the catabolic process, characterizing the end-stage of apoptosis\textsuperscript{26}. The current manuscript corroborates with this finding, suggesting that one of the mechanisms involved in DNA fragmentation observed 24 hours of exposure to 10^{-3} M AEME could be explained by the previous activation of caspase-3. Whether, the DNA fragmentation might be a consequence of this intracellular cascade activation remains to be explored.

AEME-induced neurotoxicity could be triggered by an intracellular calcium increase, in a concentration-dependent manner, observed at concentrations starting at 10^{-3} M (100 μM). The endoplasmic reticulum contains calcium release channels which can be activated by IP\textsubscript{3} through IP\textsubscript{3} receptors\textsuperscript{27}. Thus, the increase in free intracellular calcium observed at AEME concentrations starting at 10^{-3} M (0.1 μM) and reaching a maximum at 10^{-3} M (10 μM) may be triggered by the increase in inositol phosphate accumulation, in particular IP\textsubscript{3}. This outcome may be explained by the partial agonist effect at M1 and M3 mACHRs as observed in the in vitro CHO cells data for AEME. It is important to emphasize that the AEME-induced neurotoxicity investigated by Garcia et al.\textsuperscript{16} occurred after a long-term exposure, i.e., different concentrations of AEME for 12 and 24 hours.

Few studies have evaluated the effects of AEME at mACHRs\textsuperscript{13–15}. Our group was the first one to correlate the neurotoxic effects of AEME with the activation of mACHRs, as the nonspecific mACHR antagonist atropine was able to prevent AEME-induced neurotoxicity\textsuperscript{16}. Although M1 and M2 mACHRs subtypes are less expressed in the hippocampus\textsuperscript{19,20} and AEME showed an antagonist effect when interacting with these subtypes, we believe that these effects are not likely to explain AEME-mediated neurotoxicity because of the preventive atropine effect\textsuperscript{16}. However, we cannot discard their importance. M2 mACHRs subtypes are found in this brain region in cholinergic synaptic terminals controlling acetylcholine and other neurotransmitter release, e.g., glutamate, which can promote excitotoxicity at high concentrations\textsuperscript{28}. Also, some substances with an antagonist effect at the M1 and M3 mACHR subtype, when injected in the dorsal hippocampus of rats, induce retrograde amnesia, disrupting memory consolidation\textsuperscript{29}.

Several substances, e.g., the muscarinic toxins (MTs), a group of small proteins isolated from the venom of some snakes, have a high selectivity and affinity for individual mACHR subtypes with competitive antagonist, allosteric modulator, and potential agonists effects\textsuperscript{30}. MT2 toxin, for example, activates M1, M3 and M5 mACHRs, leading to a significant increase in intracellular calcium\textsuperscript{23}. According to Bashkatova et al.\textsuperscript{31}, the stimulation of M1 mACHRs by the injection of M1 agonist McN-A-343 increased nitric oxide and lipid peroxidation in the striatal tissues, the same effect observed for the psychostimulant amphetamine. Several cells, including neurons, present the constitutive form of nitric oxide synthase, which is a calcium/calmodulin-dependent enzyme rapidly activated in response to intracellular calcium increase, leading to nitric oxide production\textsuperscript{32,33}. Depending on the pathophysiological conditions, it could be overproduced, resulting in cellular toxicity and death by oxidative stress and lipid peroxidation\textsuperscript{34}. Interestingly, MT7, a potent non-competitive antagonist toxin at the M1 mACHR subtype, with no antagonist activity at the M1 or M3 mACHRs subtypes\textsuperscript{35}, was able to prevent amphetamine-induced nitric oxide generation and the lipid peroxidation process. The authors attributed that the activation of M3 mACHRs might play a critical role in the neurotoxic process induced by amphetamine\textsuperscript{31}. Our previous study showing the preventive effect of mACHRs antagonist atropine in the AEME-induced neurotoxicity\textsuperscript{16} corroborate with these findings. Moreover, we demonstrated that M1 selective antagonist (pirenzepine) and the M3 selective antagonist (p-F-HHSiD), were able to prevent or reduce AEME-induced neurotoxicity. In addition, the PLC inhibitor U73122 was also able to prevent AEME toxicity indicating that these effects are mediated by PLC activation. Taking together, these results indicate that the neurotoxic effects of AEME may involve the activation of both M1 and M3 mACHRs.

To the best of our knowledge, this is the first study demonstrating AEME mechanism of action at the mACHRs. Its partial agonist effect at M1 and M3 mACHRs may be the cause of AEME neurotoxicity, once the selective antagonists were able to prevent it. The IP\textsubscript{3} accumulation and the increase in free cytosolic calcium, as well as oxidative stress, could result in mitochondrial dysfunction, affecting the electron transfer chain, leading to ATP depletion and neuronal death by caspase activation, which causes DNA fragmentation\textsuperscript{35}. 

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This study corroborates with our previous study, reinforcing the idea that AEME is more than a crack cocaine biomarker; it may play a crucial role in several CNS disorders, including cognitive deficits of crack cocaine users\(^5\), as they are exposed to a mixture of cocaine, AEME and others (e.g. solvents) that could enhance the risk of a neurotoxic effect.

Material and Methods

**Anhydroeucgonine methyl ester (AEME).** Cocaine was gently donated by the Criminal Institute of São Paulo to the Laboratory of Toxicological Analyses (School of Pharmaceutical Sciences, University of São Paulo) for research purposes. Briefly, cocaine was purified (95%) and converted into its salt form, cocaine hydrochloride, by bubbling hydrochloric acid into mixture of purified cocaine dissolved in diethyl ether. Then, AEME was synthesized, using cocaine hydrochloride as start material and purified as previously described by our research group\(^6\). The AEME product (purity $>98\%$) was confirmed by proton nuclear magnetic resonance ($^1\text{H}-\text{NMR}$) and electrospray ionization-mass spectrometry (ESI-MS). For the muscarinic CHO cells studies, AEME was purchased from Lipomed\(^8\) (purity $>98\%$).

**Animals.** Pregnant Wistar rats, weighing 230–250 g, were obtained from Butantan Institute, São Paulo, Brazil. They were housed in plastic cages and maintained in a room with constant temperature ($22 \pm ^\circ\text{C}$) on a 12:12 hours light/dark cycle (lights on at 7:00 a.m.). Food and water were provided ad libitum. This study was performed according to NIH guidelines and approved by Animal Use Ethic Committee of Butantan Institute (protocol number 633/09) and Ethic Committee for Research Project Analysis of School of Medicine at University of São Paulo (protocol number 0841/09).

**Experiments with rat primary hippocampal cell culture.** Hippocampal neurons were dissociated from hippocampi of E18-E19 Wistar rat embryos, as described previously\(^7-10,42\). Pregnant rats were anesthetized with sodium pentobarbital 55 mg/kg and the fetuses were rapidly decapitated to remove their hippocampi. The tissue was placed into a Petri dish containing 100 U/mL penicillin and 100 $\mu\text{g/mL}$ streptomycin (Gibco) in a cooled Neurobasal medium (Gibco). Hippocampi were washed with Hank's Balanced Salt Solution (HBSS) and submitted to a mechanical fragmentation using appropriate scissors. Hippocampi fragments were then transferred to a 0.25% trypsin in Earl's Balanced Salt Solution (EBSS) solution pH 7.2–7.4 and were incubated for 10 minutes at 37°C. After the incubation period, cells were washed with an EBSS solution containing 277.5 U/mL DNase (Sigma) and 10% fetal bovine serum (FBS) (Gibco) and centrifuged at 300 $\times$ g (Eppendorf 5804R) for 2 minutes at 20°C. Neurons were isolated by mechanical dissociation in an EBSS solution (with DNase and fetal bovine serum) using Pasteur pipettes with different diameters sizes and centrifuged for 5 minutes (300 $\times$ g). The tissue was then resuspended in Neurobasal medium (Gibco) supplemented with 0.5 mM L-glutamine, 25 mM L-glutamic acid, 100 U/mL penicillin, 100 $\mu\text{g/mL}$ streptomycin and 2% B27 supplement (Gibco) to reduce glial cell proliferation\(^10,41\). The cells were seeded onto 0.01% poli-L-lysine-coated multiwell culture plate and maintained at 37°C in a humidified atmosphere of 5% CO₂, for 7–8 days, the time required for maturation of hippocampal neurons, forming a network of functional synaptic contacts\(^2\). On the second day, half of the old medium was replaced by the same volume of a fresh medium with the same composition. On the seventh day, cells were incubated with AEME in several concentrations for different time periods, depending on the experiment. Hippocampal neurons were plated on poli-L-lysine-coated 24-well culture plate at a density of $2 \times 10^5$ cells/cm². The culture cells were immunohistochemically characterized with MAP2 (neuronal marker) and GFAP (astrocytic marker) showing a predominance of 92% of neurons and 8% of astrocytes\(^16,43\).

**Effect of AEME on total $[^3\text{H}]$inositol phosphates accumulation.** Hippocampal cells ($1 \times 10^6$ cells/well) were allowed to equilibrate for 10 minutes with a nutrient solution of the following composition (mM): NaCl 157.00; KCl 5.60; CaCl₂ 0.27; MgCl₂ 3.00; NaHCO₃ 1.80; glucose 5.50 (pH 7.0–7.2) at 37°C under constant shaking. Cells were incubated with 5 $\mu\text{Ci}$ of myo-$[^3\text{H}]$inositol (specific activity 18.0 Ci/mmol) for 80 minutes, and lithium chloride (10 mM) for additional 30 minutes. Afterwards, the cells were incubated in the absence (basal level) and in the presence of carbachol (positive control) and AEME ($10^{-8}$ to $10^{-4}$ M) for 10 minutes. Cells were removed from the plate using a 0.1 M NaOH solution and washed three times with nutrient solution, transferred to 2 mL of methanol:chloroform (2:1 v/v) at 4°C and homogenized with a Ultra-Turrax T25 homogenizer at 9500 rpm. Chloroform (0.62 mL) and H₂O (0.93 mL) were added to the homogenate, and the solution was centrifuged for 5 minutes at 1,000 $\times$ g to separate aqueous and organic phases\(^44\). Total $[^3\text{H}]$inositol phosphate was measured as previously described by Ascoli et al.\(^45\) with slight modification. The aqueous layer was neutralized with 0.1 M HCl and then mixed with 1 mL anion-exchange resin (Dowex AX-8, formate form, 200–400 mesh), allowed to equilibrate for 30 minutes at room temperature, under agitation, and centrifugated at 1,000 $\times$ g, for 5 minutes at 4°C. The resin was then washed sequentially, with 10 mM myo-inositol (2 mL) and 5 mM sodium tetraborate/60 mM sodium formate (2 mL). The resin was incubated for 30 minutes at room temperature with 2 mL of 0.1 M formic acid/1 M ammonium formate. The total $[^3\text{H}]$inositol phosphate was eluted and placed in scintillation vials containing OptiPhase HiSafe 3. The amount of radioactivity was determined in scintillation $\beta$-counter (LS 6500 IC, Beckman). Total $[^3\text{H}]$inositol phosphate was expressed as percentage above basal level.
**Effect of AEME on intracellular calcium release.** Hippocampal cells were removed from the plates and previously incubated with Fluoro-AM dye (Invitrogen) dissolved in Neurobasal medium (Gibco®) for one hour. The cells were centrifuged (300 g for 5 minutes at 20 °C), washed with phosphate buffer solution (PBS) without calcium, and plated in microscopy confocal (Zeiss LSM 510, Meta) microplates pretreated with poli-L-lysine. The fluorescence was measured for 90 seconds in the absence and/or in the presence of 10⁻⁴, 10⁻³, and 10⁻² M AEME. After this period, 250 mM KCl was added in order to promote neuronal death and, thus, verify its viability. All measurements were performed with a PBS free-calcium solution. Digitalized images were analyzed through the software Origin 5.

**Effects of AEME on DNA fragmentation.** DNA fragmentation was assessed using propidium iodide following the method described by Lima et al. Hippocampal cells were incubated with 10⁻⁴ or 10⁻³ M AEME for 12 and 24 hours. Briefly, exposed cells were removed from the culture plate and centrifuged (500 g for 5 minutes) and lysed with a fragmentation buffer (100 mg/mL propidium iodide, 0.1% sodium citrate, and 0.1% Triton-X) for 2 hours at room temperature in the dark. DNA fragmentation was analyzed in a flow cytometry (FACSCalibur, Becton Dickinson, CA) using the FL2-A channel (λ_{excitation} = 585 nm and λ_{emission} = 642 nm) and 10 million cells were evaluated per experiment.

**Experiments with CHO cells.** CHO cell culture. Chinese hamster ovary (CHO) cells individually and stably expressing individual rat mAChRs (rM1 to rM5 mAChRs) were purchased from the American Type Culture Collection (ATCC) and cultured according to ATCC recommendations. To generate stable rM2 and rM4 cell lines for use in calcium mobilization assays, rM2 and rM4-expressing cells were stably transfected with a chimeric G protein (G_qi5) using Lipofectamine 2000 (Invitrogen). rM2 and rM4 cells were grown in Ham’s F-12 medium containing 10% heat-inactivated fetal bovine serum (FBS), 2 mM GlutaMAX (Gibco®), 20 mM HEPES, 500 μg/mL G418 sulfate, 200 μg/mL hygromycin B and an antibiotic-antimycotic solution (Gibco®). rM2, rM3 and rM4 mAChR-CHO cells were grown in the same medium without hygromycin B. CHO cells were cultured in a 150 mm-diameter cell culture-treated dishes (Corning®) at 37 °C and 5% CO₂ until complete confluence.

**Membrane preparation and equilibrium radioligand binding assays.** Membranes were prepared from CHO cells stably expressing individual rat mAChRs (rM2 to rM5). Briefly, media was removed, cells were washed with PBS and harvested with PBS into a centrifuge bottle. Cells were spun at 2,000 g for 5 minutes at 4 °C. After harvesting, the supernatant was gently discarded and cold membrane preparation buffer [Hanks' balanced salt solution (HBSS; Invitrogen), 4.17 mM sodium bicarbonate, 20 mM HEPES and 0.5 mM EDTA] was added (10 mL/dish). Cells were then homogenized for 5–10 seconds with a rotary homogenizer (Polytron®) and centrifuged at 20,000g for 20 minutes at 4 °C. The homogenization/centrifugation cycle was repeated twice. After the last centrifugation, the supernatant was gently discarded and the pellet was resuspended in cold membrane preparation buffer (up to 2 mL). All binding reactions were performed in a total volume of 1 mL containing 25 μg of membrane protein. Non-specific binding was determined in the presence of 1 μM atropine. [-[N-methyl-3H]scopolamine ([3H]NMS; GE Healthcare) saturation binding was performed to calculate both the index of the receptor density (B_{max}) and the NMS dissociation constant (NMS K_d) values for each subtype of mAChR. The competition binding reactions were carried out in 96-well deep-well plates with membrane protein in the presence of appropriate concentrations of AEME or vehicle, and 0.1 nM [3H]NMS for 3 hours at room temperature. The equilibrium binding was terminated by rapid filtration using a 96-well harvester (Brandel®). Filters were washed three times with ice-cold harvesting buffer and were dried overnight. Radioactivity was counted using a TopCount NXT (PerkinElmer®) and counts were normalized to the maximal specific binding in the presence of vehicle. Saturation and competition binding data were analyzed using a weighted nonlinear least-squares interactive curve-fitting program GraphPad Prism (GraphPad Prism Software version 5.0). A mathematical model for one or two binding sites was applied. The equilibrium dissociation constant (K_d) and the binding capacity (B_{max}) were determined by competition curves using the Cheng and Prusoff equation.

**Calcium mobilization assay.** CHO cells expressing rM1, rM3 and rM4 were plated at 4 × 10⁴ cells per well, whereas rM2 and rM5 were plated at 6 × 10⁴ cells per well, in standard growth media (as described previously under “cell culture”) in 96-well plates 24 hours before assay and were incubated overnight at 37 °C in 5% CO₂. On the day of the assay, media was removed, cells were washed with calcium assay buffer [Hanks’ Balanced Salt Solution (HBSS; Invitrogen), 20 mM HEPES, 4.17 mM sodium bicarbonate, 2.5 mM probenecid (Sigma), pH 7.4] and maintained in calcium assay buffer containing 2.3 μM Fluoro-AM dye (Invitrogen). Cells were incubated for 45 minutes (37 °C, 5% CO₂) for dye loading. Fluoro-AM dye was removed, cells were washed and replaced with 40 μL of calcium assay buffer. AEME concentration-response curves were determined in single-add experiments. For calcium fluorescence measurements of AEME potency (“double-add” calcium assay), 100 μM AEME was added 20 seconds after the beginning of data collection and increasing concentrations of acetylcholine were added 100 seconds later via Flexstation II (Molecular Devices). Fluorescence measurement continued for a total of 200 seconds of acquisition time using an excitation wavelength of 488 nm and an emission wavelength.
of 525 nm. For Schild analyses, fixed concentrations of AEME (10⁻⁶ to 10⁻¹⁵ M) were added before the acetylcholine concentration-response curve in the same way described previously for the "double-add" calcium assay. All compounds were dissolved in dimethyl sulfoxide (DMSO) and then in calcium assay buffer to obtain a final DMSO concentration of 0.3%. All of the peaks of the calcium response were normalized to baseline and then as a percentage of the maximum acetylcholine response. These values were fit using GraphPad Prism version 5.0 to determine EC₅₀ values and Schild slope and pA₂ in the Schild analyses.⁴³,⁵³,⁵⁴

**Effect of M₁ and M₃ selective mAChR antagonists and the PLC inhibitor on AEME-induced neurotoxicity.** Neuronal viability was evaluated in the hippocampal cell culture using the 3-(4,5-dimethylthiazol-2-yl)-2,5-diphenyltetrazolium bromide (MTT; Sigma) reduction assay, with some modification.⁶ Briefly, after 24 hours of incubation with 10⁻⁶ M AEME in the absence and in the presence of 10 nM pirenzepine hydrochloride (M₁ selective antagonist; Sigma), 10 nM p-F-HH515 (M₁ selective antagonist; Sigma) and the specific inhibitor of different isoforms of PLC U73122 (10 nM) (Sigma), all the medium was removed, and 100 μL of MTT solution, containing 5 mg/mL MTT in PBS and neumosa-1,4,5-trisphosphate production to facilitate Ca²⁺−3 M AEME in the absence and in the presence 3.5 M AEME. A control containing 0.002% DMSO was used, since the iPLC stock solution was previously dissolved in this solvent. The assay was performed in quadruplicate, and the results were expressed as a percentage of the control value.⁶

**Statistical analyses.** Data were analyzed by one-way analysis of variance (ANOVA) followed by Newman-Keuls multiple comparison post-hoc test. Acetylcholine concentration-response curve in the presence of 100 μM AEME was analyzed by Student's t-test. P < 0.05 was considered statistically significant. All data were plotted and analyzed by GraphPad Prism software version 5.0. Data were reported as mean ± SEM.

**References**

1. UNODC (United Nations Office on Drugs and Crime). World Drug Report (2014). United Nations Publication, Sales No. E.14.XI.7. Available at: http://www.unodc.org/documents/wdr2014/World_Drug_Report_2014_web.pdf (Date of access: 04/09/2014).
2. Schwartz, B. G., Rezkal, S. & Kloner, R. A. Cardiovascular effects of cocaine. Circulation 122, 2588–2596 (2010).
3. Mao, J. T. et al. Cocaine inhibits human endothelial cell IL-8 production: the role of transforming growth factor-β. Cell Immunol 181(1), 38–43 (1997).
4. Fromm, C. et al. Cocaine activates platelets and increases the formation of circulating platelet containing microparticles in humans. Heart 83(6), 688–695 (2000).
5. Vale, A., Cucatl, F., Bastos, M., de Pinho, P. G. & Carvalho, M. Contribution of oxidative metabolism to cocaine-induced liver and kidney damage. Carr Med Chem 19(33), 5601–5606 (2012).
6. Riezzo, I. et al. Side effects of cocaine abuse: multiorgan toxicity and pathological consequences. Carr Med Chem 19, 5624–5646 (2012).
7. Goel, N., Pullman, J. M. & Coco, M. Cocaine and kidney injury: a kaleidoscope of pathology. Clin Kidney J 7, 513–517, doi: 10.1093/ckj/sfu092 (2014).
8. Vale, A. Cucatl. Medicine 35(11), 607 (2007).
9. Goldstein, R. A., DesLauriers, C. & Burda, A. M. Cocaine: history, social implications, and toxicity—a review. Dis Mon 55(1), 6–38, doi: 10.1016/j.disamonth.2008.10.002 (2009).
10. Pires, A. et al. Repeated inhalation of crack-cocaine affects spermatogenesis in young and adult mice. Inhal Toxicol 24(7), 439–446, doi: 10.3109//08958378.2012.648450 (2012).
11. Fandino, A. S., Toennes, S. W. & Kauer, G. F. Studies on hydrolytic and oxidative metabolic pathways of anhydroecgonine methyl ester (methylecgonidine) using microsomal preparations from rat organs. Chem Res Toxicol 15, 1543–1548 (2002).
12. Paul, B. D., Lalani, S., Bosy, T., Jacobs, A. J. & Huestis, M. A. Concentration profiles of cocaine, pyrolytic methyl ecgonidine and thirteen metabolites in human blood and urine: determination by gas chromatography–mass spectrometry. Biomed Chromatogr 19, 677–688 (2005).
13. Erzouki, H. K., Allen, A. C., Newman, A. H., Goldberg, S. R. & Schindler, C. W. Effects of cocaine, cocaine metabolites and cocaine pyrolysis products on the pyrolysis products of the pyrolysate of cocaine. Life Sci 57, 1861–1868 (1995).
14. Wooll, J. F., Huang, L., Ishiguro, Y. & Morgan, J. P. Negative inotropic effect of methylecgonidine, a major product of cocaine base pyrolysis, on ferret and human myocardium. J Cardiovasc Pharmacol 30, 352–359 (1997).
15. Seyfedwiler, K. B., Plessinger, M. A., Shojaie, J., Wood, R. W. & Kwong, T. C. Pharmacokinetics and pharmacodynamics of methylecgonidine, a crack cocaine pyrolysate. JPET 307, 1179–1187 (2003).
16. Garcia, R. C. T. et al. Neurotoxicity of anhydroecgonine methyl ester, a crack cocaine pyrolysis product. Toxicol Sci 128, 223–234, doi: 10.1093/toxsci/kfs140 (2012).
17. Egan, R. M. Muscarinic receptor subtype pharmacology and physiology. Prog Med Chem 43, 105–136 (2005).
18. Vistoli, G. Docking analyses on human muscarinic receptors: Unveiling the subtypes peculiarities in agonists binding. Biorg Med Chem 16(6), 3049–3058 (2008).
19. Oki, T. et al. Quantitative analysis of binding parameters of [3H][N-methylscopolamine in central nervous system of muscarinic acetylcholine receptor knockout mice. Brain Res Mol Brain Res 133(1), 6–11 (2005).
20. Cardoso, C. C., Ricardo, V. P., Frassa-Filho, R., Porto, C. S. & Abdalla, F. R. Synaptic activity augments muscarinic acetylcholine receptor-stimulated inositol 1,4,5-trisphosphate production to facilitate Ca²⁺−release in hippocampal neurons. J Biol Chem 279(47), 49036–49044 (2004).
22. Levey, A. I. Immunological localization of M1-M5 muscarinic acetylcholine receptors in peripheral tissues and brain. Life Sci 52, 441–448 (1993).
23. Bradley, K. N., Rowan, E. G. & Harvey, A. L. Effects of muscarinic toxins MT2 and MT7, from green mamba venom, on m1, m3 and m5 muscarinic receptors expressed in Chinese hamster ovary cells. Toxicol 41, 207–215 (2003).
24. Shih, Y.-T. et al. Arecoline, a major alkaloid of the areca nut, causes neurotoxicity through enhancement of oxidative stress and suppression of the antioxidant protective system. Free Radic Biol Med 49, 1471–1479 (2010).
25. Dong, G. Z., Kameyama, K., Rinken, A. & Haga, T. Ligand binding properties of muscarinic acetylcholine receptor subtypes (m1-m5) expressed in baculovirus-infected insect cells. J Pharmacol Exp Ther 274, 378–384 (1995).
26. Galluzzi, L., Blomgren, K. & Kroemer, G. Mitochondrial membrane permeabilization in neuronal injury. Nat Rev Neurosci 10(7), 481–494, doi: 10.1038/nrn2665 (2009).
27. Seo, M. D., Enomoto, M., Ishiyama, N., Stathopoulos, P. B. & Ikura, M. Structural insights into endoplasmic reticulum stored calcium regulation by inositol 1,4,5-trisphosphate and ryosynode receptors. Biochim Biophys Acta doi: 10.1016/j.bbalip.2014.11.023 [Epub ahead of print] (2014).
28. Wodja, U., Salmina, E. & Kuznicki, J. Calcium ions in neuronal degeneration. Life 60(9), 575–590 (2008).
29. Jerusalinsky, D. et al. Muscarinic toxin selective for m4 receptors impairs memory in the rat. Neuroreport 9(7), 1407–1411 (1998).
30. Servent, D. & Fruchart-Gaillard, C. Muscarinic toxins: tools for the study of the pharmacological and functional properties of muscarinic receptors. J Neurochem 109, 1193–1202, doi: 10.1111/j.1471-4159.2009.06092.x (1999).
31. Bashkatova, V., Hornick, A., Vanin, A. & Prast, H. Antagonist of M1 muscarinic acetylcholine receptor prevents neurotoxicity of 6-OHDA in rat striatum via nitric oxide pathway. J Neurochem 139, 172–176, doi: 10.1166/annals.1432.004 (2008).
32. Moncada, S., Palmer, R. M. & Higgs, E. A. Nitric oxide: physiology, pathophysiology, and pharmacology. Pharmacol Rev 43(2), 109–142 (1991).
33. Garthwaite, J. Glutamate, nitric oxide and cell-cell signalling in the nervous system. Trends Neurosci 14(2), 60–67 (1991).
34. Prast, H. & Philippu, A. Nitric oxide as modulator of neuronal function. Prog Neurobiol 64(1), 51–68 (2001).
35. Cunha-Oliveira, T. et al. Mitochondrial dysfunction and caspase activation in rat cortical neurons treated with cocaine or amphetamine. Brain Res 1089, 44–54 (2006).
36. Cunha, P., Nicastri, S., Gomes, L. P., Moine, R. M. & Peluso, M. A. Neuropsychological impairments in crack cocaine-dependent inpatients: preliminary findings. Rev Bras Psiquiatr 26(2), 103–106 (2004).
37. Banker, G. A. & Cowan, W. M. Rat hippocampal neurons in dispersed cell culture. Brain Res 126, 397–442 (1977).
38. Huettner, J. E. & Baughman, R. W. Primary culture of identified neurons from the visual cortex of postnatal rats. J Neurosci 6, 3044–3060 (1986).
39. Jahn, C. E. & Stevens, C. F. Glutamate activates multiple single channel conductances in hippocampal neurons. Nature 325, 522–525 (1987).
40. Silva, R. F. M. et al. Dissociated primary nerve cell cultures as models for assessment of neurotoxicity. Toxicol Lett 163, 1–9 (2006).
41. Brewer, G. J., Torricelli, J. R., Evege, E. K. & Price, P. J. Optimized survival of hippocampal neurons in B27-supplemented Neurobasal a, a new serum-free medium combination. J Neurosci Res 35, 567–576 (1993).
42. Rigoni, M. Snake presynaptic neurotoxins with phospholipase A2 activity induce punctate swellings of neurites and exocytosis of synaptic vesicles. J Cell Sci 117, 3561–3570 (2004).
43. de Carvalho, N. D. et al. Neurotoxicity of coral snake phospholipases A2 in cultured rat hippocampal neurons. Brain Res 1552, 1–16, doi: 10.1016/j.brainres.2014.01.008 (2014).
44. Pereira, R. T., Porto, C. S., Godinho, R. O. & Abdalla, F. M. Effects of estrogen on intracellular signaling pathways linked to activation of muscarinic acetylcholine receptors and on acetylcholinesterase activity in rat hippocampus. Biochem Pharmacol 75(9), 1827–1834, doi: 10.1016/j.bcp.2008.01.016 (2008).
45. Ascoli, M., Pigñana, O. P. & Segallof, D. L. The inositol phosphate/diaclylglycerol pathway in MA-10 Leydig tumor cells. Activation by arginine vasopressin and lack of effect of epidural growth factor and human choriongonadotropin. J Biol Chem 264(12), 6674–6681 (1989).
46. Ban, J. Y. et al. Neuroprotective effect of oxyresveratrol from smilacis chine rhizome on amyloid Beta protein (25–35)-induced neurotoxicity in cultured rat cortical neurons. Bio Pharm Bull 29(12), 2419–2424 (2006).
47. Lima, R. M. et al. Cytotoxic effects of catechol to neuroblastoma N2a cells. Gen Physiol Biophys 27(4), 306–314 (2008).
48. Sheffler, D. J. et al. A novel selective muscarinic acetylcholine receptor subtype 1 agonist reduces seizures without impairing hippocampal-dependent learning. Mol Pharmacol 76, 356–368; doi: 10.1124/mol.109.056531 (2009).
49. Shirley, J. K. et al. An allosteric potentiator of M4 mAChR modulates hippocampal synaptic transmission. Nat Chem Biol 4(1), 42–50 (2008).
50. Shirley, J. K. et al. A selective allosteric potentiator of the M4 muscarinic acetylcholine receptor increases activity of medial prefrontal cortical neurons and restores impairments in reversal learning. J Neurosci 29(45), 14271–14286; doi: 10.1523/JNEUROSCI.3930-09.2009 (2009).
51. Munson, P. J. & Rodbard, D. Ligand: a versatile computerized approach for characterization of ligand-binding system. Anal Biochem 107, 220–239 (1980).
52. Cheng, Y. C. & Prassof, W. H. Relationship between the inhibition constant (Ki) and the concentration of inhibitor which causes 50 percent inhibition (IC50) of an enzymatic reaction. Biochem Pharmacol 22, 3099–3108 (1973).
53. Arunlakshana, O. & Schild, H. O. Some quantitative uses of drug antagonists. Br J Pharmacol Chemother 14(1), 48–58 (1959).
54. Brady, A. E. et al. Centrally active allosteric potentiators of the M4 muscarinic acetylcholine receptor reverse amphetamine-induced hyperlocomotor activity in rats. J Pharmacol Exp Ther 327, 941–953; doi: 10.1124/jpet.108.140350 (2008).
55. Konigame, V.C. et al. Estrogen receptors mediate total inositol phosphate accumulation in the rat endometrium. Prog Neurobiol 76, 1582–1589 (2011).
56. Doods, H. N., Willim, K. D., Bodekke H. W. & Entzeroth, M. Characterization of muscarinic receptors in guinea-pig uterus. Eur J Pharmacol 250(2), 223–230 (1993).

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Author Contributions
R.C.T.G., M.R.L.S. and T.M. conceived and designed the project. L.M.M.D. and R.C.T.G. performed all the experiments with rat primary hippocampal cell culture, supervised by M.R.L.S., S.C.A. and F.M.F.A. The DNA fragmentation assay was supervised by R.G. The experiments with CHO cells were performed and analyzed by R.C.T.G, under C.M.N. and P.J.C. supervision. J.L.C., M.Y. and R.C.T.G. provided the drug and synthesized AEME for the experiments. M.A.A.S. and M.S.B.U. performed and analyzed the experiments with M₁ and M₃ selective antagonists, under R.C.T.G. supervision. T.M. supervised the project and M.R.L.S. co-supervised it. R.C.T.G., L.H.T., F.M.F.A., S.C.A., M.Y., C.M.N, P.J.C., R.C., M.R.L.S. and T.M. wrote and edited the manuscript.

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