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Nature of The Neurotoxic Membrane Actions of Amyloid-B On Hippocampal Neurons in Alzheimer's Disease

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Abstract
The mechanism by which amyloid-β (Aβ) produces brain dysfunction in patients with Alzheimer’s disease is largely unknown. According to previous studies, Aβ might share perforating properties with gramicidin, a well-accepted membrane-disrupting peptide. Therefore, we hypothesize that the key steps leading to synaptotoxicity by Aβ and gramicidin involve peptide aggregation, pore formation, and calcium dysregulation. Here, we show that Aβ and gramicidin form aggregates enriched in β-sheet structures using electron microscopy, and Thioflavin and Congo Red staining techniques. Also, we found that Aβ and gramicidin display fairly similar actions in hippocampal cell membranes, i.e. inducing Ca\(^{2+}\) entry and synaptotoxicity characterized by the loss of synaptic proteins and a decrease in neuronal viability. These effects were not observed in a Ca\(^{2+}\)-free solution, indicating that both Aβ and gramicidin induce neurotoxicity by a Ca\(^{2+}\)-dependent mechanism. Using combined perforated patch clamp and imaging recordings, we found that only Aβ produced a perforation that progressed from a small (Cl\(^{-}\)-selective pore) to a larger perforation that allowed the entry of fluorescent molecules. Therefore, based on these results, we propose that the perforation at the plasma membrane by Aβ is a dynamic process that is critical in producing neurotoxicity similar to that found in the brains of AD patients.

Keywords
Aβ, Alzheimer’s disease, Pore formation, Membrane disruption, Aβ toxicity

1. Introduction
Alzheimer’s disease (AD) is associated with amyloid-β (Aβ) peptide aggregate accumulation in the brain parenchyma (Masters et al., 1985). There is strong evidence indicating that Aβ is able to disrupt membranes, causing pore formation (Arispe et al., 1993b). Therefore, Aβ toxicity can be explained in part on the basis of the ion dyshomeostasis induced by Aβ, either by its interaction with ligand gated ion channels (NMDA, Ca\(^{2+}\) channels) or by direct lipid disruption (Arispe et al., 1993b, Diaz et al., 2009, Sepulveda et al., 2010). This hypothesis was based on microscopic conductance changes induced by Aβ pore formation in artificial membranes that were highly complex and showed cation selectivity (Arispe et al., 1993a, Kawahara et al., 1997). Single channel measurements in lipid membranes revealed a large variability of single channel conductances (50–500 pS) (Kawahara et al., 1997). Furthermore, it was found that histidine-coordinating molecules that were able to interact with Aβ blocked the ion current flow (Diaz et al., 2009, Sepulveda et al., 2010), suggesting that Aβ amyloid pores can be pharmacologically targeted. Altogether, the data suggest that Aβ does not form a single type of conventional ion channel, but instead contributes to the formation of an array of conducting pores (Kourie et al., 2001). Interestingly, using an “oligomer-enriched” form of Aβ, an increase in lipid bilayer conductance was shown (Kayed et al., 2004). However, the latter effect might be related to the solvent action rather than to a specific effect of Aβ (Capone et al., 2009).

With the aim of characterizing functional and structural features that might explain Aβ neurotoxicity, we compared the properties of Aβ and gramicidin, a classical pore-forming peptide. Gramicidin is believed to slowly insert itself into the membrane, allowing the selective passage of cations (Andersen, 1983); and, as an antimicrobial agent, it damages the cell membrane, causing a potent cytotoxic action. Here, we show that both Aβ peptide and gramicidin form β-sheet structures. Moreover, we found that Aβ and gramicidin aggregates can perforate neuronal membranes and induce neurotoxicity by a similar calcium-dependent mechanism. Unlike Aβ,
gramicidin perforations allowed a reduced entry of fluorescent molecules, suggesting that gramicidin perforates have a smaller inner diameter than Aβ perforates.

2. Methods

2.1. Ethics statement
All animals used in this study were handled in accordance with the Institutional Animal Welfare Assurance (no. 2008100A) following Comision Nacional de Investigacion Cientifica y Tecnologica de Chile (CONICYT) and National Institutes of Health recommendations.

2.2. Cell cultures
Hippocampal neurons were obtained from 18-day Sprague Dawley rat embryos, and human embryonic kidney 293 cells (HEK) were cultured as previously described (Sepulveda et al., 2009). Briefly, the neuronal feeding medium consisted of 90% minimal essential medium (MEM; BRL Technologies, Rockville, MD), 5% heat-inactivated equine serum, 5% fetal bovine serum, and a mixture of nutrient supplements. HEK cells were cultivated in Dulbecco’s modified Eagle’s medium (D-MEM; Life Technologies, Carlsbad, CA) supplemented with 10% fetal bovine serum (Life Technologies) and streptomycin–penicillin (200 units each, Life Technologies). Cells were maintained with 5% CO₂ at 37 °C.

2.3. Aβ and gramicidin aggregation
Aβ₁₋₄₀ (r-Peptide, Bogart, GA), Aβ₁₋₄₂ (Anaspec, Fremont, CA) and gramicidin (G5002, Sigma Aldrich, St. Louis, MO) were prepared according to previous protocols (Akaike, 1996, Sepulveda et al., 2009). Briefly, Aβ₁₋₄₀ and Aβ₁₋₄₂ were dissolved in DMSO (10 mg/mL) and stored in aliquots at −20 °C. For the preparation of Aβ aggregates (80 μmol/L), aliquots of peptide stock (250 μg in 25 μL of DMSO) were added to 700 μL of PBS (Gibco, Grand Island, NY), continuously agitated (200 RPM at 37 °C) for 90 minutes, and stored at 4 °C. Gramicidin was dissolved in DMSO (50 mg/mL) and the aggregates were formed after adding PBS to obtain the desired concentration.

2.4. Electrophysiology recordings
Patch recordings were performed as previously described (Sepulveda et al., 2009). In short, Aβ and Gramicidin aggregates were used at 500 nmol/L and 100 μg/mL, respectively. Perforated recordings were obtained as follows: the perforating agent was added into the pipette solution and a 5-mV pulse was used to monitor the formation of the perforation at a holding potential of ~60 mV using an Axopatch 200B amplifier (Molecular Devices, Sunnyvale, CA). Data were displayed and stored using a 1322A Digidata acquisition board and analyzed with electrophysiological software (Molecular Devices). The external solution contained the following (in mmol/L): 150 NaCl, 5.4 KCl, 2.0 CaCl₂, 1.0 MgCl₂, 10 glucose, and 10 HEPES (pH 7.4, 330 mOsm). The standard internal solution in the patch pipette contained the following (in mmol/L): 120 KCl, 4.0 MgCl₂, 10 BAPTA, and 2.0 Na₂-ATP (pH 7.4, 310 mOsmol). Short applications of GABA (100 μmol/L) were done via lateral motion of a multi-pipette array (~200 μm in diameter). Some experiments involved an external solution without added calcium, NA7 (20 μmol/L), NA4a (20 μmol/L), or the inactive peptides NA13 and NA15 (20 μmol/L). These peptides were purchased from Peptide 2.0 (Chantilly, VA).

2.5. Western blots
The levels of proteins were detected by using the following antibodies: anti-Synapsin I (1:1000; AB1543, Chemicon, Billerica, MA) and anti-SV2 (1:200; Developmental Studies Hybridoma Bank, Iowa City, IA). Immunoreactive bands were visualized with ECL plus Western Blotting Detection System (PerkinElmer, Waltham, MA).
2.6. Transmission electron microscopy
Samples of aggregated Aβ (2.2 μg/mL) and gramicidin (100 μg/mL) in phosphate-buffered saline solution (PBS) were placed on nickel grids (Sigma Aldrich, St. Louis, MO) pretreated with 200-mesh Formvar and shaded with carbon. Phosphotungstic acid (0.2%) was used for contrast and samples were examined under a transmission electron microscope (TEM; JEOL-JEM-1200 EX II, Peabody, MA).

2.7. Thioflavin and Congo Red staining
Thioflavin T and Congo Red (Sigma Aldrich, St. Louis, MO) staining were performed as previously described (Klunk et al., 1989, LeVine, 1993). Briefly, the stainings were performed by incubating the suspension of Aβ (2.2 μg/mL) and gramicidin (100 μg/mL) with Congo Red (10 μmol/L), ThS (200 μmol/L), and ThT (200 μmol/L) in PBS with continuous agitation (200 rpm at 37 °C) for 30 minutes. The samples were mounted on a coverslip and observed under a fluorescent microscope (Eclipse TE2000; Nikon, Melville, NY, USA) equipped with a CCD camera (Andor Technology, Belfast, UK) and Workbench software 6.0 (Axon Instruments, Molecular Devices, Sunnyvale, CA). The images were processed using ImageJ software (National Institutes of Health, Bethesda, MD).

2.8. Intracellular calcium imaging
Neurons were loaded with Fluo-4 AM (1 μmol/L in pluronic acid/DMSO, Molecular Probes, Eugene, OR) for 30 minutes at 37 °C. The cells were illuminated (200 milliseconds) under an inverted fluorescent microscope (Eclipse TE; Nikon, Melville, NY, USA), using a computer-controlled Lambda 10-2 filter wheel (Sutter Instruments, Novato, CA). Fluorescent signal from regions of interest (ROI) on different neurons was collected at 1- to 2-second intervals during a continuous 5-minute period, using a SensiCam camera (PCO, Kelheim, Germany) associated to Workbench 6 software.

2.9. Immunocytochemistry
Hippocampal neurons were fixed for 15 minutes with 4% paraformaldehyde and permeabilized with 0.1% triton X-100 in PBS. Monoclonal anti-SV2 (1:200; Developmental Studies Hybridoma Bank, Iowa City, IA) and rabbit anti-MAP2 (1:300; Santa Cruz Biotechnology, Dallas, TX CA) were incubated overnight, followed by incubation with a secondary anti-rabbit IgG (1:500; Jackson ImmunoResearch Laboratories, West Grove, PA) conjugated with Cy3 and anti-mouse IgG conjugated with FITC (1:500; Jackson ImmunoResearch Laboratories) for 2 hours.

2.10. Transfection
For the NMDAR experiments, human embryonic kidney 293 (HEK293) cells were transfected with NR1–1a, NR2A, and green fluorescent protein (pGreenLantern plasmid, Invitrogen, Life Technologies, Carlsbad, CA) at a ratio of 2:2:1 using the calcium phosphate transfection kit (Invitrogen, Life Technologies). Expression of green fluorescent protein was used as a marker for positively transfected HEK293 cells and recordings were made after 16 to 20 hours.

2.11. Neuronal viability
3-(4,5-Dimethylthiazol-2-yl)-2, 5-diphenyltetrazolium bromide (MTT; Sigma-Aldrich, St. Louis, MO) was added to each well, and the plates were incubated for 6 hours. The Formazan crystals were then dissolved in dimethyl sulfoxide (DMSO; Sigma-Aldrich), giving a spectrophotometrically measurable purple solution. Optical density was read at 570 nm on the NOVOstar plate reader (BMG Labtech, Ortenberg, Germany).

2.12. Microscopic current recordings and analysis
The methodology for single-channel recordings in cell-attached configuration was done using patch pipettes coated with R6101 elastomer (Dow-Corning, Midland, MI) that had a tip resistance of 6 to 10 megaohms after fire polishing. Cells were voltage clamped at −50 mV, and the current was filtered (1-kHz low-pass 8-pole Butterworth) and acquired at 5 to 20 kHz using pClamp software (Molecular Devices). NMDA receptor agonist
and blockers were applied to cells using a stepper motor-driven rapid solution exchanger (Warner Instrument Corp, Hamden, CT). Cells were maintained in extracellular medium containing the following (in mmol/L): 150 NaCl, 5 KCl, 2 CaCl₂, 10 HEPES, 10 glucose; pH 7.4. The intracellular recording solution contained (in mmol/L): 120 KCl, 2 MgATP, 10 BAPTA, and 10 HEPES, pH 7.2.

2.13. Simultaneous cell imaging and electrophysiological recordings
Hippocampal neurons were mounted on an inverted microscope (TE200U, Nikon, Melville, NY, USA) equipped with EMCCD IonX 16-bit camera (Andor, Belfast, UK) plus a patch clamp set-up. The imaging was carried out using a 40X/1.35 objective, and the fluorescent signal for ethidium bromide and a non-hydrolyzable glucose analog (6-NBDG) were obtained using excitation wavelengths of 480 nm and 545 nm, respectively. The emissions of these fluorescent compounds were collected at 520 nm and 590 nm, respectively. To diminish the vibration of the stage caused by the movement of the excitation source, a shutter and filter wheel were used to automatically shift the excitation wavelengths. The xenon lamp housing and the filter wheel assembly were coupled to the microscope with a bent optical fiber.

2.14. Data analysis
Nonlinear analysis was performed using OriginPro 7.0 (Microcal, Origin Lab, Northhampton, MA). Membrane charge was calculated by integrating the transient capacitative current after subtracting the pipette capacitance; under this condition, the area under the current represented the membrane charge transferred (fC). Moreover, we fitted the current with a standard exponential function using Chebyshev algorithm to calculate the tau for the decay of the current. Using this strategy, we calculated the total charge transferred to the membrane under the patch pipette. All-points amplitude histograms of single channel experiments were generated and fitted with Gaussian functions using ClampFit. The values were expressed as mean ± standard error of the mean (SEM) from at least 5 to 8 neurons or cells. Statistical differences were determined using 1-way analysis of variance (ANOVA) or paired Student t tests, followed by the Bonferroni post hoc test in some cases.

3. Results
3.1. Aβ and gramicidin aggregate in aqueous solution
We previously hypothesized that a key step in inducing membrane perforation and synaptotoxicity by Aβ and gramicidin might involve protein aggregation and a subsequent calcium dysregulation (Aguayo et al., 2009). Therefore, we studied the ultrastructural characteristics of Aβ and gramicidin in a saline aqueous solution, and found that Aβ and gramicidin formed fibrillar aggregates (Fig. 1A). To gain further insight into the common features of these structures, we stained them with Thioflavin T and S, two compounds that react with proteins enriched in β-sheet structures (LeVine, 1993). Both peptides were positive for these amyloid-staining compounds (Fig. 1B). In addition, both were congophilic and displayed green birefringence under polarized light, which is characteristic for amyloid (Fig. 1C). The presence of β-sheet structures has been well described for Aβ aggregates (Klunk et al., 1989, LeVine, 1993); however, this is the first time that this structural feature has been associated with gramicidin aggregates.
Fig. 1. Aβ and gramicidin form amyloid aggregates. (A) Ultrastructural microphotographs of Aβ (2.2 μg/mL) and gramicidin (100 μg/mL) aggregates in phosphate-buffered saline solution (PBS). Aβ was agitated at 200 RPM at 37 °C for 90 minutes, and gramicidin was added directly to PBS before observation. (B) Microphotographs show ThS-positive (left panel) and ThT-positive (right panel) Aβ and gramicidin aggregates. The aggregates were obtained as described in A and stained with ThS (200 μmol/L) and ThT (200 μmol/L). (C) Aβ and gramicidin aggregates (obtained under the same conditions as described in A) stained with Congo Red and photographed under polarized light, demonstrating the congophilic-positive nature of both amyloid aggregates. Bars correspond to 500 nm (A), 10 μm (B), 5 μm (C, top panels), and 2 μm (C, bottom panels).

3.2. Aβ and gramicidin altered membrane conductance
To better understand the phenomenon of membrane perforation induced by Aβ, we compared the properties of Aβ and gramicidin on neuronal and HEK293 membranes. Aβ and gramicidin aggregates displayed rapid perforating actions (Fig. 2A). In hippocampal neurons, for example, Aβ and gramicidin increased the capacitative current during cell attached recordings (175 ± 25 and 185 ± 28 fC, n = 5–7 cells). In comparison, the recordings were stable with a control solution in the patch pipette (i.e., 30 minutes) (Fig. 2B). The analysis of charge transferred during the capacitative response and membrane resistance showed that the effect of Aβ and gramicidin on hippocampal neuron and HEK membranes was very similar (Fig. 2C and D). On the other hand, this membrane effect was not produced by the reverse Aβ peptide (Aβ40*-), a non-aggregating peptide, supporting the idea that Aβ aggregation leads to membrane pore formation, which we previously demonstrated to play a key role in synaptotoxicity induced by Aβ (Parodi et al., 2010, Peters et al., 2013, Sepulveda et al., 2010). In addition, the data show that the effects of Aβ on membrane conductance were also blocked by NA7 (5 ± 5 fC, n = 5), a mini-peptide that interferes with Aβ perforations, but not by NA13 (135 ± 12 fC; Fig. 2C and D). The lack of activity found with NA13 is due to the substitution of key His residues by Ala (Diaz et al., 2009, Parodi et al., 2010, Sepulveda et al., 2010).
3.3. Aβ and gramicidin aggregates induce neurotoxicity by a calcium-dependent mechanism

We previously showed that Aβ was able to decrease the level of a pre-synaptic protein by a Ca\(^{2+}\)-dependent mechanism (Parodi et al., 2010, Sepulveda et al., 2010). In the present study, we found that Aβ and gramicidin significantly decreased the levels of SV2 (Fig. 3A, green signal), as shown by a diminished number of puncta (18 ± 1 vs. 10 ± 2 and 9 ± 1 puncta/20 μm) and a decrease in the fluorescence associated with SV2 in hippocampal neurons (Fig. 3B and C). In addition, this synaptotoxic effect of Aβ and gramicidin in hippocampal neurons was confirmed by Western blot analysis (Fig. 3D and E). Interestingly, using fluorescence measurements with Fluo-4, we found that both Aβ and gramicidin increased intracellular calcium in hippocampal neurons (Fig. 4A and B) at times when they were disrupting the cell membrane (i.e., 15 to 20 minutes) (Fig. 2A). Aβ and gramicidin reduced neuronal viability, measured with an MTT assay (78% ± 3% and 74% ± 6% of control at 24-hour treatments), and these effects were blocked by a reduction in external calcium from 2 to 0.01 mmol/L (Fig. 4 C and D) supporting a calcium-dependent mechanism. The data also show that culturing hippocampal cells in the absence of external calcium for 24 hours was without effects on viability. Similar results were obtained by adding BAPTA-AM, a calcium chelator (data not shown), supporting the idea that the neurotoxicity induced by Aβ and gramicidin is through a calcium-dependent mechanism.
3.4. Changes in microscopic membrane conductance induced by Aβ and gramicidin

The effect of a pore-forming peptide to produce the increase in capacitative current results from the sum of small changes in membrane conductances (Ebihara et al., 1995). Therefore, we wanted to detect the channel-like microscopic current induced by Aβ and gramicidin during cell-attached recordings to determine the degree of similarities. These experiments were done in HEK cells, because neurons present too many ion channels.

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Fig. 3. Aβ and gramicidin induce synaptoxicity. (A) Confocal microphotographs obtained in control and after culturing hippocampal neurons with Aβ (500 nmol/L) and gramicidin (50 nmol/L) for 48 hours. Green signal shows SV2 immunoreactivity, and red signal is MAP2, a neuronal marker. Bar scales correspond to 10 μm (A, left panels) and 2 μm (A, right panels). (B and C) Graphs illustrate the number of fluorescent SV2 puncta/20-μm length and intensity of the fluorescent signal associated with SV2, respectively, under the same conditions. (D) Western blots show the inhibitory effects of Aβ and gramicidin on SV2 levels measured by chemoluminescence techniques (α-tubulin and β-actin were used as internal controls). (E) Graphs summarize SV2 levels under the conditions described in (D) (from 3 different experiments). * p < 0.05; ** p < 0.01.

Fig. 4. Toxic effects of Aβ and gramicidin on hippocampal neurons were calcium dependent. (A) Graph shows the effect of exposing hippocampal neurons to 1 μmol/L Aβ (filled symbols) and 1 μmol/L gramicidin (open symbols) on intracellular Ca^{2+}. Arrow indicates the beginning of the application. (B) Effect of Aβ and gramicidin on intracellular Ca^{2+} levels in hippocampal neurons measured after 20 minutes of application. (C and D) Aβ and gramicidin neurotoxicity is calcium dependent, as measured by hippocampal neuron viability using an MTT assay after 24 hours of treatment. Data also show that removing extracellular calcium was without effect on viability. Bars are mean ± standard error. n ≥ 3 cells. * p < 0.05; *** p < 0.01.
precluding this objective. Using normal internal solution (140 mmol/L CsCl), these cells did not display any events reminiscent of single channel currents (not shown). On the other hand, in cells recorded with Aβ (n = 32) or gramicidin (n = 13), the recording progressed to a perforated configuration, without detectable single channel behavior. Moreover, in some cells, Aβ and gramicidin induced conductance changes that were not like typical single channel behavior (Sepulveda et al., 2010). Fig. 5A, for example, shows representative traces obtained in the presence of Aβ (left) and gramicidin (right) during establishment of perforated conformations. The noisy nature of the microscopic current events produced by both peptides did not allow clear discriminations between open and closed states (Fig. 5B) and plots of all-point current distributions from different patches (Aβ, n = 7; gramicidin, n = 8) showed multiple levels of peak conductance (100–600 pS), suggesting the occurrence of several membrane disruptions by Aβ and gramicidin (Fig. 5C). Furthermore, it was noteworthy that gramicidin was unable to induce the single channel behavior described in artificial lipid membranes (Andersen, 1983).
Under the same conditions, NR1 and NR2A NMDA-induced channels (45 pS) were clearly defined (Fig. 5B, inset).

3.5. Reversal potential of GABA<sub>A</sub>R Cl<sup>−</sup>-current recorded with Aβ and gramicidin cell perforations indicate differential ionic selectivity

Previous studies showed that the ionophore action of Aβ during the first 15 minutes of pore formation was cation selective, as it behaved similarly to gramicidin, a recognized cation-selective pore-forming peptide (Sepulveda et al., 2010). In the present study, we compared GABA current–voltage (I–V) relationships produced by Aβ and gramicidin with recordings at early (15–20 minutes) and late (30–40 minutes) states of membrane perforation (Fig. 6A). The data show that Aβ (squares), but not gramicidin (circles) produced a membrane perforation that progressed from cation-selective–like to non-selective. At 15 to 20 minutes (filled symbols), for example, the GABA<sub>A</sub>-activated current reversed at a potential near the Cl<sup>−</sup> reversal potential (~40 mV) with either Aβ or gramicidin in the pipette, supporting the idea that the perforations allow the passage of cations with short periods of applications. Interestingly, at 30 to 40 minutes of application with Aβ (open squares; see arrows), but not gramicidin (open circles), the Cl<sup>−</sup> current reversed near 0 mV, which is close to the expected value for a non-selective pore, supporting the idea that Aβ, but not gramicidin, is able to produce a membrane perforation that changes its selectivity with time. The current traces show that Aβ, but not gramicidin, caused a shift of the reversal potential for the Cl<sup>−</sup>-induced GABA current to 0 mV (Fig. 6B).
Fig. 6. Aβ pores have a time-dependent transition from cationic-selective to non-selective. (A) Graph shows GABA-induced anionic current–voltage relationships obtained with perforated recordings at 15 and 30 minutes for Aβ (squares) and gramicidin (circles). Symbols represent the mean ± standard error of the mean from 3 or more neurons. Reversal potentials were close to −50 and 0 mV at 15 to 20 minutes (filled symbols) and 30 to 40 minutes (open symbols) of recording. Note the change on reversal potential for Aβ (red arrows), but not gramicidin, at 30 minutes. (B) Traces show currents activated by 100 μmol/L GABA at two times of perforated recordings and holding potentials of 0 and −60 mV. Trace recorded at 30 to 40 minutes of Aβ-induced perforated recording reverses at 0 mV, indicating that the Cl− gradient was lost.

3.6. Aβ perforations permitted entry of small molecules into hippocampal neurons

Using a technique that combined perforated patch clamp recordings and cell imaging with a fluorescent non-hydrolyzable glucose analog (6-NBDG, ~1 nm) in the patch pipette, we found that small fluorescent probes that have molecular weight and sizes comparable to those of other important cellular metabolites, such as glucose (~0.9 nm) and nucleotides like ATP (~1.5 nm), did not pass through intact hippocampal neuron membranes. The control data recorded in the absence of Aβ show that 6-NBDG in the pipette was unable to get into hippocampal neurons (Fig. 7A, left). However, disruption of the integrity of the membrane under the patch pipette when the whole-cell configuration was established with a small negative pressure produced the rapid entry of 6-NBDG into the neuron detected within 30 sec (Fig. 7A, right). Thus, the presence of intracellular fluorescence was used as an indicator for membrane leakiness induced by Aβ or gramicidin using 2 small fluorescent molecules with a Van der Waals diameter near ~1 nm (6-NBDG, em/ex = 490/520) and ~1.3 nm (EtBr, em/ex = 545/565). The data show that Aβ-induced perforations allowed the entry of 6-NBDG and EtBr into hippocampal neurons as assessed by the fluorescence increase (Fig. 7C and D), in parallel with the increase in the capacitative current (Fig. 7E and F). The effect of Aβ was dependent on the functionality of the Aβ pores because the fluorescence increase was blocked by NA7 or NA4a, peptides designed to block changes in membrane conductance associated with amyloid perforations (Fig. 7C and D, middle). On the other hand, although gramicidin increased the charge transfer, the pore formed did not produce the entry of EtBr into the neurons (red squares). The data also show that gramicidin caused a small increase in the fluorescent signal (46 ± 6 URF) associated with 6-NBDG compared with the effect induced by Aβ. In support of the idea that Aβ membrane perforation is a time-dependent process, progressing from a small (ion-selective pore) to a large membrane perforation, we found that the time course for the capacitative response, mainly associated with the passage of ions through the
membrane, increased significantly before uptake of EtBr and 6-NBDG through Aβ perforations (Fig. 7E and F), suggesting a late formation of a membrane perforation with a larger pore size than that of gramicidin (Fig. 8).

Fig. 7. Aβ perforations allow the entry of small molecules into hippocampal neurons. (A) Left: Cell-attached recording in control conditions using a patch pipette filled with 6-NBDG. Right: Whole-cell configuration allowed rapid entry of 6-NBDG into the neuron. (B) Left: EtBr (diameter ~1.3 nm) and 6-NBDG (diameter ~1 nm) 2-dimensional structures. Right: Schematic representation of the electrical and fluorescence changes expected under cell-attached and whole-cell configurations. (C) Time-dependent increase in intracellular EtBr fluorescence in the presence of Aβ in the pipette (upper panel) was blocked by the NA7 mini-peptide (middle panel). Gramicidin did not produce the entry of EtBr (lower panel). (D) Time-dependent intracellular increase of 6-NBDG in the presence of Aβ in the pipette (upper panel) was blocked by NA7 mini-peptide (middle panel). Gramicidin allowed a discrete entry of 6-NBDG into hippocampal neurons (lower panel). (E) Graphs show the quantification of membrane charge transferred (capacitative current) and the EtBr fluorescence induced by Aβ (left) and gramicidin (right) perforations. Aβ, but not gramicidin, perforations allowed the entry of EtBr. (F) Graphs show the quantification of membrane charge transferred (capacitative current) and the 6-NBDG fluorescence entry induced by Aβ (left) and gramicidin (right) perforations. * p < 0.05.

Fig. 8. Simplified model showing the micro- and macro-perforations induced by Aβ. After association with membrane components, the oligomers/aggregates cause disruptions characterized by small ionic pores, allowing cationic influx. Longer exposure of Aβ in the membrane increases the perforation, allowing the entry or exit of large molecules such as EtBr and 6-NBDG. The resulting cellular dyshomeostasis subsequently leads to synaptic failure and apoptosis.
4. Discussion

4.1. Lack of single channel behavior of Aβ and gramicidin amyloid-like aggregates in native membranes

It was previously found that gramicidin and Aβ formed pores in artificial lipid membranes having complex single channel conductance (100–1000 pS) and kinetic properties, supporting the idea of complicated molecular behavior (Arispe et al., 1993a, Ebihara et al., 1995). In addition, the channel time course was well described by clear open and closing kinetics for gramicidin (Andersen, 1983) and for Aβ (Kourie, 2001). However, unlike artificial membranes (Andersen, 1983, Arispe et al., 1993a), we found no channel behavior with either ionophore in native membranes. In a few recordings, the response more reminiscent of single channel behavior was an increase in “noise” that we interpreted as a restricted and temporal membrane disruption. The lack of single channel behavior for gramicidin was somewhat surprising, because based on single channel studies in lipid membranes, it was proposed that 2 monomers of gramicidin inserted into the membrane form a functional cationic-selective channel (Kelkar and Chattopadhyay, 2007). In their working model, it was assumed that gramicidin existed only as a monomer in aqueous solution. However, under our working saline conditions, it was evident that gramicidin formed complex and stable oligomeric structures to produce membrane perforations supporting the idea of complex effects on cellular membranes. These structures were of amyloid nature because they reacted with thioflavin (T and S) and Congo Red, known to be β-sheet binding compounds (Klunk et al., 1989, LeVine, 1993). Thus, the present study supports the conclusion that both neurotoxic aggregates, which can form membrane pores, present this type of secondary structure in aqueous solution. In addition, the gramicidin aggregates were more stable than Aβ aggregates, because the former aggregates were proteinase K resistant (data not show).

4.2. Considerations for Aβ perforations and its toxic effects

From the perforated patch clamp recordings obtained at different times, we postulate that Aβ and gramicidin actions on cell membranes are fairly different. The gramicidin action appeared to be stable and selective for cations, because the Cl− gradient was not affected, providing a reversal potential near ~40 mV for the GABA_A receptor–mediated current. For Aβ, on the other hand, using the GABA_A–mediated reversal potential as a readout, we detected a transition state of Aβ pores from cationic-selective pores to non-selective pores, suggesting that the Cl− gradient was lost during prolonged interaction of the peptide with the membrane. The data obtained with single cell imaging analysis indicate that Aβ caused the entry of EtBr and 6-NBDG into the cell, providing additional support for this lack of ionic selectivity during the latter stage of membrane perforation (Fig. 7). For gramicidin pores, on the other hand, a modest entry of small fluorescent molecules (6-NBDG), but not for large fluorescent molecules (EtBr), into the cell suggests pore sizes smaller than with Aβ. These data imply that, upon binding to the cell membrane, Aβ may initiate its own nucleation, producing a large perforation allowing the entry of the larger molecules (Fig. 8). Interestingly, the entry of EtBr started minutes after the formation of the pore, indicating that these molecules were taken up by these Aβ structures. In fact, an inhibitory peptide (NA7) that blocked these pores was also able to block the EtBr uptake, indicating that the Aβ pores were allowing the passage of small molecules. The mechanism for this disrupting effect of Aβ on neuronal membranes is largely unknown, but it is possible that the peptide interacts directly with the lipid components or by means of other neuronal proteins such as NMDA receptors (Ronicke et al., 2011, Shankar et al., 2007) or Ca2+ channels (Mezler et al., 2012).

4.3. Neurotoxicity induced by Aβ and gramicidin aggregates in hippocampal neurons

We have recently shown that the synaptotoxicity produced by Aβ is a consequence of a membrane perforation that leads to a vesicular depletion caused by a calcium-dependent mechanism (Parodi et al., 2010, Sepulveda et al., 2010). In the present study, we found that Aβ aggregates can form membrane perforations, leading to a
decrease in hippocampal neuron viability, which was blocked by reducing extracellular calcium. It is noteworthy that both aggregates decreased the presynaptic protein SV2, indicating that Aβ and gramicidin aggregates produced a synaptic failure, which could be part of the early changes leading to AD (Aguayo et al., 2009). Gramicidin has been widely described as capable of producing extensive bacterial death. However, it has not been shown that gramicidin can produce neurotoxicity. Interestingly, it has recently been suggested that Aβ actually possesses antimicrobial activity, and may function as an effector molecule of innate immunity in the brain (Soscia et al., 2010).

In conclusion, we demonstrate that the membrane effects of Aβ have several similarities to those of gramicidin, explaining its calcium-dependent induced synaptotoxicity. We postulate that the inhibition of these effects should render Aβ inactive.

Disclosure statement
The authors declare that they have no conflicts of interest.

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