Free-floating adult human brain-derived slice cultures as a model to study the neuronal impact of Alzheimer’s disease-associated Aβ oligomers

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ABSTRACT

Background: Slice cultures have been prepared from several organs. With respect to the brain, advantages of slice cultures over dissociated cell cultures include maintenance of the cytoarchitecture and neuronal connectivity. Slice cultures from adult human brain have been reported and constitute a promising method to study neurological diseases. Despite this potential, few studies have characterized in detail cell survival and function along time in short-term, free-floating cultures.

New Method: We used tissue from adult human brain cortex from patients undergoing temporal lobectomy to prepare 200 μm-thick slices. Along the period in culture, we evaluated neuronal survival, histological modifications, and neurotransmitter release. The toxicity of Alzheimer’s-associated Aβ oligomers (AβOs) to cultured slices was also analyzed.

Results: Neurons in human brain slices remain viable and neurochemically active for at least four days in vitro, which allowed detection of binding of AβOs. We further found that slices exposed to AβOs presented elevated levels of hyperphosphorylated Tau, a hallmark of Alzheimer’s disease.

Comparison with Existing Method(s): Although slice cultures from adult human brain have been previously prepared, this is the first report to analyze cell viability and neuronal activity in short-term free-floating cultures as a function of days in vitro.

Conclusions: Once surgical tissue is available, the current protocol is easy to perform and produces functional slices from adult human brain. These slice cultures may represent a preferred model for translational studies of neurodegenerative disorders when long term culturing in not required, as in investigations on AβO neurotoxicity.

1. Introduction

The sporadic form of Alzheimer’s Disease (AD) accounts for more than 95% of AD cases worldwide (Alzheimer’s Association, 2016). Mounting evidence implicates soluble aggregates of the Aβ peptide, along with hyperphosphorylated Tau, as the main neurotoxins in AD.
(reviewed in Glabe, 2006; Ferreira and Klein, 2011; Viola and Klein, 2015). However, the complexity of the molecular and cellular mechanisms underlying AD has hampered the development of effective diagnostic and therapeutic approaches (Mucke and Selkoe, 2012; Nelson and Alafuzoff, 2012; Puzo and Gulisano, 2015a, 2015b). Conceivably, this limited progress is in part due to the extensive use of disease models such as transgenic mice overexpressing the amyloid precursor protein, APP, which do not fully recapitulate the complexity of sporadic AD mechanisms (Puzo and Gulisano, 2015a, 2015b). In this context, the availability of slice cultures from adult human brain is of great interest.

Organotypic cultures, herein termed slice cultures, have been defined as three-dimensional tissue cultivated on, or within, a gel support (e.g., collagen or artificial ECM) or free floating in culture medium (Hoffman, 2001). Slice cultures from different tissues have been reported, including skin, lung and brain (Hoffman, 2001; Steinstraesser and Rittig, 2009; Hum pel, 2015). The main advantage of slice cultures over dissociated cell cultures is the preservation of the original tissue architecture and connections. This feature has key implications for the application of slice tissue cultures in bioscience. Notably, slice cultures are more resistant to the action of some drugs or toxins than dissociated, 2D-cultures (Miller and Miller, 1985; Elkayam and Amitay-Shaprut, 2006; Olinga and Schuppnan, 2013), and respond differently to physiological stimuli as insulin compared to organotypic cultures from liver tissue (Dash and Figler, 2017).

Classic protocols for culturing post-natal rodent brain slices were described many years ago (Gahwiler, 1981; Stoppini and Buchs, 1991), and have been the basis for the development of innovative protocols. More recently, successful cultures of adult human brain tissue using either postmortem or surgical explants have been reported (Brewer and Espinosa, 2001; Verwer and Hermens, 2002; Dai and Buijs, 2004; Freiman and Surges, 2006; Risher and Lee, 2011; Sebollela and Freitas-Correa, 2012; Eugene and Cluzeaud, 2014). Notably, amygdala-hippocampoctomy has been the source of tissue in most of the papers using slice cultures from non-postmortem adult human brain (e.g. Brewer and Espinosa, 2001; Sebollela and Freitas-Correa, 2012; Eugene and Cluzeaud, 2014; Schwarz and Hedrich, 2017), which is mostly due to the necessary removal of a non-epileptogenic area (temporal cortex) to provide access to the mesial structures to be resected. In those studies, tissue remained alive in culture for days to weeks, and two studies showed preserved electrical activity (Eugene and Cluzeaud, 2014). Despite some efforts aimed at characterizing the health and functionality of those slice cultures (reviewed in Jones and da Silva, 2016), limited information is currently available on adult human brain-derived histocultures. This is surprising considering that cultured adult human brain tissue should represent a preferred model for studying age-associated neurodegenerative diseases (Przedborski and Vila, 2003), as they are potentially free from artifacts introduced when postmortem tissue is used (Born and Matos, 2017) or derived from loss of the original neuronal connectivity in dissociated cultures.

Here our goal was to characterize the cell content, viability and neural activity in adult human brain-derived slices cultivated free-floating for short-term. This method distinguishes from previously reported protocols focused on long-term culturing, which usually require the use of customized media and semipermeable membranes. In addition, we evaluated the potential of such cultures to serve as a platform to study molecular events in age-associated human neuropathologies, using as a model the impact of AD-associated Aβ oligomers (Viola and Klein, 2015). We found that the majority of neurons in cultured slices are alive and neurochemically active up to at least four days in vitro, which allowed the detection of binding of Aβ oligomers. This study may help to broaden the application of slice cultures derived from adult human brain in studies of mechanisms of pathogenesis, with the potential to enhance the translational impact of novel therapeutics for human brain disorders.

2. Methods

2.1. Human samples

Cortical tissue was obtained from adult patients submitted to amygdalo-hippocampectomy for the treatment of refractory temporal lobe epilepsy (TLE), in collaboration with the Epilepsy Surgery Center (CirEp) at the Clinical Hospital of the Ribeirão Preto Medical School - University of São Paulo, or at the University Hospital at the Federal University of Rio de Janeiro. Preoperative mapping of epileptogenic foci was carried out by magnetic resonance imaging and video coupled to electroencephalography (video-EEG). In most cases, the epileptogenic focus was restricted to the mesial zone of the temporal lobe. In one patient, epileptogenic activity (EEG) was also observed at the lateral (neocortical) zone of the temporal lobe. Tissue was obtained from 33 adults (55% male) averaging 40.5 ± 11.3 years old, and comprised a 1 cm² sampling of the lateral temporal cortex which is removed to access the mesial structures of the temporal lobe. This study was approved both by the Ribeirão Preto Medical School and the Federal University of Rio de Janeiro Ethics Committees (HCRP #17578/15 and UFRJ #0069.0.197.000-05).

2.2. Tissue culture

The following procedure is a modification of the protocol described in Sebollela and Freitas-Correa (2012). Briefly, a fragment of lateral temporal cortex was collected at the surgical room immediately after resection, immersed in ice-cold oxygenated medium (50% v/v HBS [Hank’s Balanced Salt Solution] in Neurobasal A (Gibco) supplemented with 10 mM Hepes and glucose 3 mg/ml) and transported to the laboratory in approximately 15 min. Tissue was immersed in ice-cold oxygenated HBSS, pia mater was carefully removed, and the remaining fragment was sliced at 200 μm in a VT1000 s automatic vibratome (Leica) in such a way that each slice contained all the cortical layers and white matter. Slices were transferred to a Petri dish with cold oxygenated HBSS for trimming in an approximate ratio of 1/3 white matter to 2/3 grey matter, and plated in 24-well plates (1 slice/well) filled with 600 μL of Neurobasal A (Gibco) supplemented with 1% Glutamax (Gibco), 1% Penicillin/Streptomycin (Gibco), 2% B27 (Gibco) and 0.25 μg/mL Amphotericin B (Gibco). Free-floating cultures were incubated at 36 °C and 5% CO₂. By the end of day 0, 200 μL of the medium were replaced by fresh medium supplemented with 50 ng/mL BDNF (Sigma Aldrich). At day in vitro (DIV) 1, the volume of medium per well was reduced to 400 μL and thereafter one third of the volume was replaced by BDNF-supplemented fresh medium every 24 h.

2.3. Cell viability in cultured slices

Viability was assessed using the MTT (3-(4,5-dimethylthiazol-2-yl)-2,5-diphenyl tetrazoline bromide] assay (Mosmann, 1983). Briefly, MTT (50 μg/mL) was added at the indicated times and allowed to be metabolized for 3 h at 36 °C. Next, the slices were washed with PBS and frozen. Just before spectrophotometric reading, slices were homogenized in 200 μl isopropanol/HCl 0.01 N using a motorized pestle (Kimble Chase) and centrifuged at 1000 rpm for 1 min. Supernatants were collected and the optical density (O.D.) at 540 nm was determined using a microplate reader (Molecular Devices). O.D. values were normalized by the mass of each slice.

2.4. Histological analysis

Slices were fixed in buffered paraformaldehyde (4%) overnight at 4 °C, cryoprotected with 30% sucrose, frozen and cut at 30 μm in a freezing microtome (Leica). Sections were transferred to 24-well plates for free-floating Nissl staining with thionin as previously described (Kadar and Wittmann, 2009) or immunostaining. Immunostainings
were performed as previously described (Horta-Júnior Jde and Lopez, 2008) with modifications. Briefly, sections were incubated for 40 min in 2% normal donkey serum in phosphate buffer and incubated overnight with primary antibodies, anti-NeuN (Millipore, MAB377; 1:1000 dilution), anti-GFAP (Novocastra, NCL-GFAP-G5; 1:1000 dilution) or anti-β-tub1a (Abcam, ab178846; 1:2000 dilution). After incubation with avidin-biotin secondary antibodies (Vectorstain, Kit Standard, PK-4000, Vector), immunoreactivity was revealed using DAB + 0.04% nickel ammonium. Finally, sections were mounted on gelatin coated slides, air dried, dehydrated in increasing ethanol solutions, delipidified in xylene, and imaged.

2.5. Western blotting

Extracts from 1 to 2 slices were prepared in RIPA buffer (50 mM Tris—HCl, pH 7.4; 150 mM NaCl; 1.5 mM MgCl₂; 1.5 mM EDTA; 1% Triton X-100; 10% glycerol; supplemented with protease and phosphatase inhibitors) using a motorized pestle. Homogenates were centrifuged at 4 °C for 10 min at 16,000x g and supernatants were collected. Protein content was determined using the Bradford reagent (BioRad) and 50 μg were resolved on a 12% SDS-PAGE. The proteins were transferred to a nitrocellulose membrane using the transblot TURBO apparatus (Bio-Rad). The membrane was blocked at RT for 1 h with 5% non-fat dry milk in TBS (0.05 M Tris, 0.15 M NaCl, pH 7.5) plus 0.1% Tween 20 (Sigma-Aldrich), and incubated overnight with anti-NeuN (1:1000; MAB377, Millipore), and β-actin (1:10,000; MAB1501, Millipore). Primary antibodies were diluted in TTBS with 2.5% BSA (Sigma-Aldrich). After washing and incubation with HRP-conjugated secondary antibodies (GE Healthcare), membranes were developed using ECL-Prime (GE Healthcare). When necessary, membranes were stripped with 1% SDS, 100 mM glycine, 0.1% NP40, pH 2.0 for 40 min at RT and re-blocked before incubation with primary antibody. Quantitative analyses were made using NIH ImageJ software.

2.6. Neurotransmitter release assay

Neuronal depolarization was induced by transferring the slices from HBSS (15 min at 37 °C, corresponding to basal levels of neurotransmitter) to HBSS plus 80 mM of KCl (15 min at 37 °C, corresponding to KCl-induced released neurotransmitters). When indicated, slices were incubated with Ca²⁺/Mg²⁺-free HBSS. Media were collected and used for analysis of neurotransmitter levels. Both basal and KCl-evoked released GABA and L-glutamate were measured in slice culture media by high-performance liquid chromatography coupled to electrochemical detection (HPLC-ED, L-ECD-6 A module; Shimadzu), using a reverse-phase C-18 column (Kromasil; 5 μm particle size, 250 mm × 4.6 mm). OPA-sulfite derivatives of amino acids were generated immediately before sample injection (50 μL), as previously described (Monge-Acuna and Fornaguera-Trias, 2009). Standard curves were generated using L-glutamate and GABA standards (Sigma-Aldrich).

2.7. Binding of Aβ oligomers to human brain cultures

Aβ oligomers (AβOs) were prepared using Aβ1–42 (American Peptide) as in Chromy and Nowak (2003). At DIV 2, cortical slices were treated with AβOs 250 nM or vehicle for 1 h at 37 °C, washed twice with medium, and immediately after either fixed with 4% paraformaldehyde for IHC or homogenized under non-denaturing conditions for ELISA. In both cases, samples were probed using the anti-Aβ scFv-type antibody, NUsc1 (Velasco, Heffern et al. 2012). For IHC, fixed tissue was paraffin-embedded and cut at 5 μm. Thin sections were incubated with phage-bound NUsc1 (pβNUsc1) followed by HRP-conjugated anti-phage M13 (GE). Reactivity was revealed using DAB. ELISA was performed as in Sebella et al. (2017). Briefly, 96-well plates were coated overnight with 100 μL monoclonal anti-pan Aβ antibody (6E10, Covance) at 1.0 μg/mL in PBS at 4 °C, washed and blocked with 200 μL of 2% NFD milk/PBS. Detergent-free brain slice extracts were added at 0.5 mg/mL (total protein) diluted in 2% NFD milk/PBS and incubated overnight at 4 °C, followed by washing with PBS containing 0.05% Tween-20, and incubated with NUsc1 (diluted 1:100 in 2% milk-PBS) for 2 h at RT. Detection of bound antibody was achieved by incubation with HRP-conjugated anti-phage M13 (GE). Plates were developed using TMB (Sigma) as substrate. OD was read at 450 nm in a plate reader (Molecular Devices).

2.8. Total tau and p-tau levels in human brain cultured slices exposed to Aβ oligomers

Human brain slices at DIV7 were incubated with AβOs (500 nM) for 24 h, washed and homogenized in RIPA buffer containing protease inhibitors cocktail (Sigma-Aldrich) and 100 mM of the phosphatase inhibitor, sodium orthovanadate (Sigma-Aldrich), with a motorized pestle (Kimble Chasse). A 12% SDS-PAGE was loaded with 20 μg of total protein. Western blotting was carried out as described above, and membranes were incubated with anti-p-Tau Ser396 rabbiy polyclonal (1:1000; Santa Cruz Biotechnology), p-Tau Ser202/Thr205 mouse monoclonal (1:1000; Thermo Fisher Scientific), Tau-1 mouse monoclonal (1:1000; Merck-Millipore) and β-Actin mouse monoclonal (1:10,000; Santa Cruz Biotechnology). Quantitative analysis was made using ImageJ software (Schneider et al., 2012). The basal level (100%) of Tau phosphorylation was determined in slices treated with vehicle.

2.9. Statistical analysis

Statistical analysis was performed using Graph Pad Prism 6 (Graph Pad Software). Means were compared using either Student’s t-test or one-way ANOVA, as indicated in the legends for the figures. The number of donors has been established as the experimental unit (referred to as “N” throughout the text).

3. Results and discussion

3.1. Cell viability and tissue structure in cultured adult human brain slices

We initially employed the MTT assay, which has been successfully used to assess cell viability in a variety of histocultures (Tobita and Izumi, 2010; Wang and Peng, 2010; Paterniti and Impellizzeri, 2013), to evaluate cell viability in adult human brain slices along four days in culture. MTT reduction was not altered up to 4 days in vitro (DIV), indicating no extensive reduction in cell viability (Fig. 1A). A trend of decrease (albeit not statistically significant) in MTT reduction was verified at 4 DIV, suggesting that extending the culture period longer could result in reduced cell viability. Indeed, preliminary MTT data at DIV9 (N = 2 donors) indicate that viability at this stage is approximately 50% compared to DIV0 (data not shown). Preservation of cell viability up to 4 DIV obtained with the current protocol is likely a consequence of BDNF supplementation (Supp. Fig. 1) and the reduction in the thickness of the slices to 200 μm, compared to often used 300–400 μm slices (Sebella and Freitas-Correa, 2012; Eugene and Cluzeaud, 2014), which may have a positive impact on diffusion of O₂ and nutrients through the tissue. We have also observed that the rate of medium consumption (determined by acidity) dropped markedly by the end of DIV1, likely as a consequence of the metabolic recovery from processing and stabilization to the in vitro environment. Thus, we preferentially used DIV2 or later for treatments.

In line with preservation of cell viability, tissue architecture was not affected by tissue processing or by cultivation (Fig. 1B-D). Neuronal content in slices was assessed by both IHC and Western blotting (WB) for neuronal nuclear protein NeuN, a typical marker of mature and healthy neurons (Mullen and Buck, 1992). Neuronal morphology and tissue organization were preserved in cultured slices (Fig. 2A-D). In addition, WB analysis revealed no significant differences in NeuN levels
together, these data indicate that our free re-organization upon long-term cultivation in newborn rodent brain-preserved, were both con the presence of astrocytes and microglia, expected if tissue is indeed quantification. In combination, IHC and WB results indicate preserva-
flattening has not been reported in recent works using long-term adult human brain-derived slice cultures (Eugene and Cluzeaud, 2014; Andersson and Avaliani, 2016), mor-
phofunctional alterations triggered by plasticity-like events associated to the adaptation to the in vitro environment should not be completely ruled out. Supporting this notion is the observation by Eugene and Cluzeaud (2014) of a slight difference in electrophysiological patterns between slices at 9DIV and 24DIV. Altogether, it is possible to suggest that short-term human brain slice cultures might model with more accuracy the actual response of the mature human brain to neurotoxic challenges, such as the accumulation of Aβ oligomers in AD, than long-
dergan KCl depolarization in Ca2+-free medium. Results indicate that neurotransmitter release using this protocol was mainly synaptic, since release in the absence of calcium was signiﬁcant. Global neurotransmitter release measured under our conditions, we depolarization by KCl induced a greater than 5-fold increase in the release of the major excitatory and inhibitory neurotransmitters, glutamate and GABA, respectively (Fig. 3). Importantly, KCl-evoked re-
depolarization was signiﬁcantly reduced at both 1 and 2 DIV (Fig. 3, white bars). Collectively, these data suggest that neuronal activity in cultured slices is preserved at least until 4 DIV. In addition, considering the known role of astrocytes in the modulation of synaptic neurotransmitter release (Wallach and Lalouette, 2014; Sobieski and Jiang, 2015; Cazemier and Clascá, 2016), and of micro-
globin content and the formation and maintenance of neuronal synapses (Harry, 2013; Lim and Park, 2013), these functional data suggest that the main cell connections in the cortex are preserved in adult human-derived cultures (Humpel, 2015; Gilbride, 2016), and changes in gene/protein expression in the rodent mature brain (Staal and Alexander, 2011). Therefore, although flattening has not been reported in recent works using long-term adult human brain-derived slice cultures (Eugene and Cluzeaud, 2014; Andersson and Avaliani, 2016), mor-
along the period in culture (Fig. 2E-F). We note that the NeuN labeling pattern varied amongst donors, in some cases presenting bands mi-
gating at MW higher than 66 kDa, believed to be non-speciﬁc (Darlington and Goldman, 2008) and, for this reason, not included in quantification. In combination, IHC and WB results indicate preservation of neuronal number and viability in cultured slices. Importantly, the presence of astrocytes and microglia, expected if tissue is indeed preserved, were both conﬁrmed in slices at DIV4 (Fig. 2 G–H). Taken together, these data indicate that our free floating slices resemble the original in vivo environment, a key motivation for developing the current method. This is particularly relevant considering that slices cultured on membranes are known to undergo flattening and synaptic re-organization upon long-term cultivation in newborn rodent brain-
3.2. Functional analysis in cultured human brain tissue
In order to assess neurotransmitter release in the cultures, slices were subjected to KCl-induced depolarization followed by determination of neurotransmitter release to the medium. As expected, at 0 DIV, depolarization by KCl induced a greater than 5-fold increase in the release of the major excitatory and inhibitory neurotransmitters, glutamate and GABA, respectively (Fig. 3). Importantly, KCl-evoked release of neurotransmitters was preserved up to 4 DIV (Fig. 3).
To analyze the contribution of vesicular-dependent release to the global neurotransmitter release measured under our conditions, we carried out KCl depolarization in Ca2+-free medium. Results indicate that neurotransmitter release using this protocol was mainly synaptic, since release in the absence of calcium was signiﬁcantly reduced at both 1 and 2 DIV (Fig. 3, white bars). Collectively, these data suggest that neuronal activity in cultured slices is preserved at least until 4 DIV. In addition, considering the known role of astrocytes in the modulation of synaptic neurotransmitter release (Wallach and Lalouette, 2014; Sobieski and Jiang, 2015; Cazemier and Clascá, 2016), and of micro-

![Fig. 1. Cell viability and tissue structure in slice cultures from adult human cortex.](image)

**Fig. 1.** Cell viability and tissue structure in slice cultures from adult human cortex. (A) Viability in slices kept in culture for the indicated times. DIV: days in vitro. Viability was evaluated by the MTT assay. Representative slices incubated (dark blue) or not (white) with MTT are shown. Absorbance values were normalized by the mass (mg) of each slice. (N = 4–7 individual tissue donors for each time point). Data are plotted as means ± standard error. p-values for all comparisons were > 0.05 (one-way ANOVA). (B–D) Cytoarchitecture was assessed by Nissl staining at days 0, 2 and 4. Images correspond to cortical layers II/III. Scale bar = 50 μm. (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article).

![Fig. 2. Neural cell content in adult human brain slices in culture.](image)

**Fig. 2.** Neural cell content in adult human brain slices in culture. Neurons were identified by NeuN immunohistochemistry in cultured slices at day 0 (A and B), 1 (C) and 4 (D). Neuronal content along period in culture was quantified by Western Blotting for NeuN. A representative membrane is shown in E, and quantification (normalized by β-actin levels) is shown in F. (N = 3–7 independent human tissue donors for each time point). p-values for all comparisons were > 0.05 (one-way ANOVA). Astrocytes and microglia were also identiﬁed by GFAP (G) or Iba1 (H) immunostaining, respectively, at day 4. All images correspond to cortical layer V. Scale bars correspond to 100 μm (A, G, H) or 25 μm (B, C, D).
Fig. 3. Neurotransmitter release in adult human brain slices in culture. GABA and glutamate release were determined after KCl depolarization. Slices incubated in the presence of KCl (black bars) presented higher levels of both neurotransmitters compared to the basal release (indicated by the horizontal dashed line) at all time points tested. When KCl depolarization was performed in Ca++-free medium (white bars), the KCl-evoked neurotransmitter release was markedly reduced at both DIV 1 and 2. N = 3–8 individual tissue donors for each time point. *p < 0.05; **p < 0.01 (one-way ANOVA).

A major concern with the use of cortical tissue ressected from TLE cases to prepare slice cultures is the possibility that neurophysiological alterations might have spread from the hippocampal focus of epileptic activity to the neocortical region (lateral) which is the source of the cultivated tissue. The concept of epilepsy as a network disease, or the concept of focal epilepsy, with potential secondary spreading of epileptic activity to the neocortical region would be useful in the study of several neurological diseases. Since it is generally accepted that sporadic AD is linked to brain accumulation of Aβ oligomers (AβOs), which in turn leads to Tau hyperphosphorylation (De Felice and Wu, 2008; Forny-Germano and Lyra e Silva, 2014; Selkoe and Hardy, 2016), we exposed cultures to exogenous AβOs, an approach that has been widely applied to either neural cultures or animals (by icv injection) to model AD pathology (reviewed in Benilova and Karran, 2012; Walsh and Teplow, 2012; Viola and Klein, 2015). AβO binding to the tissue was measured by ELISA in extracts prepared from slices incubated with vehicle or Aβ oligomers. Robust binding of AβOs to cultured brain slices was evident (Fig. 4A). Binding was specific for Aβ oligomers, as we have used as detecting antibody the conformational scFv-type antibody NUsc1, shown to be highly specific towards AβOs (Velasco and Heffern, 2012; Sebollela and Cline, 2017). AβO binding to adult human brain slices was further detected by IHC using tissue from a different donor. Intense NUsc1-reactive labeling associated to cells was observed only in tissue exposed to Aβ oligomers (Fig. 4B-C). Interestingly, this labeling pattern resembles that observed in AD human brain sections probed with anti-Aβ oligomer antibodies (Lacor and Buniel, 2004).

Fig. 4. Toxicity of Alzheimer’s-associated Aβ oligomers on adult human brain slices has been previously attempted. Alterations in gene expression (Sebollela and Freitas-Correa, 2012) and activity. Combined with our current results showing the expected response in terms of K+ -induced release of glutamate and GABA, those previous observations suggest that cortical slices (distal to the epileptic foci) prepared from TLE patients are neurochemically normal and thus hold potential to be used as a preferred model for studying age-associated brain diseases. It is important to note, however, that the feasibility of using short-term cultures in free floating format in electrophysiological studies needs to be further investigated, and compared with the encouraging results obtained using customized media and semiperous membranes reported by Eugene and Cluzeaud (2014) and Schwarz and Hedrich (2017).

3.3. Modeling Alzheimer’s disease using cultured slices from adult human brain

Since we were able to culture adult human brain slices from different donors, and to keep them alive and synaptically active for at least 4 DIV, we asked whether this histoculture could be used as a platform to model sporadic AD in bona fide human brain tissue. A key motivation for this analysis comes from the work from Verhoog and Gorionova (2013) showing that acute (used within 15 h post-surgery) neocortical slices derived from epileptic patients presented no sign of spontaneous epileptiform activity, suggesting that slice cultures from this brain region would be useful in the study of several neurological diseases. Since we were able to culture adult human brain slices from different donors, and to keep them alive and synaptically active for at least 4 DIV, we asked whether this histoculture could be used as a platform to model sporadic AD in bona fide human brain tissue. A key motivation for this analysis comes from the work from Verhoog and Gorionova (2013) showing that acute (used within 15 h post-surgery) neocortical slices derived from epileptic patients presented no sign of spontaneous epileptiform activity, suggesting that slice cultures from this brain region would be useful in the study of several neurological diseases. Since it is generally accepted that sporadic AD is linked to brain accumulation of Aβ oligomers (AβOs), which in turn leads to Tau hyperphosphorylation (De Felice and Wu, 2008; Forny-Germano and Lyra e Silva, 2014; Selkoe and Hardy, 2016), we exposed cultures to exogenous AβOs, an approach that has been widely applied to either neural cultures or animals (by icv injection) to model AD pathology (reviewed in Benilova and Karran, 2012; Walsh and Teplow, 2012; Viola and Klein, 2015). AβO binding to the tissue was measured by ELISA in extracts prepared from slices incubated with vehicle or Aβ oligomers. Robust binding of AβOs to cultured brain slices was evident (Fig. 4A). Binding was specific for Aβ oligomers, as we have used as detecting antibody the conformational scFv-type antibody NUsc1, shown to be highly specific towards AβOs (Velasco and Heffern, 2012; Sebollela and Cline, 2017). AβO binding to adult human brain slices was further detected by IHC using tissue from a different donor. Intense NUsc1-reactive labeling associated to cells was observed only in tissue exposed to Aβ oligomers (Fig. 4B-C). Interestingly, this labeling pattern resembles that observed in AD human brain sections probed with anti-Aβ oligomer antibodies (Lacor and Buniel, 2004).

Modeling the neurotoxicity of AD-associated Aβ oligomers using adult human brain slice cultures has been previously attempted. Alterations in gene expression (Sebollela and Freitas-Correa, 2012) and
energy metabolism (Seixas da Silva and Melo, 2017) induced by AβOs in adult human brain slice cultures have been reported. In AD brains, it is though that brain accumulation of AβOs leads to hyperphosphorylation of the microtubule associated protein, Tau, which is a hallmark of AD (Selkoe, 2001; Kowalska, 2004; Iqbal and Alonso Adel, 2005). This prompted us to investigate whether exposure of cultured human brain slices to AβOs would trigger Tau hyperphosphorylation. Results showed a decreased in total Tau levels compared to β-actin in slices exposed to AβOs (Fig. 4F-G). Interestingly, pTau/Tau ratio was higher in AβO-treated slices, ranging from 142 to 225% depending on the phosphoepitope examined (Fig. 4H). It is important to point out that this increase was observed at phosphorylation sites strongly associated with AD (Iqbal and Alonso Adel, 2005). These results suggest that adult human brain slice cultures prepared as described herein are amenable to model molecular/cellular alterations in AD. Further, such cultures represent an advance over previously described cultures prepared from both fetal or postmortem human brains, in which either cell connections or the molecular machinery are quite different from that in adult live brain parenchyma, considering that fetal tissue presents cellular connections not yet fully developed (Mountcastle, 1997; Rakic, 2009; Lui and Hansen, 2011), while the later undergoes biochemical and neuroanatomical changes from death to tissue collection (Fountoulakis and Hardmeier, 2001; Lavenex and Lavenex, 2009). Studies on the feasibility of using these slice cultures to model other age-associated neurodegenerative diseases, such as Parkinson Disease, should now be pursued.

4. Conclusions

In the present work, we describe a detailed protocol for culturing adult human brain slices obtained from surgical explants in free-floating format, which renders viable and neurochemically functional slices for at least four days in culture. We have found that these slices are sensitive to Alzheimer’s disease-associated Aβ oligomers, suggesting that this adult CNS model may be a useful tool to study molecular and cellular mechanisms of neurodegenerative diseases in a more clinically relevant context.

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Appendix A. Supplementary data

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