Augmentation of catecholamine release elicited by an *Eugenia punicifolia* extract in chromaffin cells


1Instituto Teófilo Hernando, Departamento de Farmacología, Facultad de Medicina, Universidad Autónoma de Madrid, Spain,
2Servicio de Farmacología Clínica, Hospital Universitario de la Princesa, Madrid, Spain,
3Departamento de Biología Celular e Molecular, Universidade Federal Fluminense, Brazil,
4Departamento de Bromatologia, Faculdade de Farmácia, Universidade Federal Fluminense, Brazil,
5Programa de Pós-graduação em Química Orgânica, Instituto de Química, Universidade Federal Fluminense, Brazil,
6Departamento de Farmácia e Administração Farmacêutica, Faculdade de Farmácia, Universidade Federal Fluminense, Brazil.

**Abstract:** Plant extracts of *Eugenia punicifolia* (Kunth) DC., Myrtaceae, are used in Amazon region of Brazil to treat diarrhea and stomach disturbances, and as hypoglycemic medicine. We have recently shown that an aqueous extract of *E. punicifolia* augmented cholinergic neurotransmission in a rat phrenic nerve-diaphragm preparation. In this study, we investigated the effects of an *E. punicifolia* dichloromethane extract (EPEX) in a neuronal model of cholinergic neurotransmission, the bovine adrenal chromaffin cell. EPEX augmented the release of catecholamine triggered by acetylcholine (ACh) pulses but did not enhance ACh-evoked inward currents, which were inhibited by 30%. Since EPEX did not cause a blockade of acetylcholinesterase or butyrylcholinesterase, it seems that EPEX is not directly activating the cholinergic system. EPEX also augmented K+-elicited secretion without enhancing the whole-cell inward calcium current. This novel and potent effect of EPEX in enhancing exocytosis might help to identify the active component responsible for augmenting exocytosis. When elucidated, the molecular structure of this active principle could serve as a template to synthesise novel compounds to regulate the exocytotic release of neurotransmitters.

**Keywords:** calcium channels, catecholamine release, chromaffin cells, *Eugenia punicifolia*, nicotinic receptors

**Introduction**

Historically, there is a long tradition investigating the pharmacological profile of plant extracts and looking for new active compounds with potential therapeutic interest. This strategy has given rise to many new compounds that lead to the synthesis of new chemical entities to treat several diseases (Marcaurelle & Johannes, 2008). Our interest is in determining natural product extracts with pharmacological activity, and we recently became interested in studying the cholinergic effects of *Eugenia punicifolia* (Kunth) DC., Myrtaceae, a plant that grows in the Amazonia forests of Brazil. We studied the effects of a 5% (w/v) aqueous extract of *E. punicifolia* on cholinergic neurotransmission at the muscle endplate (rat phrenic nerve-diaphragm preparation), and we found that this extract augmented neurotransmission mediated by nicotinic receptors for acetylcholine (nAChR) (Grangeiro et al., 2006).

Our laboratory has a long-standing interest in the search for compounds with potential capabilities for cholinergic neurotransmission enhancement. With this enhancement, we expect these compounds to improve cognition in Alzheimer’s disease (AD) patients, as is the case for galantamine, rivastigmine or donepezil, three inhibitors of acetylcholinesterase (AChE) and...
butyrylcholinesterase (BuChE) (Marco-Contelles et al., 2006; Marco & do Carmo Carreiras, 2006). For the last decade, we have been designing and synthesising novel hybrid compounds with multi-target profiles, such as inhibiting AChE and/or BuChE, the modulation of nAChR and neuronal calcium (Ca$^{2+}$) signalling, or neuroprotection against different neurotoxic stimuli (de los Rios et al., 2002; Leon et al., 2005; Orozco et al., 2006; de los Rios et al., 2010). In this context, we hypothesised that the/Eugenia punicifolia extract that augmented the release of catecholamines from bovine chromaffin cells stimulated with either acetylcholine (ACh) or with high K$^+$ concentrations. In elucidating the mechanisms involved in such augmentation, we also investigated the effects of EPEX on ACh-elicited inward currents ($I_{ACh}$), Na$^+$ channel currents ($I_{Na}$), Ca$^{2+}$ channel currents ($I_{Ca}$) and cytosolic Ca$^{2+}$ concentrations ([Ca$^{2+}$]). Furthermore, we also investigated EPEX effects on the activity of both purified AChE and BuChE enzymes.

Material and Methods

Plant material and preparation of crude extracts

Eugenia punicifolia (Kunth) DC., Myrtaceae, was kindly supplied by Centro de Instrução de Guerra na Selva (Manaus-AM, Brazil) and was identified at Museu Nacional, UFRJ (Brazil), where voucher specimens were kept for future reference. The official authorisation to investigate this plant was given by Instituto Brasileiro de Meio Ambiente e Recursos Renováveis, Brazil, and it was registered under the number 16602-1. The plant was successively extracted at room temperature with solvents of increasing polarity beginning with n-hexane, then dichloromethane and finally methanol. The extracts were concentrated under reduced pressure to yield oily (hexane extract) or solid (dichloromethane and methanol extracts) residues, and they were stored at 4 °C. Stock solutions were prepared at 1 mg/mL in 10$^{-2}$ M DMSO. Appropriate dilutions of the solutions were made for each experiment. All experiments in this study used the dichloromethane extract (EPEX). Appropriate controls with this solvent were made and no significant effects were observed in the presence of DMSO in any of the experimental protocols (see below) used in this study.

Measurement of AChE activity

The effect of EPEX on acetylcholinesterase (AChE) activity was determined following the method of Ellman et al. (Ellman et al., 1961) using AChE from Electrophorus electricus (Electric eel; eeAChE) and acetylthiocholine iodide (0.35 mM) as substrate. The reaction took place in a final volume of 3 mL and consisted of a phosphate buffer solution at pH 8.0 containing 0.035 U of eeAChE and 0.35 mM of 5,5'-dithiobis-2-nitrobenzoic acid (DTNB), at 25 °C. The yellow anion 5-thio-2-nitrobenzoic acid is produced as the enzymatic reaction proceeds. After a 10-min preincubation of the enzyme with different EPEX concentrations, the substrate was added and the enzymatic reaction was allowed to proceed for 15 additional min. A sample without EPEX was always present as a negative control to determine 100% of enzymatic activity, along with a sample containing 100 nM tacrine as a reference compound (Guillou et al., 2000). After the 15-min incubation period, the production of colour as an indication of enzymatic activity was evaluated by measuring absorbance at 410 nm in a spectrophotometer plate reader (iEMS Reader MF, Labsystems).

Measurement of BuChE activity

The effect of EPEX on butyrylcholinesterase (BuChE) was also determined following the method of Ellman et al. (Ellman et al., 1961) using BuChE from horse serum (eqBuChE) and butyrylthiocholine iodide (0.5 mM) as the substrate. The reaction took place at a final volume of 3 mL and consisted of a phosphate buffer solution at pH 8.0 containing 0.05 U of eqBuChE and 0.35 mM DTNB. The yellow anion 5-thio-2-nitrobenzoic acid is produced as the enzymatic reaction proceeds. The enzyme was pre-incubated for 10 min with different concentrations of EPEX. After this time, the substrate was added and the enzymatic reaction was allowed to proceed for an additional 15 min, at 25 °C. A sample without EPEX was always present, as a negative control, to determine the 100% of enzymatic activity, along with a sample containing 100 nM tacrine as a reference compound (Guillou et al., 2000). After the 15-min incubation period, the production of colour as an indication of enzymatic activity was evaluated by measuring absorbance at 410 nm in a spectrophotometer plate reader (iEMS Reader MF, Labsystems).
Isolation and culture of adrenal medulla chromaffin cells

Bovine adrenal glands were obtained from a local slaughterhouse. Adrenal medulla chromaffin cells were isolated following standard methods (Livett, 1984) with some modifications (Moro et al., 1990). Cells were suspended in DMEM supplemented with 5% foetal calf serum, 10 μM cytosine arabinoside, 10 μM fluoro-deoxyuridine, 50 IU/mL penicillin, and 50 μg/mL streptomycin. For the patch-clamp studies, cells were plated on 1-cm diameter glass coverslips at a low density (5x10⁴ cells per coverslip). For the catecholamine release measurements, cells were plated on 5-cm diameter Petri dishes at 5x10⁶ cells per dish. To study changes in cytosolic Ca²⁺ levels ([Ca²⁺]c), cells were plated at a density of 2x10⁵ cells/well in 96-well black dishes.

On-line measurements of catecholamine release

Bovine chromaffin cells were scraped off carefully from the bottom of the Petri dishes with a rubber policeman and centrifuged at 800 rpm for 10 min. The cell pellet was resuspended in a 200 μL Krebs-HEPES solution at pH 7.4 (in mM: 144 NaCl, 5.9 KCl, 1.2 MgCl₂, 11 glucose, 10 N-(2-hydroxyethyl) piperazine-N'-(2-ethanesulfonic acid) (HEPES)). Cells were placed in a microchamber for their superfusion at the rate of 2 mL/min at 37 °C in the Krebs-HEPES solution. Under these conditions, the cell superfusion fluid emanating from the microchamber was measured with a thermistor probe and showed a temperature of 35 °C. For the detection of released catecholamines, the liquid flowing from the superfusion chamber reached an electrochemical detector (model CH-9100; Metrohm AG, Herisau, Switzerland) equipped with a glassy carbon working electrode, an Ag/AgCl reference electrode and a gold auxiliary electrode. Catecholamines are oxidised at +0.65 V and the oxidation current was recorded on a PC computer (Borges et al., 1986). Cells were stimulated to secrete with short pulses (5 s) of a Krebs-HEPES solution containing either 35 or 70 mM K⁺ or 100 μM acetylcholine, in the absence or presence of different extract concentrations (See Results). Solutions were rapidly exchanged through electrovalves operated by a computer.

Patch-clamp current measurements and analysis

Inward currents through voltage-gated Ca²⁺ channels (IₐCa), voltage-gated Na⁺ (IₐNa), and nAChR (IₐACH) were recorded using the whole-cell configuration of the patch-clamp technique (Hamill et al., 1981). Whole-cell recordings were made with fire-polished electrodes (resistance 2-5 MΩ when filled with the standard intracellular solutions) that were mounted on the headstage of an EPC-9 patch-clamp amplifier (Heka Electronic, Lambrecht, Germany), allowing cancellation of capacitative transients and compensation of series resistance. Data were acquired with a sample frequency of 20 kHz by using PULSE 8.74 software (Heka Elektronik, Lambrecht, Germany). Linear leak and capacitative components were subtracted by using a P/4 protocol and series resistance was compensated by 90%. The data analysis was performed with Igor Pro (Wavemetrics, Lake Oswego, OR) and PULSE programs (Heka Elektronik).

Coverslips containing the cells were placed on an experimental chamber mounted on the stage of a Nikon Diaphot inverted microscope. Cells were continuously superfused with a control Tyrode solution at pH 7.4 containing (in mM): 137 NaCl, 1 MgCl₂, 2 CaCl₂, 10 HEPES/NaOH. Once the patch membrane was ruptured and the whole-cell configuration of the patch-clamp technique had been established, the cell was locally, rapidly and constantly superfused with an extracellular solution of similar composition to the chamber solution, but containing nominally 0 mM Ca²⁺ to measure IₐNa, 2 mM Ca²⁺ to measure IₐCa, or 10 mM Ba²⁺ (instead of Ca²⁺) to measure IₐACH (see Results for specific experimental protocols). Cells were internally dialysed with an intracellular solution containing (in mM): 100 CsCl, 14 EGTA, 20 TEA.Cl, 10 NaCl, 5 Mg-ATP, 0.3 Na-GTP, and 20 HEPES/CsOH (pH 7.3). The external solutions were rapidly exchanged using electronically driven miniature solenoid valves coupled to a multi-barrel concentration-clamp device, the common outlet of which was placed within 100 μm of the cell to be patched. The flow rate was 1 mL/min and regulated by gravity.

Cells were held at -80 mV; Na⁺ currents were generated by 15 ms depolarizing pulses to -10 mV, Ba²⁺ currents were generated by 50 ms depolarizing pulses at 0 or +10 mV; nAChR currents were generated by the application of 250 ms ACh pulses (100 μM). All experiments were performed at room temperature (24±2 °C) on cells from 2 to 4 days after culture.

Measurement of [Ca²⁺]c

Cells were plated at a density of 2x10⁵ cells per well into 96-well plates, and the experiments were performed 48 h later. Cells were loaded with a Krebs-HEPES solution at pH 7.4 (in mM: 144 NaCl, 5.9 KCl, 1.2 MgCl₂, 2 CaCl₂, 11 D-glucose, and 10 HEPES) containing 10 μM fluo-4-AM and 0.2% pluronic acid. The cells were incubated in this solution for 45 min at 37 °C in the dark. After this incubation period, cells were washed twice with the Krebs-HEPES solution at room temperature in the dark. Changes in fluorescence (excitation 485 nm, emission 520 nm) were measured using a fluorescent plate reader (Fluostar, BMG Labtech).
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Offenburg, Germany). Basal levels of fluorescence were monitored before adding the stimulation solution (containing 100 μM acetylcholine) with an automatic dispenser. After stimulation of the cells, changes in fluorescence were measured for 40 s. To normalise fluo-4 signals, responses from each well were calibrated by measuring maximum and minimum fluorescence values. At the end of each experiment, 3% Triton X-100 (Fmax) was added followed by 0.2 M MnCl2 (Fmin). Data were calculated as a percentage of Fmax-Fmin.

**Data analysis**

The results are presented as mean±SE. The measurements of peak current and charge that flow through the channel after its activations were calculated with the program Clampfit 5.03. IC50 values were estimated through non-linear regression analysis using Origin software (OriginLab Corporation, Northampton, USA). Comparisons between means of group data were performed by one-way analysis of variance (ANOVA) followed by the Duncan post-hoc test when appropriate. p≤0.05 was taken as the limit of significance.

**Results**

**Effects of EPEX on catecholamine release elicited by repeated ACh pulsing in chromaffin cells**

Because EPEX was previously shown to enhance neurotransmission at the muscle endplate (rat phrenic nerve-diaphragm preparation) (Grangeiro et al., 2006), we first explored whether the extract had the capability of augmenting the exocytotic release of catecholamines. Thus, an electrochemical detector working under the amperometric mode was used to measure the real-time rate of catecholamine release from bovine chromaffin cell populations (around 5 million cells per experiment). These cells were trapped in a microchamber and fast-perfused with a Krebs-HEPES solution containing 2 mM Ca2+. In each experiment, cells were initially perfused for a 5 to 10-min period to allow the stabilisation of basal secretion (around 10 nA). Accumulated experience in our laboratory shows that these cells give highly reproducible amperometric secretory responses upon repeated pulses applied at regular intervals, using solutions containing supramaximal concentrations of ACh (Santos et al., 2001; Cuchillo-Ibáñez et al., 2002; Tapia et al., 2009). We used 3-s pulses of solutions containing 100 μM ACh (100 ACh), which in a previous study were found to fully deplete the ready-releasable vesicle pools under these experimental conditions and produce a maximal peak secretory response (Cuchillo-Ibáñez et al., 2002).

Figure 1A shows a prototypical experiment using repeated 3-s 100 ACh pulses given at 5-min intervals. Note that the peak of the initial secretory spike of 200 nA was maintained fairly constant throughout the experiment. Using this protocol, we surprisingly found that at the low concentration of 0.1 ng/mL, EPEX gradually augmented the ACh-evoked, catecholamine release response, reaching an amplitude 2.5 fold higher than the initial response (Figure 1B). As shown in Figure 1B, basal catecholamine release was not affected by EPEX perfusion; however, upon EPEX washout, the response quickly returned to initial levels and was enhanced once more upon reperfusion of the extract. Pooled data plotted in Figure 1C shows EPEX causing a gradual increase in catecholamine secretion, which was maintained for 15 min and reversed upon EPEX washout. However, we did not find an obvious concentration-dependence for the secretion-enhancing effects of EPEX that seemed to augment this response even more at 0.1 ng/mL than at 1 and 5 ng/mL, although there were no statistical differences between the three concentrations (Figure 1D).

**Effects of EPEX on ACh-evoked inward currents**

In trying to elucidate the mechanism involved in the enhancement of ACh-evoked catecholamine release elicited by EPEX, we explored the possibility that the extract was modifying the ACh-evoked inward currents (I\text{ACh}). Thus, we performed experiments in voltage-clamped cells using the whole-cell configuration of the patch-clamp technique.

Figure 2A shows an example of the original I\text{ACh} obtained upon repeated application of 100 μM ACh pulses to the same cell. In this cell, the initial I\text{ACh} peak amplitude was 968 pA. In subsequent ACh pulses, I\text{ACh} was maintained fairly well. Figure 2B shows a prototype experiment in which the initial I\text{ACh} was reduced by 37% in the presence of 1 ng/mL EPEX, and this current reduction was readily reversible upon the extract washout. Figure 2C shows the pooled averaged results of experiments performed in 34 cells. At EPEX concentrations within the range of 0.01-1 ng/mL, I\text{ACh} progressively decreased to...
63% of the initial current, and higher concentrations (10-100 ng/mL) did not produce additional blockade. EPEX washout produced a partial current recovery.

We also tested whether EPEX could affect other ionic channels by measuring the TTX-sensitive sodium current (I_{Na}) in voltage-clamped (-80 mV) cells. We found no effect of EPEX on such current at 0.1 and 1 ng/mL (data not shown).

**Effects of EPEX on the activities of AChE and BuChE**

Since the effects of EPEX in enhancing ACh-stimulated catecholamine release did not seem to be associated with a direct action on nAChR, we wanted to know whether EPEX affected the activities of two enzymes involved in the rapid degradation of ACh at synaptic sites, AChE and BuChE (Kutty, 1980; Buchwald, 2001). We performed experiments on the basis of the following previous findings: i) augmentation of synaptic transmission at the muscle endplate elicited by EPEX could be due to inhibition of AChE and BuChE (Grangeiro et al., 2006); ii) drugs inhibiting AChE and BuChE have proven to have beneficial effects in Alzheimer’s disease patients (Schmidt et al., 2008; van Marum, 2008) and iii) inhibition of AChE and BuChE could explain the EPEX-elicited augmentation of ACh-evoked catecholamine release responses.

Figure 3 shows the concentration response curves for enzyme inhibition by tacrine, a well-known inhibitor of both AChE and BuChE (Summers et al., 1986). As expected, tacrine inhibited both enzymes in a concentration-dependent manner, with an IC50 for BuChE at 2.0 nM, and 25 nM for AChE. In contrast, the concentrations of EPEX that augmented ACh-evoked

![Figure 1](image-url)
secretion (0.1-1 ng/mL, Figure 1 B and D) or even 1000-fold higher concentrations of EPEX, did not affect the activities of AChE or BuChE.

**Figure 2.** EPEX causes a concentration-dependent mild inhibition of the inward whole-cell current generated by ACh pulses. Cells were voltage-clamped at -80 mV. Upon breaking into the cell, ACh pulses (100 μM, 250 ms) were sequentially applied to each individual cell at 60 s intervals. (A) An example cell stimulated with ACh pulses that produced inward $I_{ACh}$. (B) An example cell showing the $I_{ACh}$ traces obtained before (initial), 3 min after perfusion with EPEX, and 3 min after extract washout. (C) Pooled results obtained in cells that were sequentially stimulated with ACh pulses before (initial $I_{ACh}$) and 3 min after perfusion with increasing concentrations of EPEX (bottom horizontal bar). Each concentration was tested in a different cell. Data were normalised as the percent of $I_{ACh}$ amplitude obtained initially and are represented as mean±SE of 34 cells from five different cultures. *$p<0.05$, **$p<0.01$, and ***$p<0.001$, with respect to initial $I_{ACh}$ amplitude.

**Effects of EPEX on catecholamine release responses triggered by repeated application of high-K+ pulses**

The sequence of events leading to the coupling between ACh stimulation and the secretory response (Douglas, 1968) include cell depolarization, opening of voltage-activated Ca$^{2+}$ channels (VACC) and exocytosis (Garcia et al., 2006). Because EPEX did not seem to augment secretion through a cholinergic-mediated pathway, we explored the possibility that the extract could act on VACC to enhance secretion. We therefore triggered secretion with direct depolarization of the cell membrane with a solution containing 70 mM K$^+$ (70K$^+$). At this concentration, the bovine chromaffin cell membrane potential depolarizes to near 0 mV (Orozco et al., 2006) and opens all VACC (Garcia et al., 2006).

**Figure 3.** EPEX did not affect the activities of acetylcholinesterase (AChE) and butyrylcholinesterase (BuChE). Purified AChE from *Electrophorus electricus* or BuChE from horse serum were incubated with EPEX at the concentrations indicated on the abscissa. Tacrine was used as a positive control. In each individual experiment, the maximal enzyme activities (100%) were determined in the presence of vehicle, and enzyme activities were determined in the presence of EPEX or tacrine, expressed as the percent of maximal activity. Data are represented as mean±SE from eight wells from at least two different experiments.

**Figure 4A** shows the secretory spikes obtained in a representative experiment with cells challenged with repeated 5-s, 70 mM K$^+$ pulses given at 3 min intervals. The initial secretory responses of 200 nA showed an amplitude similar to spikes elicited by ACh pulses (Figure 1A). These responses gradually declined with time to amplitudes of around 70% of the initial responses. Similar to ACh, EPEX also augmented the K$^+$-evoked responses, and this increase developed more rapidly than with ACh, reaching about 2.5 times the initial spike amplitude (Figures 1B and 4B). The reason for such a difference may be explained because we used 5 ng/mL EPEX in the K$^+$ experiments, a concentration 50-fold higher than that used in the ACh experiments. Figure 4C shows the pooled results of the effects of EPEX on K$^+$-induced catecholamine secretion. The augmentation of the secretion elicited by EPEX seems to be gradual, similar to that of ACh (Figures 1C and 4C). Figure 4D shows pooled results of experiments performed with increasing concentrations of EPEX, and the 0.1 and 1 ng/mL concentrations that enhanced ACh responses had no effect on the K$^+$ responses. In contrast, 5 ng/mL of EPEX almost doubled the initial K$^+$ secretory response.
Effects of EPEX on inward currents through Na⁺ and Ca²⁺ channels

High K⁺ concentrations will induce a cell membrane depolarization allowing the opening of voltage-gated Na⁺ channels and VACC to enhance Ca²⁺ entry, [Ca²⁺], and catecholamine secretion. Thus, EPEX-elicited enhanced secretion could be due to augmentation of Ca²⁺ entry through VACC. The most direct way of testing this possibility was measuring the whole-cell inward currents through these channels. Thus, we performed experiments aimed to test the possible effects of EPEX on voltage-gated Na⁺ and Ca²⁺ channels by recording whole-cell inward currents through these channels.

Figure 5A shows original recordings of I_{Na} (left panel) and I_{Ba} (right panel) elicited by 15 or 50 ms depolarizing pulses (see protocol on top or the original recordings), in the absence and/or the presence of EPEX (0.1 and 1 ng/mL). The peak Na⁺ current density in cells from different batches was 42.15±6.65 pA/pF (n=8) and the peak Ba²⁺ current density in cells from different batches was 45.63±4.13 pA/pF (n=15). Averaged results obtained with these experimental protocols showed that I_{Na} was not significantly modified in the presence of EPEX (left panel in Figure 5B), and only a mild (around 10%), not significant, decrease of I_{Ba} at 0.1-1 ng/mL EPEX was observed. Figure 5C shows current-voltage curves generated by increasing voltage-step depolarizations. Peak current was reached at 0-+10 mV and was reduced by 5-10% in the presence of 1 ng/mL EPEX; however, no shift of the current-voltage (I-V) curve was found during cell treatment with the extract.

Figure 4. EPEX augmented the release of catecholamines from cells repeatedly stimulated with K⁺-enriched solutions. Cells were initially perfused with Krebs-Hepes solution and, once baseline secretion was stable, cells were challenged for 5 s every 3 min with a solution containing 70 mM K⁺ (with low Na⁺) (dots at the bottom). (A) An example cell showing initial secretion spikes of 200 nA. This spike amplitude was maintained during the first five initial pulses and declined by 20% at the end of the experiment. (B) An example cell perfused with EPEX during the period indicated by the bottom horizontal bar. (C) Pooled data obtained in different experiments made with the protocols shown in panels A (control) and B (EPEX-treated cells). (D) The effects of increasing concentrations of EPEX on K⁺-evoked secretion. Data in panels C and D are normalised as a percent of the initial response in each individual experiment and are represented as mean±SE. ***p<0.001, with respect to the corresponding time in control cells (panel C) and **p<0.01, with respect to the initial secretion (panel D).
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Figure 5. EPEX did not significantly modify the whole-cell inward currents through voltage-gated sodium ($I_{Na}$) and calcium ($I_{Ca}$) channels. Cells were voltage-clamped at -80 mV and step-depolarising pulses (15 ms to -10 mV for $I_{Na}$, and 50 ms to +10 mV for $I_{Ca}$; see protocol on top of figure) were sequentially applied at 10-s intervals. Panel (A) shows original recordings of $I_{Na}$ (left panel) and $I_{Ca}$ (right panel) recorded in the absence and/or the presence of EPEX (0.1 and 1 ng/mL). In panel (B) averaged results obtained with these experimental protocols show that $I_{Na}$ was not significantly modified in the presence of EPEX (left panel; n=8), and only a mild (around 10%; right panel; n=15), not significant, decrease of $I_{Ca}$ at 0.1-1 ng/mL EPEX was observed. Panel (C) shows current-voltage curves generated by increasing voltage-step depolarizations. Peak current was reached at 0-+10 mV and was reduced by 5-10% in the presence of 1 ng/mL EPEX (n=11); No shift of the current-voltage (I-V) curve was found during cell treatment with the extract. Data are mean±SE.
Effects of EPEX on the elevations of cytosolic calcium concentrations ([Ca^{2+}]_c) elicited by ACh

In bovine chromaffin cells, ACh pulses cause depolarisation and action potentials that give rise to transient [Ca^{2+}]_c elevations (de Diego et al., 2008). Such Ca^{2+} signals are the result of Ca^{2+} entry through nAChR via VACC, or from Ca^{2+} release from the endoplasmic reticulum (Garcia et al., 2006). Thus, we tested whether EPEX could alter this ACh-elicited Ca^{2+} signal to explain its ability to augment depolarization-evoked secretion.

Fluo-4-loaded cells were challenged with ACh pulses in the absence or the presence of increasing concentrations of EPEX. Figure 6A shows two fluorescence records indicating [Ca^{2+}]_c elevations elicited by ACh (bottom black horizontal bar), before (left) and during the application of 0.1 ng/mL EPEX, as indicated by the bottom horizontal grey bar. EPEX slightly augmented the rate and amplitude of the ACh-evoked [Ca^{2+}]_c elevation. Pooled results are given in Figure 6A, showing that at 0.1 and 0.3 ng/mL, EPEX augmented the [Ca^{2+}]_c elevation by 7 and 12%, respectively (not significant). At greater concentrations, the extract did not change the ACh-evoked [Ca^{2+}]_c elevation.

Discussion

In this study, we found that an extract of the plant Eugenia punicifolia (EPEX) caused an augmentation of catecholamine release from bovine adrenal medullary cells challenged with ACh (Figure 1) or high K+ (Figure 4). Both type of stimuli are widely used to trigger secretion (both induce cell membrane depolarization and subsequently Ca^{2+} entry through voltage-gated Ca^{2+} channels, but they differ in the fact that ACh-induced catecholamine secretion implies the activation of nAChR and indirectly depolarize cell membrane, while high K+ serves to directly depolarise cell membrane (Garcia et al., 2006). We also used electrophysiological and Ca^{2+} imaging techniques to determine the mechanisms underlying this potentiation.

Augmentation of ACh-evoked secretion could be due to a direct action of EPEX on nAChR by behaving as an allosteric modulator of α7 nAChR (Faghih et al., 2007) that have been identified in bovine chromaffin cells (Quik et al., 1987; Geertsen et al., 1988; Criado et al., 2007) that have been identified in bovine chromaffin cells. EPEX could alter this ACh-elicited Ca^{2+} signal to explain its ability to augment depolarization-evoked secretion.

Although we have not yet uncovered the mechanism of action of EPEX, this study has served to unravel an entirely new biological activity for this plant extract, augmentation of exocytosis. In Brazil, Eugenia punicifolia extracts are employed in Amazon region to treat diarrhea and stomach disturbances, and as hypoglycemic medicine (Brito et al., 2007; Bopp et al., 2009). The mechanism underlying pharmacological properties of the genus Eugenia are partially attributed to flavonoids (myricitrin, querectin, and quercetrin), steroids, terpenoids, tanines, and anthraquinones (Consolini & Sarubbo, 2002). We keep doing experiments on the dichloromethane extract used in the presents tudy to gain further information on the composition and to identify the component/s that could be responsible of the observed effects on the cholinergic neurotransmission.

Its use for the treatment of diabetes could be linked to the augmentation of the exocytotic release of insulin in response to hyperglycemia. Also, it is interesting that hypoglycaemic drugs block a type of ATP-dependent potassium channel, and we cannot rule out that a ligand present in the Eugenia punicifolia extract may bind to these potassium channels to elicit an effect. Unfortunately, this type of channel is not present in bovine chromaffin cells and hence we should not explore such possibility. Related to the possibility that EPEX could be acting on other K+ channel types to regulate the catecholamine secretory response, our laboratory found that SK (small conductance calcium-activated K+ channels) and BK (big conductance calcium-activated K+ channels) were involved in the modulation of exocytosis, due to an indirect effect, by Ca^{2+} entering through VDCCs (Uceda et al., 1994; Lara et al., 1995). However, we did not find any effect of EPEX on such currents (not shown).

We have discovered a simple and efficient biological test for isolating and chemically characterising the active compound responsible for the potentiation of exocytosis by EPEX. Given the high sensitivity of this test to subnanogram concentrations of EPEX and, since
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Submilligram concentrations are sufficient to analyse exocytotic release of catecholamines from bovine chromaffin cells, testing different fractions of purified extracts can be performed.

In addition to the identification of the active compound in EPEX, our findings have also interested us to search for new biological targets and ligands with potential therapeutic interest in neurodegenerative diseases, particularly in Alzheimer’s disease (Rafii & Aisen, 2009). A compound that augments exocytosis might also enhance neurotransmission, synaptic plasticity and neuronal survival. If the molecular structure of the active compound present in EPEX could eventually be identified, it could become a prototype for the design and synthesis of new compounds capable of improving synaptic plasticity and neuronal communication.

In conclusion, we have discovered that an *Eugenia punicifolia* extract augments the exocytotic catecholamine release from bovine adrenal chromaffin cells stimulated with ACh or K⁺. Using catecholamine release as a biological probe in subsequent studies, could facilitate the isolation of the active compound and the elucidation of its molecular structure. This could be used as a template to synthesise new derivatives with therapeutic potential as cognition enhancers in Alzheimer’s disease patients.

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*Correspondence
Luis Gandia
Instituto Teófilo Hernando de I+D del Medicamento, Facultad de Medicina, Universidad Autónoma de Madrid
Arzobispo Morcillo, 4; 28029 Madrid, Spain
luis.gandia@uam.es
Tel.: 34 91 497 53 96
Fax: 34 91 497 36 80