Cardiolipin Profile Changes are Associated to the Early Synaptic Mitochondrial Dysfunction in Alzheimer’s Disease

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Abstract. Brain mitochondria are fundamental to maintaining healthy functional brains, and their dysfunction is involved in age-related neurodegenerative disorders such as Alzheimer’s disease (AD). In this study, we conducted a research on how both non-synaptic and synaptic mitochondrial functions are compromised at an early stage of AD-like pathologies and their correlation with putative changes on membranes lipid profile, using 3 month-old nontransgenic and 3xTg-AD mice, a murine model of experimental AD. Bioenergetic dysfunction in 3xTg-AD brains is evidenced by a decrease of brain ATP levels resulting, essentially, from synaptic mitochondria functionality disruption as indicated by declined respiratory control ratio associated with a 50% decreased complex I activity. Lipidomics studies revealed that synaptic bioenergetic deficit of 3xTg-AD brains is accompanied by alterations in the phospholipid composition of synaptic mitochondrial membranes, detected either in phospholipid class distribution or in the phospholipids molecular profile. Globally, diacyl- and lyso-phosphatidylcholine lipids increase while ethanolamine plasmalogens and cardiolipins content drops in relation to nontransgenic background. However, the main lipidomic mark of 3xTg-AD brains is that cardiolipin cluster-organized profile is lost in synaptic mitochondria due to a decline of the most representative molecular species. In contrast to synaptic mitochondria, results support the idea that non-synaptic mitochondria function is preserved at the age of 3 months. Although the genetically construed 3xTg-AD mouse model does not represent the most prevalent form of AD in humans, the present study provides insights into the earliest biochemical events in AD brain, connecting specific lipidomic changes with synaptic bioenergetic deficit that may contribute to the progressive synapses loss and the neurodegenerative process that characterizes AD.

Keywords: Alzheimer’s disease, brain bioenergetics, cardiolipin, mitochondrial lipidomics

INTRODUCTION

Lipids are a structurally high diverse group of amphiphilic molecules with a key biological role in energy storage and demand, in the dynamic structure of biomembranes, and in several cell signaling pathways [1, 2]. In multicellular organisms, the spatial
and temporal regulation of lipidome reaches a high level of complexity, which includes a transversal (lipid asymmetry) and a lateral (lipid domains) dynamic membrane organization [1, 2]. Therefore, the cornerstone of the highly complex eukaryotic organisms (as humans) is a continuous high energy flux required to the dynamic regulation (in time and space) of a vastly expanded repertoire of lipid species. Mitochondria, which supply more than 90% of the ATP used by animal cells, must be a key player in the homeostatic control of lipidome. Thus, it is not surprising that human diseases connected with aberrant lipid metabolism (e.g., Alzheimer’s disease, AD) are also associated with mitochondrial dysfunction [3, 4].

AD is the most common neurodegenerative disorder predominantly characterized at the brain level by extracellular deposition of amyloid-β (Aβ) peptide, intracellular neurofibrillary tangles containing hyperphosphorylated tau protein, and synaptic dysfunction [5]. Since Dr. Alois Alzheimer described this disorder over 100 years ago, a number of molecular mechanisms underlying altered biochemical pathways have been identified as potential causes of AD [6–9]. Abnormal enzymatic processing of amyloid-β protein precursor (AβPP) raised the first hypothesis for AD, the amyloid cascade hypothesis, which postulates that generation of Aβ fragments and the formation of senile plaques trigger AD pathology [9]. AβPP is a transmembrane glycoprotein implicated in neural plasticity and synapse formation. The mechanism by which AβPP is cleaved is not fully understood, but two pathways have been reported. Normally, AβPP follows the constitutive nonamyloidogenic pathway via α-secretase, releasing the neurotropic ectodomain (soluble AβPP) in the extracellular space, but it can also undergo an amyloidogenic process via BACE (β-site cleaving enzyme), leading to Aβ formation with consequent neurotoxic effects [10]. It is reported that AβPP and α-secretase are mainly located in lipid disordered domains (Ld or Lα phase) while its proteolytic enzyme (BACE) resides in lipid ordered domains (Lo phase or rafts) [11]. Thus, the regulation of lateral compartmentalization of the membrane is essential to prevent the neurodegenerative process induced by Aβ. However, the soluble Aβ levels are better correlated with dementia severity than amyloid deposits [12]. Soluble Aβ itself can interact with membrane lipids, promoting changes in the dynamic organization of cell membranes, including mitochondrial membranes [13]. In fact, profound alterations on mitochondrial enzyme activity were associated with AD and mitochondrial dysfunction was also proposed as the primary event leading to AD pathology [14, 15]. Brain mitochondrial dysfunction, characterized by reduced ATP synthesis and enhanced reactive oxygen species (ROS) generation [16, 17], may underlie oxidative stress, the loss of cellular Ca2+ homeostasis and the disruption of the functional membrane structure, triggering both Aβ and tau pathologies.

Mitochondria functional integrity is highly dependent on phospholipid composition, mainly cardiolipin (CL) content and diversity [18–20]. CL, a unique anionic tetra-acyl phospholipid class in mammalian cells, is fundamental to the functional membrane assembly of redox complexes of the respiratory system and phosphorylation apparatus, regulating mitochondrial bioenergetics [21, 22]. In this context, we aimed to evaluate the bioenergetic alterations and the mitochondrial membrane lipid profile at an early stage of AD in order to better understand how the putative mitochondrial phospholipid profile changes are reflected in brain mitochondrial bioenergetics, in brain ATP levels and brain redox state usually associated with AD pathology.

MATERIAL AND METHODS

Animal care

All animal experiments were conducted according with the National (DL 129/92, DL 197/96, P 1131/97) and European Convention for the Protection of Animals used for Experimental and Other Scientific Proposes and related European Legislation (2010/63/EU). Male nontransgenic (nonTg; C57BL6/129S) and triple transgenic (3xTg-AD) mice for AD were used, since they express three major genes associated with familial AD, namely APP, PS1M146V, and tauP301L, and develop the pathological hallmarks of disease in an age-dependent manner [23, 24]. Thus, Aβ and tau pathologies are detected at 6 and 12 months of age, respectively. Animals were obtained from Center for Neuroscience and Cell Biology (CNC) Animal Facilit, University of Coimbra and maintained at veterinary hospital of the University of Trás-os-Montes e Alto Douro and were housed in polycarbonate cages on a 12 h light/dark cycles. Room temperature was maintained at 24 ± 2°C with relative humidity 55 ± 5%. Food and water were provided ad libitum. Although the experiments performed in the present work were carried out in 3-month old presymptomatic male mice, the neuropathological hallmarks were assessed in 14-month old male nonTg and 3xTg-AD. The extracellular Aβ deposits and intracellular deposits of hyperphosphorylated tau protein were detected by
Fig. 1. Phosphorylated AMPK (pAMPK) to AMPK total ratio. Representative western blots of pAMPK and AMPK total (upper panel). Bar graphs show pAMPK/AMPK total ratio, determined by western blot analysis, in brain mitochondria-free extracts obtained from 3 months-old nonTg and 3xTg-AD mice. Error bars represent standard deviation for \( n \geq 3 \) independent experiments using 3 animals in each one. *Significantly different from nonTg group, with \( p \leq 0.05 \).

Determination of brain ATP, ADP, and AMP levels

Adenine nucleotides (ATP, ADP, and AMP) were extracted from nonTg and 3xTg-AD brains using a modified tissue extraction protocol as described elsewhere [25]. ATP, ADP, and AMP were quantified by high performance liquid chromatography (HPLC) (UltiMate 3000, Dionex) using standard curves obtained with the chromatogram peak areas corresponding to different concentrations of each adenine nucleotide. The results were expressed as \( \mu \text{mol/g wet brain} \).

Determination of enzymatic and non-enzymatic antioxidant defences in brain mitochondria-free cytosolic fraction

Brain enzymatic (superoxide dismutase, SOD; catalase, CAT) and non-enzymatic (glutathione, GSH; glutathione disulfide, GSSG) antioxidant defences as well ROS generation were measured in mitochondria-free cytosolic fraction from nonTg and 3xTg-AD mice brains. Brains were homogenized in ice-cold buffer (160 mM sucrose, 10 mM Tris-Cl, pH 7.4) (1:10, w/v) using a glass-Teflon Potter Elvins hom. Brain mitochondria-free cytosolic fractions were obtained after centrifugation at 16 000\( \times g \) for 20 min at 4\( ^\circ \)C (Sigma 2K-16). The final concentration of protein was determined by the Biuret method [26].

Copper-zinc SOD (CuZn-SOD) activity was evaluated using 0.2 mg of protein into phosphate buffer (50 mM KH\(_2\)PO\(_4\), 1 mM EDTA, pH 7.4) supplemented with 0.1 mM nitro blue tetrazolium (NBT) and 0.1 mM hypoxantine [27]. Absorbance was measured in a Cary 50 UV-Vis spectrophotometer (Agilent Technologies, Santa Clara, CA, USA) at 560 nm after 0.023 U mol\(^{-1}\) xanthine oxidase addition. The values were expressed as U/min/mg protein in which U is defined as the enzyme activity that inhibited the reduction of NBT to blue formazan by 50%.

CAT activity was determined polarographically following the oxygen production using a Clark-type oxygen electrode (Hansatech, Norfolk, UK) resulting from \( \text{H}_2\text{O}_2 \) decomposition [28]. The reaction was conducted in 1 mL of 50 mM sodium phosphate buffer (pH 7.0) and initiated by adding 1 M \( \text{H}_2\text{O}_2 \). CAT activity was expressed in terms of nmol O\(_2\) produced per minute per milligram protein.

GSH and GSSG levels were determined by spectrofluorimetric assay, using a Varian Eclipse fluorescence spectrophotometer (Agilent Technologies, Santa Clara, CA, USA) setting excitation at 350 nm and emission at 420 nm [29]. To determine GSH content, samples (0.5 mg of protein) were incubated at room temperature in 2.5 mL of phosphate buffer (100 mM potassium phosphate, 5 mM EDTA, pH 8.0) supplemented with 200 \( \mu \text{g} \) fluorescein ortho-phthalaldehyde (OPT) for 15 min. To determine GSSG content, samples (0.5 mg of protein) were first incubated at 22\( ^\circ \)C with 12 mM \( \text{N} - \text{ethylmaleimide} \) (NEM) for 45 min. After that 1.66 mL of 0.1 M NaOH and 0.7 mL of phosphate buffer supplemented with 200 \( \mu \text{g} \) OPT were added to the reaction followed by 15 min incubation at 22\( ^\circ \)C. Concentrations were calculated using standard curves prepared with different concentrations of GSH and GSSG which underwent the same sample treatment. The results were expressed as GSH/GSSG ratio.

ROS production in brain mitochondria-free cytosolic fractions was estimated using the fluorescent probe 2,7'-dichlorofluorescein diacetate (DCFH-DA) [30]. Samples (0.1 mg of protein) were incubated with DCFH-DA (5 \( \mu \text{M} \)) at 37\( ^\circ \)C for 45 min. Fluorescence intensity of 2,7'-dichlorofluorescein (DCF) was measured in a Varian Eclipse fluorescence spectrophotometer (Agilent Technologies, Santa Clara, CA, USA) at 480 nm excitation and 535 nm emission.
wavelengths. ROS content was determined using a calibration curve obtained with DCF standards. Results were expressed as nmol DCF per milligram of protein.

**Western blots analysis of 5′ adenosine monophosphate-activated protein kinase (AMPK)**

AMPK total and phosphorylated AMPK levels in nonTg and 3xTg-AD mice brains were evaluated by western blots analysis [31]. Western blots were performed in the mitochondria-free cytosolic fractions obtained as described above and supplemented with phosphatase inhibitors cocktail (Roche). The samples were resolved by electrophoresis in 10% SDS–polyacrylamide gels and transferred to polyvinylidene difluoride (PVDF) membranes. The blots were subsequently incubated with specific primary antibodies [p-AMPK alpha (T172) Antibody from Cell Signaling catalog #2451 (1:1,000); AMPK alpha Antibody from Cell Signaling catalog #2532] overnight at 4°C. After quick washes with TBS containing 0.1% Tween (TBS-T), the membranes were incubated with secondary antibodies for 2 h at room temperature and again washed with TBS-T. Specific bands of immunoreactive proteins were visualized in a VersaDoc Imaging System (Bio-Rad) after 5-min incubation with enhanced chemifluorescence. The density of protein bands was calculated using the Quantity One Program (Bio-Rad). Results were expressed as the ratio of phosphorylated AMPK to AMPK total.

**Determination of acetylcholinesterase (AChE) activity in brain mitochondria-free cytosolic fraction**

AChE activity was evaluated in brain mitochondria-free cytosolic fraction, as previously described [19]. Brieﬂy, 2 mL of phosphate buffer (0.1 M KH₂PO₄, pH 7.4) supplemented with 10 μL of brain homogenate, 0.1 mM ethopropazine and 10 μL of Ellman’s reagent (0.15 mM 5,5-dithio-bis-2-nitrobenzoic acid (DTNB)) in 0.1 M phosphate buffer, pH 7.4) were incubated at 25°C for 2 min. The reaction was initiated by adding substrate acetylthiocholine and the catalytic activity was measured by following the increase in absorbance at 412 nm for 3 min, using a Cary 50 UV-Vis spectrophotometer (Agilent Technologies, Santa Clara, CA, USA). AChE activity was evaluated as a function of substrate concentration, using seven concentrations of acetylthiocholine iodide, from 16 to 2000 μM, to determine the Michaelis–Menten kinetic parameters Kᵣ (Michaelis constant) and Vₘₐₓ (maximal velocity).
Triton X-100, 10 μM DTNB, 1 mM oxaloacetate, and 0.2 mg of mitochondrial protein. The reaction was initiated by adding 0.37 mM acetyl-CoA and monitored at 412 nm in a Cary 50 UV-Vis spectrophotometer (Agilent Technologies, Santa Clara, CA, USA). The specific activity of CS was calculated by DTNB (extinction coefficient 13,600 M\(^{-1}\) cm\(^{-1}\)) reduction rate, determined in the linear range of the plot and expressed as nmol/min/mg protein.

Complex I (NADH-ubiquinone oxidoreductase) was evaluated spectrophotometrically following reduced nicotinamide-adenine dinucleotide (NADH) oxidation. NADH (50 μM) fluorescence intensity was evaluated before and after adding 162.5 μM decylubiquinone to the reaction medium (25 mM KH\(_2\)PO\(_4\), 10 mM MgCl\(_2\), pH 7.4) supplemented with 1 mM KCN and 0.3 mg of mitochondrial suspension. The reaction was followed in a Varian Eclipse fluorescence spectrophotometer (Agilent Technologies, Santa Clara, CA, USA) at 450 nm, setting excitation at 366 nm. Enzyme activity was determined by the difference between the slopes before and after 3.0 μM rotenone (specific complex I inhibitor) addition. This value was expressed in arbitrary units/min/mg protein.

Complex II (Succinate Dehydrogenase) activity was assessed monitoring 2,6-dichlorophenolindophenol (DCIP) reduction at 600 nm in a 25 mM potassium phosphate buffer pH 7.4 containing 0.2 mg of mitochondrial protein, 2 mM KCN, 6.5 μM antimycin A, 0.05 mM DCIP, and 0.1 mM decylubiquinone. The reaction was initiated by adding 20 mM succinate and stopped 5 min later by adding 0.5 mM oxaloacetate (complex II inhibitor). The extinction coefficient used for DCIP was 21,000 M\(^{-1}\) cm\(^{-1}\).

Complex IV (cytochrome c oxidase) activity was measured by the oxygen consumption in a Clark-type oxygen electrode (Hansatech, Norfolk, UK) associated to cytochrome c oxidation promoted by 5 mM ascorbate/2.5 mM N,N,N',N'-tetramethyl-1,4-benzendiamine dihydrochloride (TMPP). The reaction was initiated by addition of ascorbate/TMPD into standard buffer (130 mM sucrose, 50 mM KC\(_2\), 5 mM MgCl\(_2\), 5 mM KH\(_2\)PO\(_4\), and 5 mM HEPES; pH 7.2) supplemented with 0.2 mg mitochondrial suspension, 3 μM rotenone, 0.1 μM antimycin A (inhibitor of complex III) and 15 μM cytochrome c. KCN (2 mM; specific inhibitor of complex IV) was used to stop the reaction. Complex IV activity was expressed as nmol O\(_2\)/min/mg protein.

\(P_{\text{FoF}}-\text{ATPase}\) activity was determined using a Crison pH evaluation system (Barcelona, Spain) connected to a Kipp and Zonen recorder (Omni Instruments, Dundee, UK), as previously described [34]. Assay buffer (130 mM sucrose, 60 mM KCl, 0.5 mM Hepes, and 2.5 mM MgCl\(_2\); pH 7.0) was supplemented with 3 μM rotenone and 0.2 mg of mitochondrial protein. The reaction was initiated by adding 2 mM ATP-Mg and stopped by adding 1 μg oligomycin, since ATP hydrolysis results in transmembrane proton translocation. At the end of each reaction assay system calibration was performed using a standard HCl solution. \(P_{\text{FoF}}-\text{ATPase}\) activity was expressed as nmol H\(^+\)/min/mg protein.

### Lipid extraction and phospholipid quantification

Phospholipids extraction from non-synaptic and synaptic mitochondria was performed using a double extraction procedure with a solvent combination of methanol/chloroform/water at final proportions of 2:1:0.8 (v/v/v) [33]. A volume of mitochondrial suspension corresponding to 5 mg of protein was reconstituted in 0.8 mL aqueous buffer. Internal standards were added based on mitochondrial protein concentration (per mg of protein): 0.7 μg phosphorylcholine, PC (14:0/14:0); 0.2 μg sphingomyelin, SM (d18:1/12:0); 0.2 μg lyso-phosphatidylcholine, LPC (14:0/0:0), 0.3 μg phosphatidylinositol, PI (16:0/16:0); 0.45 μg phosphatidylserine, PS (14:0/14:0); 0.35 μg phosphatidylinositol, PI (14:0/14:0); 0.34 μg cardiolipins, CL (14:0/14:0/14:0); and 0.3 μg phosphatidylglycerol, PG (14:0/14:0). Mitochondrial samples with internal standards were mixed with 2 mL methanol for 5 min and then 1 mL chloroform was added. The mixture was blended for more 5 min and then centrifuged at 2000 g for 5 min at 4°C in order to obtain a two-phase system: an aqueous top phase and an organic bottom phase from which lipids were collected. The palette obtained in the first centrifugation was used to perform lipid re-extraction.

In order to quantify membrane phospholipids content, phosphorus assay was performed according to the method outlined by Bartlett and Lewis [35]. Briefly, 0.65 mL of concentrate perchloric acid (70% v/v) were added to lipid samples (50–100 μL) previously dried with nitrogen stream and then incubated at 180°C for 2 h. 3.3 mL of H\(_2\)O, 0.5 mL of 1% (v/v) ammonium molybdate, and 4% (v/v) ascorbic acid were added to all samples, followed by incubation at 100°C.
for 10 min in a water bath. Standards from 0 to 250 nmol phosphate underwent the same sample treatment. Standards and samples absorbance was measured at 800 nm in a Varian Cary 50 spectrophotometer. After quantification, extracted lipids were split per glass ampoules in pools of 322.5 nmol (10 μg phosphorous). Samples were dried under nitrogen atmosphere and stored at −20°C in sealed glass ampoules. Before use, each sample was dissolved with 100 μL 2:1 chloroform/methanol (v/v) mixture.

Separation and quantification of phospholipid classes by high pressure liquid chromatography - mass spectrometry (HPLC-MS)

Phospholipid classes were solved on an HPLC system (Waters Alliance 2690) coupled to an electrospray linear ion trap mass spectrometer (ThermoFinnigan, San Jose, CA, USA) [36]. The mobile phase A consisted of 10% water and 55% acetonitrile with 35% (v/v) methanol. The mobile phase B consisted of acetonitrile 60%, methanol 40% with 10 mM ammonium acetate. Total lipid extract was diluted in the mobile phase A (1:1, v/v) and introduced into an Ascentis Si HPLC Pore column (15 cm × 1.0 mm, 3 μm) (Sigma-Aldrich). Solvents gradient started with 0% of phase A followed by a linear increase to 100% of phase A during 20 min, and held isocratically for 35 min, returning to the initial conditions in 5 min. The flow rate through the column was 16 μL/min. obtained using a pre-column split (Acurate, HPLC Packings, USA).

Mass electrospray analysis was done using the following parameters: source voltage, 5 kV, positive mode, 4.7 kV, negative mode, source temperature, 275°C, sheath gas flow, 8 U, source current, 100 ua, capillary voltage, 10 V, positive mode, −43 V negative mode. An isolation with of 0.5 Da was used with a 30-ms activation time for MS/MS experiments. Data acquisition and treatment of results were carried out with an Xcalibur data system (V2.0, ThermoFinnigan, San Jose, CA, USA). Quantitation of individual molecular phospholipid species were calculated by determining the ratio of the peak area of each molecular species to the corresponding internal standard.

Separation of cardiolipin by thin layer chromatography (TLC) and its analysis by electrospray mass spectrometry (ESI-MS)

Phospholipid classes from the total lipid extracts were separated by TLC using silica gel plates [19]. Prior to separation, plates were washed with chloroform/methanol (1:1, v/v) and treated with 2.3% (w/v) boric acid in ethanol. The plates with spots containing between 20 and 30 μg of phosphorous were developed in solvent mixture chloroform/ethanol/water/triethylamine (35:30:7:35, v/v/v/v). Lipid spots on TLC plates were detected with a UV lamp (λ = 254 nm) after primuline (50 μg/mL in acetone/water, 80:20, v/v) exposure. CL was identified by using a CL standard and correspondent TLC spot was scraped off the plates to lipid extraction and further electrospray mass spectrometry analysis. CL lipid extracts were directly analyzed by ESI-MS in negative mode in an electrospray linear ion trap mass spectrometer (ThermoFinnigan, San Jose, CA, USA), following the conditions indicated above for lipid analysis in negative ion mode. CL MS analyses were performed in three independent samples.

Statistical analysis

All the data presented in this study correspond to the mean value of n independent experiments ± SD, with n ≥ 3. In each independent experiment at least three animals were used. Comparison of two data sets was performed with one-way ANOVA test. Data sets were considered statistically different when p < 0.05.

RESULTS

Non-synaptic and synaptic mitochondrial bioenergetics state

Decline of glucose metabolism has been identified as a characteristic phenotype in both AD patients and animal models for AD [37, 38]. Therefore, the overall brain adenine nucleotides levels and non-synaptic and synaptic mitochondria activities were evaluated using animals in presymptomatic condition (3 months of age) in order to assess whether brain bioenergetics is compromised at an early stage of the disease. Although the global levels of adenine nucleotides is similar in both groups, significant differences were detected in ATP (nonTg: 2.030 ± 0.134, 3xTg-AD: 1.620 ± 0.102) and AMP (nonTg: 0.275 ± 0.024, 3xTg-AD: 0.385 ± 0.042) levels, leading to an increase of AMP/ATP ratio in 3xTg-AD brains (Table 1). This increased AMP/ATP ratio may has functional consequences, since a parallel increase of phosphorylated AMPK/AMPK total ratio was detected (Fig. 1). Brain energy charge and ATP/ADP ratio were also determined and, despite the absence of significant differences, a slight decrease in these bioenergetic indicators was observed in 3xTg-
AD brains (Table 1). These results indicate specific alterations in AD brain, considering all brain areas. However, brain is a spatial and cellular regionalized metabolic organ and AD pathology exhibits a specific spatial brain progression [39]. Thus, the bioenergetic changes that were detected may not be the result of brain random effects, but rather reflect significant alterations in brain regions preferentially affected by the disease. On the other hand, cellular metabolic regionalization emerges as a consequence of several brain cell types and the existence of two brain mitochondria populations. Brain mitochondria fall into non-synaptic (in neuronal and glial cell bodies) and synaptic (in synaptic bottom of neurons) which differ in both structural and functional features, such as bioenergetic activity and mitochondrial membrane lipid composition [32].

In this view, non-synaptic and synaptic brain mitochondria were isolated and purified in order to discriminate their contribution to the bioenergetic deficit detected in 3xTg-AD mice. The purity degree of both non-synaptic and synaptic mitochondria populations (Supplementary Table 1) from 3xTg-AD animals exhibit lower activity than their age-matched controls. However, at this age (3 months old) these differences are of no statistical significance.

Table 1

<table>
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<th></th>
<th>Non-synaptic mitochondria</th>
<th>Synaptic mitochondria</th>
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<tr>
<td></td>
<td>NonTg</td>
<td>3xTg-AD</td>
</tr>
<tr>
<td>ATP (μmol/g wet brain)</td>
<td>2.03 ± 0.134</td>
<td>1.62 ± 0.102*</td>
</tr>
<tr>
<td>AMP (μmol/g wet brain)</td>
<td>0.67 ± 0.078</td>
<td>0.63 ± 0.097*</td>
</tr>
<tr>
<td>ADP (μmol/g wet brain)</td>
<td>0.27 ± 0.024</td>
<td>0.38 ± 0.042*</td>
</tr>
<tr>
<td>ATP+ADP+AMP (μmol/g of wet brain)</td>
<td>3.18 ± 0.157</td>
<td>2.82 ± 0.146</td>
</tr>
<tr>
<td>ATP/ADP</td>
<td>2.31 ± 0.255</td>
<td>1.96 ± 0.263</td>
</tr>
<tr>
<td>AMP/ATP</td>
<td>0.115 ± 0.015</td>
<td>0.237 ± 0.029*</td>
</tr>
<tr>
<td>Ratios</td>
<td>Brain energy charge</td>
<td>0.776 ± 0.066</td>
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<td></td>
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<td>0.718 ± 0.056</td>
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Results are expressed as mean ± std of three independent experiments. *Significantly different from nonTg, with p<0.05.
activity, since their ability to couple ATP hydrolysis, associated to transmembrane proton translocation, is similar to that detected in nonTg non-synaptic and synaptic mitochondria, respectively (Fig. 2). Differences between nonTg and 3xTg-AD mice, however, cannot be attributed to technical variability ensuing from isolation effects putatively associated with the number of independent assays, since studies on bioenergetics using 6 to 12 month old animals revealed that synaptic mitochondrial deficits detected at the age of 3 months are exacerbated with aging and extended to non-synaptic brain mitochondria [42].

Taken together, these data indicate that synaptic mitochondrial dysfunction is mainly associated with a decay in complex I activity that precedes non-synaptic mitochondrial dysfunction and may compromise the ATP-dependent pathways supporting synaptic transmission.
Table 2

<table>
<thead>
<tr>
<th></th>
<th>NonTg</th>
<th>3xTg-AD</th>
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<tbody>
<tr>
<td>CAT (nmol O₂/min/mg protein)</td>
<td>5.61 ± 0.10</td>
<td>4.78 ± 0.05</td>
</tr>
<tr>
<td>SOD (Uact/min/mg protein)</td>
<td>5.06 ± 0.22</td>
<td>4.99 ± 0.32</td>
</tr>
<tr>
<td>GSH/GSSG</td>
<td>1.87 ± 0.58</td>
<td>1.99 ± 0.21</td>
</tr>
<tr>
<td>ROS (nmol DCF/mg protein)</td>
<td>8.76 ± 0.34</td>
<td>9.98 ± 0.20  *</td>
</tr>
</tbody>
</table>

Results are expressed as mean ± s.d of three independent experiments. *Significantly different from nonTg, with p<0.05.

Phospholipid content of brain mitochondria

In order to assess mitochondrial membrane lipid profile and evaluate its putative relationship with synaptic bioenergetic dysfunction detected in an AD animal model, phospholipids were extracted from non-synaptic and synaptic mitochondria and analyzed by means of lipidomic approaches. Mitochondrial membrane phospholipid composition was accomplished through HPLC-MS.

Phospholipid profile of synaptic brain mitochondria

Lipid extracts of synaptic and non-synaptic mitochondria were analyzed using HPLC-MS/MS in both positive- and negative-ion modes. Typical positive-ion ESI mass spectra and the relative abundance of most representative molecular species of choline-containing-phospholipids (i.e., PC, LPC and SM) from synaptic mitochondrial populations of 3xTg-AD and NonTg matched controls (Fig. 3). The relative abundance of phospholipid classes of non-synaptic mitochondrial membranes is similar in healthy and 3xTg-AD mouse brains and is characterized by PC, PE, and CL dominance which together represent more than 75% of all phospholipids (Fig. 3A). On the other hand, 3xTg-AD synaptic mitochondria exhibited an altered phospholipid composition detected in choline phospholipids (PC and LPC) and CL contents (Fig. 3B). PC and LPC relative abundance is significantly higher in 3xTg-AD group (PC: nonTg 31.91 ± 1.41, 3xTg-AD 39.84 ± 3.34; LPC: nonTg 0.78 ± 0.29, 3xTg-AD 1.63 ± 0.68). Instead, synaptic mitochondrial CL content is significantly decreased in 3xTg-AD animals regarding healthy controls (nonTg 18.20 ± 1.38, 3xTg-AD 11.97 ± 1.31).

Fig. 3. Relative abundance (mol%) of each phospholipid classes in total lipid extract obtained from non-synaptic (A) and synaptic (B) brain mitochondria isolated from nonTg (dark bars) and 3xTg-AD (gray bars) mice. PC phosphatidylcholine, LPC lyso-phosphatidylcholine, SM sphingomyelin, PE phosphatidylethanolamine, PS phosphatidylserine, CL cardiolipin, PA phosphatidic acid, PG phosphatidylglycerol, PI phosphatidylinositol. Error bars represent standard deviation for n=3 independent experiments using 3 animals each in one. The amount of each phospholipid class was calculated by the sum of its individual species identified and quantified by HPLC-MS/MS, as described in methods section. Considering the results as total nmols of phospholipids per mg of protein: non-synaptic mitochondria: nonTg 366.9 ± 33.2 and 3xTg-AD 384.2 ± 30.3; synaptic mitochondria: nonTg 334.0 ± 13.9 and 3xTg-AD 351.2 ± 14.3. *Significantly different from nonTg group, with p<0.05.
Fig. 4. Molecular profile of choline lipids extracted from nonTg (black bars) and 3xTg-AD (gray bars) synaptic mitochondria. A) Typical ESI-MS in positive mode with formation of [M+H]^+ ions of PC, LPC, and SM. Distribution of (B) PC, (C) LPC and SM molecular species. Error bars represent standard deviation for n=3 independent experiments using 3 animals in each one. *Significantly different from nonTg group, with p ≤ 0.05.

Identification of m/z 496 (16:0), 731 (36:1), and 760 (34:1) as the predominant ions of LPC, SM, and PC phospholipid classes, respectively. Relative abundance of PC molecular ions at m/z 760 (34:1), 782 (36:4), and 810 (38:4) is increased in 3xTg-AD group compared to nonTg group (Fig. 4B). Indeed, PC molecular ion at m/z 760 (34:1) comprises higher fatty acid diversity in 3xTg-AD PC (16:0/18:1) and PC (16:1/18:0) than in nonTg PC (16:0/18:1). In contrast, 3xTg-AD PC ions at m/z 782 (36:4) and 810 (38:4) lose fatty acid diversity by lacking PC (18:0/18:4) and PC (18:1/20:3), respectively, as indicated in Supplementary Table 3. The relative abundance of LPC molecular ions at m/z 496 (16:0), 518 (18:3), 550 (20:1), and 568 (22:6) and of SM molecular ions at m/z 703 (34:1) and 733 (36:0) was found to have increased in 3xTg-AD synaptic mitochondria (Fig. 4C). Fatty acid composition of LPC and SM molecular ions was preserved in 3xTg-AD synaptic mitochondrial membranes compared to healthy controls (Supplementary Table 3). On the other hand, choline-containing phospholipids profile of 3xTg-AD non-synaptic brain mitochondria was similar both in composition and diversity to that detected in nonTg matched controls (Supplementary Figure 2).

HPLC-MS/MS analyses were also performed in negative-ion mode in order to access the molecular profile of phospholipid classes PS, PE, PA, PG, PI, and CL from nonTg and 3xTg-AD synaptic mitochondrial membranes (Figs. 5 and 6). Typical ESI-MS
Fig. 5. Molecular profile of PS, PE, PA, PG, and PI extracted from nonTg (black bars) and 3xTg-AD (gray bars)-synaptic mitochondria. Typical ESI-MS of (A) PS, (B) PE, (E) PA, (F) PG, and (H) PI obtained in negative ion-mode. Distribution of (C) PS and (D) PE and (G) PA, PG, and PI molecular species. Error bars represent standard deviation for n = 3 independent experiments using 3 animals in each one. *Significantly different from nonTg group, with p \leq 0.05.
Fig. 6. Molecular profile of CL extracted from nonTg (black bars) and 3xTg-AD (gray bars) synaptic mitochondria. A) Typical ESI-MS of CL obtained in negative mode with formation of [M-H]⁻ ions. B) Distribution of CL molecular species with differences between nonTg and 3xTg-AD groups emphasized into insert (b). Error bars represent standard deviation for n = 3 independent experiments using 3 animals in each one. *Significantly different from nonTg group, with p ≤ 0.05.

Spectra and correspondent quantifications of PS and PE molecular species are displayed in Fig. 5A–D. Data are consistent, revealing that ions at m/z 834 PS (40:6), 766 PE (38:4), and 774 PE plasmalogen (p40:6) are the most representative molecular species within the respective phospholipid class in synaptic
brain mitochondria of both groups. Although PS profile of synaptic mitochondria was similar in both animal groups (Fig. 5A, C), PE profile showed significant changes, especially visible in the relative abundance of diacyl-PE species at m/z 762 (38:6) and plasmalogen species at m/z 726 (p36:2), 748 (p38:5, o38:6), 750 (p38:4), 772 (p40:7), and 774 (p40:6) in the 3xTg-AD group (Fig. 5D), in which they significantly decreased. Regarding the minor anionic phospholipids (Fig. 5E–H), they showed similar PA, PG, and PI profiles for both animal groups, although the relative abundance of PA ions at m/z 723 (16:0/22:4, 18:0/20:4) and 727 (18:1/20:1, 18:0/20:2) was higher and lower respectively, in 3xTg-AD mice. The fatty acid diversity of each PS, PE, PA, PG, and PI molecular ions obtained from synaptic mitochondria is listed in Supplementary Tables 4–6.

Non-synaptic mitochondrial membranes showed no significant differences in the quantitative analysis of PS, PE, PA, PG, and PI molecular species (Supplementary Figure 5) with the exception of PE plasmalogen ion with m/z 726 (p36:2), whose relative abundance declined in the 3xTg-AD group. In this mitochondria population, the pathological condition did not affect the fatty acid diversity in the phospholipid classes analyzed. Furthermore, the essential nature of fatty acid diversity within each phospholipid class was similar in both nonTg brain mitochondrial populations.

**CL analysis**

CL is a unique tetra-acyl phospholipid found almost exclusively in the inner mitochondrial membranes and is essential for maintaining mitochondrial functionality and membrane integrity [43]. Specifically, CL has shown to be required for the proper structure and activity of mitochondrial electron transport chain enzymes [22, 44]. Not surprisingly, due to its importance in mitochondrial function, alteration of normal CL metabolism has been associated with several pathologies, including neurodegenerative diseases [43]. CL profiling observed in MS spectra of nonTg and 3xTg-AD synaptic mitochondria are shown in Fig. 6A. NonTg singly charged [M-H]– ions, correspondent to CL molecular species, are represented at least by seven major clusters with a variety of fatty acid residues in synaptic mitochondria. Interestingly, this pattern was lost in MS spectrum obtained for CL of 3xTg-AD synaptic mitochondria, mainly due to a decline in the most representative molecular species within each cluster, as reflected in CL molecular species quantification (Fig. 6B, b). Additionally, in MS spectra of Fig. 6A, monolysocardiolipins were not detected in synaptic mitochondria of either nonTg and 3xTg-AD mice, since on m/z region, between 1100 and 1380, only the species with m/z at 1239 (internal standard, CL(14:0)4) reach the minimum intensity to be identified. Despite the aberrant CL profile exhibited by 3xTg-AD synaptic mitochondria, the molecular species diversity was preserved (Supplementary Table 7).

Detailed analyses were performed in order to better understand the putative functional impact of this aberrant CL profile on mitochondrial membranes, taking into consideration the number of unsaturated bounds (Fig. 7A) and the carbon chain length (Fig. 7B), as well as the differences between the aberrant CL profile of synaptic mitochondria isolated from nonTg (black bars) and 3xTg-AD (gray bars) mice. *Significantly different from nonTg group, with p ≤ 0.05.
profile exhibited by 3xTg-AD synaptic mitochondria and the predicted CL profile, bearing in mind that the decreased CL content exhibited by synaptic mitochondria of 3Tg-AD mice occurs only as a function of the relative abundance of each CL species in nonTg mice (Supplementary Figure 6). These data show a decrease of specific CL species regarding the number of unsaturated bonds and carbon chain length, which points to a tendency toward normalization of its relative abundance. In other words, the predominant CL species with a specific number of unsaturated bonds or with a specific carbon chain length that were found in the nonTg group disappeared in the 3Tg-AD group, reaching a relative abundance that is similar to the one observed among minor CL species. Therefore, the abnormal CL profile of synaptic mitochondria of 3Tg-AD mice does not result from a regular decrease associated with low levels of total CL content in this group.

CL profile of non-synaptic mitochondria was also analyzed by MS; the differences between nonTg and 3Tg-AD groups found in this mitochondria population are not as extensive as those found in synaptic mitochondria (Supplementary Figure 4). In fact, the CL profile of non-synaptic mitochondria is just affected by the pathological situation of some CL molecular species, namely the ones that correspond to [M-H]- ions at m/z 1399 (68:4); 1421 (70:7); 1423 (70:6); 1447 (72:8); 1453 (72:5); 1517 (76:1); and 1531 (78:8), whose relative abundance differs from that of the nonTg group. Additionally, fatty acid diversity within each CL species is similar in both non-synaptic and synaptic brain mitochondria from nonTg mice (Supplementary Table 7).

**Brain oxidative stress state and acetylcholinesterase activity**

In order to characterize the overall redox state of 3-month old nonTg and 3Tg-AD mice brains, we evaluated the activity of antioxidant enzymes (SOD and CAT), GSH and GSSG contents, and ROS levels generated by brain mitochondria-free cytosolic fractions. Table 2 shows that SOD and CAT activities and GSH and GSSG levels are similar in nonTg and 3Tg-AD groups. Nevertheless, in 3Tg-AD brains, a significant increase in ROS levels was observed (13%) compared to nonTg group, which indicates oxidative stress events are more likely to occur in 3Tg-AD group.

AChE activity was also evaluated in the brain mitochondria-free cytosolic fractions, since its pharmacological modulation is a usual therapeutic strategy used in AD. Although AChE activity was slightly increased in 3Tg-AD brains, the Michaelis–Menten enzymatic kinetics parameters (Vmax and Km) were not significantly different from nonTg matched controls, as shown in Supplementary Figure 5.

**DISCUSSION**

The dynamic regulation of brain functions requires a high energy flux dependent on both glucose metabolism and mitochondrial function [45]. In this way, brain bioenergetic decline has been involved in the neurodegenerative pathways and consequent synaptic failure in AD [37, 46]. In the present study, bioenergetic analyses revealed the existence of dysfunctional synaptic mitochondria associated with the decay of complex I activity which contributes to ATP deficit and an increased phosphorylated AMPK/AMPK total ratio detected in 3Tg-AD brains (Table 1 and Fig. 1). This brain energy deficit promotes AMPK activation, triggering not only a metabolic reprogramming to restore the brain energy homeostasis but also changes in cell signaling pathways connected to mitochondrial biogenesis and autophagy [47]. Therefore, at an early stage of AD, brain mitochondrial dysfunctions are primarily detected in synaptic mitochondria that precede the extracellular deposition of Aβ and the formation of intracellular neurofibrillary tangles [24]. This is concurrent with the report by Yao and collaborators of a brain mitochondrial respiration impairment and decreased pyruvate dehydrogenase protein levels in 3-month old 3Tg-AD female mice [48]. Alterations of brain mitochondrial functions were also detected in 3-month old Thy-1 AβPP mice, characterized by a decrease in mitochondrial transmembrane potential and ATP levels [49]. A more detailed study, using the same mice model, showed that brain mitochondria functionality disruption associated with declined complex IV activity is earlier detected in synaptic population [46]. Postmortem analysis of brain tissue from AD patients also detected anomalous complex IV and pyruvate dehydrogenase expression and activities [50], highlighting the role of mitochondrial dysfunctions in disease pathogenesis.

The differences in CS normalized complex I activity detected in the synaptic mitochondria of nonTg and 3Tg-AD mice may result from several factors, including: i) selective changes in the relative complex I membrane content as a consequence of downregulation of protein expression and/or synthesis; and ii) changes...
in the lipid composition of mitochondrial membranes which regulate the membrane assembling and activity of mitochondrial enzymes complexes [19, 20, 33]. Complex I (similarly to complexes III and IV and FoF1-ATPsynthase) is a multi-subunit enzyme with some subunits encoded by mitochondrial DNA and others (majority) encoded by nuclear genome, which are mainly synthesized in perinuclear cellular regions. The spatiotemporal distribution of mitochondria in neurons is dynamically regulated by mitochondrial traffic through axon that includes mitochondria either moving primarily anterogradely and retrogradely or appearing to be stationary for an extended period in cellular regions with high ATP demand (e.g., synapses) [51]. Since complex I activity in non-synaptic mitochondria is similar in both nonTg and 3xTg-AD mice, problems affecting protein expression and/or synthesis can be ruled out. Thus, it was our concern to ascertain whether 3xTg-AD brain bioenergetic deficit associated with decreased complex I activity was correlated with changes in the phospholipid composition of synaptic mitochondrial membranes. Data from lipidomic analyses show that 3xTg-AD synaptic mitochondria exhibit an altered phospholipid composition at two levels of complexity, characterized by: i) decrease of CL content with a concomitant increase of choline phospholipids (PC and LPC), ii) altered PC, LPC, PE (mainly plasmalogens), and CL molecular profiles. It is interesting to note that changes of phospholipid class content were not random but resulted from modifications in the relative abundance of specific molecular species, as is clearly evidenced by LPC (Fig. 4), ethanolamine plasmalogens (Fig. 5), and CL (Fig. 6) profiles.

The higher LPC content suggests that intracellular phospholipases A2 (iPLA2) may be upregulated in synaptic bottom of neurons of transgenic animals with an expected impact on lipid singling processes. Detailed analysis of LPC diversity (Supplementary Table 3) supports the idea that increased membrane LPC at m/z 496, 520, 522, 524, and 546 may result from hydrolysis of PC at m/z 782, 806, 808, 810, and 832 by iPLA2 with the consequent release of free arachidonic acid (C20:4), a well-known precursor of pro-inflammatory lipid signaling molecules, such as prostaglandins and thromboxanes. In fact, increased LPC content and oxidized derivatives of arachidonic acid have been involved in brain inflammatory processes which are characteristic of AD [54].

Ethanolamine plasmalogens seem to be essential to preserve the functional organization of the membrane and protect it against oxidative stress [54]. The functional relevance of membrane plasmalogens is also well evidenced by the positive relationship between low levels of brain membrane plasmalogens and severe human dementia [55]. Brain mitochondrial plasmalogens comprise about 50% of PE content; however, their specific role in brain mitochondria function remains poorly understood. Profile changes in ethanolamine plasmalogens found in synaptic mitochondria of 3xTg-AD mice, occurring simultaneously with mitochondrial bioenergetic deficits, may contribute to the characteristic metabolic phenotype of AD.

CL, a specific mitochondrial phospholipid, is synthesized from condensation of PG and cytidinediphosphate-diacylglycerol (CDP-DAG) to form immature CL [18]. However, the functional mitochondrial CL profile results from remodeling processes that involve sequential phospholipase (deacylation) and transacylase/acyltransferase activities and use PE, PC, and Acyl-CoA as fatty acids donor to produce mature species [56]. CL content and diversity are intimately linked to functional membrane assembly of redox complexes of the respiratory system and phosphorylation apparatus regulating cells’ energetic requirements [20, 44]. Besides, the brain exhibits a CL profile with the highest molecular diversity, comprising about 100 different species organized in at least seven clusters [19, 57]. These are eventually necessary to support the highly dynamic energy flux required for its proper functioning. 3xTg-AD synaptic mitochondria lose the CL cluster-organized profile due to a decline of the most representative molecular species, which may have functional consequences. In this view, decreased CL content shown in 3xTg-AD synaptic mitochondria (Fig. 6) may reflect a decrease in CL synthase activity and/or expression, or enhanced CL degradation pathways. Since CL profile of non-synaptic brain mitochondria of 3xTg-AD mice is similar to that exhibited by nonTg mice (Supplementary Figure 4), CL synthase expression and/or activity should, therefore, not be affected. Therefore, the decline of more representative CL species may result from changes in remodeling pathways or increased degradation of selective species.

The pathological role of altered CL metabolism associated with mitochondrial dysfunction and increased mitochondrial apoptotic cascade pathway has been documented in several disorders, such Barth Syndrome. In those cases, the alteration of CL profile is believed to be the primary molecular manifestation emerging from tafazzin deficiency [58]. Along with iPLA2, tafazzin is considered a key enzyme in CL remodeling process and its cellular deficiency is...
characterized by an increase of monolysocardiolipin levels and a decrease of CL content and of the average number of double bonds per cardiolipin molecule [59]. Although 3xTg-AD synaptic mitochondria exhibit an aberrant CL profile and lower CL content, the average number of double bonds per cardiolipin molecule (7.6) is similar to that found in nonTg synaptic brain mitochondria (Fig. 7).

CL content loss also contributes to the age-related phenotype generally associated with mitochondrial dysfunction and oxidative stress [60]. Our results indicate that 3-month old 3xTg-AD mice are able to preserve brain redox state as suggested by GSH/GSSG ratio and CAT and SOD activities (Table 2). In turn, the augmented ROS generation detected in 3xTg-AD brains (Table 2) could lead to a ROS-dependent oxidation of selective molecules, with a putative contribution to the abnormal CL profile. However, the mitochondrial accumulation of oxidized CL should not be particularly relevant, since those species were neither detected by HPLC-MS/MS analyses of total lipid extracts nor by direct ESI-MS/MS analyses of TLC obtained CL.

Taken together, bioenergetic and lipidomic data support the idea that synaptic mitochondrial dysfunction, resulting essentially from a decay of complex I activity, and an aberrant mitochondrial membrane phospholipid composition, mainly detected in CL profile, are interconnected key players contributing to the energetic deficit detected in early stages of AD.

Lipid and mitochondrial bioenergetic metabolisms are functionally interdependent and our data, obtained at the age of three months, do not allow us to discriminate where the dysfunction first occurs. Thus, synaptic mitochondrial lipid changes and bioenergetic deficit may compromise the ATP-dependent biochemical pathways with several functional consequences, including the loss of lipid asymmetry in synaptic membrane. The loss of lipid asymmetry increases the concentration of aminophospholipids (PS and PE) in the outer leaflet of the membrane, leading to a decrease in the lateral membrane organization, making PP accessible to hydrolysis by BACE and causing Aβ formation (Aβ pathology); and ii) decrease of the negative charge density on the cytosolic surface of the synaptic membrane, lessening its ability to bind and regulate intracellular Ca2+ fluxes that are instrumental in releasing neurotransmitters and an increase of free cytosolic Ca2+ and the associated neurotoxicity (e.g., tau pathology and upregulation of calcium-dependent iPLA2).

To conclude, our study is pioneer insofar as it shows, for the first time, that synaptic mitochondrial defects concomitant with extensive alterations in CL profile represent key early alterations in the development of AD. In fact, dysfunction of synaptic mitochondria and energy depletion associated to a loss of lipid asymmetry contribute to Aβ accumulation and cell dysfunction.

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SUPPLEMENTARY MATERIAL

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