Iron accumulation in microglia triggers a cascade of events that leads to altered metabolism and compromised function in APP/PS1 mice

RUNNING TITLE: Iron accumulation impairs microglial function

Allison McIntosh*, Virginia Mela*, Conor Harty, Aedin M Minogue, Derek A Costello, Christian Kerskens and Marina A. Lynch§
Trinity College Institute for Neuroscience,
Trinity College,
Dublin 2,
Ireland.

*Joint first authors

§Corresponding author
Email: lynchma@tcd.ie
Tel: 353-1-896-8531
Fax: 353-1-896-3545
ABSTRACT

Among the changes that typify Alzheimer’s disease (AD) are neuroinflammation and microglial activation, amyloid deposition perhaps resulting from compromised microglial function, and iron accumulation. Data from Genome Wide Association Studies (GWAS) identified a number of gene variants that endow a significant risk of developing AD and several of these encode proteins expressed in microglia, and proteins that are implicated in the immune response. This suggests that neuroinflammation and the accompanying microglial activation are likely to contribute to the pathogenesis of the disease. The trigger(s) leading to these changes remain to be identified. In this study we set out to examine the link between the inflammatory, metabolic and iron-retentive signature of microglia in vitro and in transgenic mice that overexpress of amyloid precursor protein (APP) and presenilin 1 (PS1; APP/PS1 mice), a commonly-used animal model of AD. Stimulation of cultured microglia with interferon (IFN)γ and amyloid-β (Aβ) induced an inflammatory phenotype and switched the metabolic profile and iron handling of microglia so that the cells became glycolytic and iron-retentive, and the phagocytic and chemotactic function of the cells was reduced. Analysis of APP/PS1 mice by magnetic resonance imaging (MRI) revealed genotype-related hypointense areas in the hippocampus consistent with iron deposition and immunohistochemical analysis indicated that the iron accumulated in microglia, particularly in microglia that decorated Aβ deposits. Isolated microglia prepared from APP/PS1 mice were characterised by a switch to a glycolytic and iron retentive phenotype and phagocytosis of Aβ was reduced in these cells. This evidence suggests that the switch to glycolysis in microglia may kick-start a cascade of events that ultimately leads to microglial dysfunction and Aβ accumulation.

Key words: Glycolysis, iron, microglia, amyloid-β (Aβ), neuroinflammation, phagocytosis, APP/PS1 mice.
INTRODUCTION

Optimal brain function requires iron. Iron catalyses several enzymes, is necessary for myelination, to support mitochondrial energy generation and it plays a role in cell cycling. Regulation of cell iron is therefore critical and both deficiency and iron excess have detrimental effects on brain function (33). Iron transport into and out of cells is determined by the iron import protein, divalent metal transporter1 (DMT1), and the export protein, ferroportin (9) both of which are regulated by iron regulatory protein 1 (IRP1) (39). A link between neuroinflammation and iron dysregulation in the brain has been described, particularly in Parkinson’s disease (PD). A significant body of research suggests that the pathogenesis of PD is, at least in part, driven by inappropriate microglial activation (32) and in an animal model of PD, areas of iron deposition were associated with marked microgliosis in the substantia nigra (51). Indeed increased iron loading in microglia has also been reported (47) while maximal deposition has been reported in the substantia nigra (19), where neurodegeneration is most profound. It has been proposed that increased iron may be a contributing factor in the pathogenesis of the disease (2, 38).

Inflammatory cytokines like interleukin (IL)-6 (50) and tumour necrosis factor (TNF)α (34) increase expression of DMT1 in microglia and therefore promote iron sequestration in the cells. Iron sequestration is also promoted by a downregulation of cell surface expression of ferroportin, which is regulated by the 25-amino acid liver-derived peptide, hepcidin; the evidence indicates that hepcidin binds to and internalizes ferroportin and thereby decreases iron efflux from cells (46, 48). Both IL-6 or TNFα, increase expression of hepcidin in microglia (42) and affect iron in cells by increasing influx and decrease efflux. These findings establish a link between inflammation and iron handling. Our recent findings demonstrated that IFNγ-induced changes in microglia included production of inflammatory cytokines together with iron retention, perhaps driven by an upregulation in mRNA expression of ferritin and a downregulation of ferroportin (15) and this provides further evidence that iron accumulation occurs in activated microglia.

Increased iron staining has been found in proximity to Aβ plaques in formalin-fixed tissue from individuals with Alzheimer’s disease (AD), (22) and it has been reported that iron accumulation correlates with Aβ and tau pathology (43). Iron accumulation has also been observed in AD by magnetic resonance imaging (MRI) (4) but, at present, the impact of iron accumulation in AD remains unclear.

We have recently reported that IFNγ-induced inflammatory changes and iron retention in microglia is accompanied by a metabolic switch towards glycolysis (15) and this indicated that the well-described link between inflammation and glycolysis in macrophages (12) extended to microglia.
These data are corroborated by the finding that, in immortalized microglia triggered to adopt the so-called M1 activation state, took up non-transferrin-bound iron (21).

Here we set out to determine whether exposure of microglia to IFN\(\gamma\) and A\(\beta\) in vitro and in vivo triggers a phenotype that can be identified by an inflammatory, iron retentive and metabolic signature. We report that IFN\(\gamma\) and A\(\beta\) increased glycolysis and glycolytic enzymes, including PFKFB3 in microglia, and that iron retention in cells was increased. These changes were accompanied by compromised function, specifically chemotaxis and phagocytosis. In a mouse model of AD, characterized by overexpression of amyloid precursor protein (APP) and presenilin 1 (PS1; APP/PS1 mice) and increased expression of IFN\(\gamma\) and A\(\beta\), MRI analysis revealed accumulation of iron in brain that immunohistochemical analysis showed was microglia-associated. These iron-laden microglia decorated A\(\beta\) plaques and we found that they were PFKFB3+, and therefore glycolytic. The evidence suggests that glycolytic, iron-retentive microglia lose their phagocytic capability perhaps reflecting the metabolic inefficiency of glycolysis. The possibility exists that this metabolic profile contributes to accumulation of A\(\beta\) in AD.

MATERIALS AND METHODS

Animals

Male and female transgenic APPswe/PS1dE9 mice (on a C57BL/6;C3H genetic background), littermate wild type mice and C57BL/6 mice were used in these experiments. They were sacrificed at 19-20 months of age. All experiments were performed under license from the Health Products Regulatory Authority of Ireland in accordance with EU regulations and with local ethical approval (Trinity College Dublin). Animals were housed under controlled conditions (20–22°C, food and water ad lib) and maintained under veterinary supervision.

MRI

MRI was conducted using a 7-T small animal Bruker Biospin MRI system and we used susceptibility weighted imaging to assess iron deposition (13) as this method is considered to offer a significant benefit in terms of sensitivity (27). Animals were anaesthetised (isoflurane 5% induction, 1-2% maintenance) and placed on a cradle using ear bars to stabilise the head. A built-in water pump-driven temperature regulator (SA Instruments Inc., Stony Brook, NY, USA) was used to maintain rectal temperature at 37°C ± 1°C. A mechanical ventilator (Ugo Basile, Comerio, VA, Italy) was used to deliver adequate inflowing gas to the facemask and the respiration signal was monitored using custom hardware and software (SA Instruments Inc., Stony Brook,
NY, US). Respiratory rate was maintained at 60-70 breathes per minute throughout the session. To ensure accurate positioning an initial, rapid pilot image was recorded followed by a T2 phase contrast scan triggered by respiratory rate [spatial resolution, 0.0625 x 0.0625 x 0.3 mm³; matrix, 256x256; FOV, 16x16; TR, 800ms; TE (ms), 3.09, 7.09, 11.09, 15.09, 19.09, 23.09, 27.09, 31.09; flip angle, 50°; repetitions, 1; averages, 10; trigger delay, 150ms; scan time, 25m 36s], high resolution susceptibility weighted image (SWI) for iron depositions [spatial resolution, 0.0417 x 0.0417 x 0.3 mm³; matrix, 384x384; FOV, 16x16; TR, 505.58ms; TE, 11.99ms; flip angle, 30°; repetitions, 1; averages, 6; scan time, 19m 24s], T1-weighted image [spatial resolution, 0.0625 x 0.0625 x 0.3 mm³; matrix, 256x256; FOV, 16x16; TR, 225.74ms; TE, 3.5ms; flip angle, 30°; repetitions, 1; averages, 10; scan time, 6m 28s] and a continuous arterial spin labelling sequence for qualitative cerebral perfusion [spatial resolution, 0.164 x 0.25 x 1 mm³; matrix, 128x64; FOV, 20x16; TR, 3000ms; TE, 12.95ms; flip angle, 90°; repetitions, 1; averages, 24; scan time, 2m 24s]. Areas of hypointensity in the susceptibility weighted images were quantified manually using Image J region of interest tool.

**Tissue preparation**

Mice were anaesthetised with sodium pentobarbital (Euthanimal) and transcardially perfused with saline. The brain was dissected and one hemisphere fixed in 4% paraformaldehyde for 24 h for later preparation of sections for immunohistochemical analysis. The second hemisphere was used for isolation of microglia. Following fixation, the tissue was incubated in 30% sucrose for 48 h for cryoprotection and frozen in isopentane. Tissue was mounted in optimal cutting temperature (OCT) solution and 20 μm coronal sections taken using a cryostat. Sections were stored at -80°C in freezing solution (30% ethylene glycol, 30% sucrose in PBS) until required for immunohistochemistry. Sections were stained with Iba 1 to identify microglia and for Aβ, iron and PFKFB3.

**Preparation of microglia from WT and APP/PS1 mice**

Tissue was homogenised using the gentle-MACS Dissociator (Miltenyi Biotec, UK) in combination with the AdultBrain Dissociation Kit (Miltenyi Biotec UK) according to manufacturer’s instructions. The dissociated tissue was filtered, washed and centrifuged and the pellet containing the microglia was resuspended, stained with CD11b microbeads (Miltenyi Biotec. UK) and magnetically-separated using the QuadroMACS separator (Miltenyi Biotec, UK) according to manufacturer’s instructions. Samples were resuspended and the final microglial pellet was resuspended in phosphate-buffered saline (PBS; 75 µl), cells were seeded in 48-well plates (2.5x10⁴ cells/well; final volume 200 µl) and cultured in cDMEM supplemented with M-
CSF (100 ng/ml; R&D Systems, UK) and GM-CSF (100 ng/mL; R&D Systems, UK). Microglia were allowed to acclimatise for 5 days and media was changed every second day. On day 5, cells were fixed with 4% paraformaldehyde (30 min) and stored in PBS at 4°C until use. Some coverslips were incubated on day 5 with Aβ1-40 (5 μM) for 2 h prior to fixation to investigate phagocytosis.

**Preparation of microglia from neonatal mice**

Microglia were prepared as previously described (6). Briefly, isolated mixed glia from cortical tissue of C57BL/6 neonatal mice were cultured in T25 cm² flasks in cDMEM supplemented with macrophage colony stimulating factor (M-CSF; 100 ng/ml; R&D Systems, UK) and granulocyte macrophage colony stimulating factor (GM-CSF; 100 ng/ml; R&D Systems, UK) for 10-12 days, after which time non-adherent microglia were seeded in 24-well plates (1x10⁵ cells/well) and cultured for a further 2 days. Medium was replaced with fresh cDMEM ± IFNγ (50 ng/ml; Enzo Life Sciences, UK) and amyloid-β (Aβ1; 10 μM comprising Aβ1-40 (4.8 μM) + Aβ1-42 (5.8 μM)) for 24 h. We used this Aβ combination because cells are exposed to both peptides in vivo and previous experiments indicated that it triggered inflammatory changes (20) and synergised with LPS to trigger neuroinflammatory and metabolic changes (36). To prepare Aβ, lyophilized Aβ1-40 and Aβ1-42 peptides were dissolved in HPLC grade water to provide a 6 mg/ml stock solution, which was diluted to 1mg/ml using sterile PBS and allowed to aggregate (24 h, 220 rpm, 37°C). This preparation contains both oligomers and fibrils as evidenced by enhanced Thioflavin T binding (data not shown). This In separate experiments, to assess phagocytosis and chemotaxis, cells were seeded in 48-well plates (6x10⁴ cells/well) and cultured for 24 h. Medium was replaced with fresh cDMEM ± IFNγ (50 ng/ml; Enzo Life Sciences, UK) and amyloid-β (Aβ1-40 (4.2 μM) and Aβ1-42 (10 μM)) for 24 h and harvested for analysis as described below. In separate experiments, cells were incubated in the presence or absence of FeCl₃ (100μM; Sigma, UK) for 6 h to assess the effect of iron loading on glycolysis as assessed by PFKFB3.

**PCR**

mRNA expression of TNFα and iNOS was assessed in microglia from neonatal mice by RT-PCR. RNA was isolated from microglia using the Nucleospin® RNAII KIT (Macherey-Nagel, Duren, Germany) and cDNA was prepared using High-Capacity cDNA RT kit according to the manufacturer’s instructions (Applied Biosystems, UK). Real-time PCR was performed with predesigned Taqman gene expression assays (TNFα (Mm0043258_m1; NOS2 (Mm00440502_m1; Applied Biosystems, UK)) using an Applied Biosystems 7500 Fast Real-Time PCR machine (Applied Biosystems, Germany). Samples were assayed as previously
described (7) with β-actin as the endogenous control to normalize gene expression data. Gene expression was calculated relative to the endogenous control samples and to the control sample giving an RQ value (2\(^{-\Delta\Delta CT}\), where CT is the threshold cycle).

**Western immunoblotting**

Microglia, or samples of hippocampal homogenate in the case of IFN\(\gamma\) treatment, were prepared for western immunoblotting as previously described (8). Briefly, cells were incubated in lysis buffer (composition in mM: Tris-HCl 10, NaCl 50, Na\(_2\)PO\(_4\)-H\(_2\)O 10, NaF 50, containing 1% each of Igepal, phosphatase inhibitor cocktail I and II, and protease inhibitor; Sigma, UK), equalized for protein, added to 4x SDS sample buffer (composition: Tris-HCl 100mM, pH 6.8, 4% SDS, 2% bromophenol blue, 20% glycerol; Sigma, UK), boiled (95°C, 5 min) and applied to 8-12% SDS gels. Proteins were transferred to nitrocellulose membrane, non-specific binding was blocked (5% Marvel in TBS containing 0.05% Tween 20) and membranes were incubated overnight at 4°C with the antibodies raised against ferritin (1:1000), PFKFB1 (1:400), PFKFB3 (1:1000), hexokinase II (1:500), PKM2 (1:1000) (AbCam, US), HIF-1α (1:1000; Novus Biologics) or IFN\(\gamma\) (1:1000; R&D Systems, UK) (all IgG, raised in rabbit, diluted in 5% non-fat dried milk/TBS-T). Membranes were washed and incubated (room temperature, 2 h) with a secondary HRP-linked anti-rabbit antibody (1:5000 in 5% milk in TBS-T). Immunoreactive bands were detected using WesternBright ECL chemiluminescent substrate (Advansta, US). Images were captured using the Fujifilm LAS-4000 imager and densitometry analysis was carried out using ImageJ (http://rsb.info.nih.gov/).

**Metabolic analysis**

The SeaHorse Extracellular Flux (XF24) Analyser (SeaHorse Bioscience, US) was used to carry out real-time metabolic analysis of cells. Microglia (1x10\(^5\) cells/well) were seeded (100 μl/well) on SeaHorse cell culture microplates and treated with IFN\(\gamma\) (50ng/ml) +Aβ (10μM) for 24 h as described above. The sensor cartridge was hydrated by adding SeaHorse XF Calibrant solution (1 ml; SeaHorse BioScience, US) to each well of the utility plate and left overnight in a CO\(_2\)-free incubator at 37°C. Prior to the assay, cells were washed twice with the appropriate assay medium (1 ml) according to the manufacturer's instructions, and assay media added to give a final volume of 500 μl/well and the plate was incubated in a CO\(_2\)-free incubator (37°C, 1 h). For the mitochondrial stress test, oligomycin (20 μM; AbCam, UK), carbonyl cyanide-4-(trifluoromethoxy)phenylhydrazone (20 μM; FCCP; Sigma-Aldrich, UK) and antimycin A (40 μM; Sigma-Aldrich, UK) were loaded into the appropriate ports for sequential delivery. For the glycolytic flux test, glucose (10 mM), oligomycin (20 μM) and 2-deoxy-D-glucose (2-DG; 500
mM; all Sigma-Aldrich, UK) were prepared in glycolytic flux assay media and similarly loaded into the appropriate ports. Following calibration, oxygen consumption rate (OCR) and extracellular acidification rate (ECAR) were measured every 8 min for 96 min during which time the appropriate compounds were injected sequentially at 24 min intervals. ECAR and OCR were automatically calculated using the SeaHorse XF24 software and 4-6 replicates were assessed for each separate sample.

**Iron accumulation in microglia from neonatal mice**

Iron accumulation was assessed in microglia from neonatal mice that were incubated in the presence or absence of IFNγ+Aβ. Cells were then washed and fixed (4% PFA, 30 mins) for subsequent iron staining. The Prussian blue reaction for iron staining was carried out using the Hematognost Fe® kit (Merck Millipore, Germany). Briefly, after washing with PBS, cells were incubated in the staining solution (1:1 ratio of 4.78% potassium hexacyanoferrate (II) and 5% hydrochloric acid) for 20 min at 37°C, washed with distilled H₂O and counterstained with nuclear fast red solution (0.1%, 5 min). Coverslips were then dehydrated through graded alcohol and immersed in 100% xylene (VWR International UK) before mounting onto slides using DPX (RL Lamb, UK). Images were taken using an Olympus IX51 light microscope with a built-in camera (Olympus, Japan). Ten fields of view (FOV) were averaged per replicate resulting in a total of 12381 cells analysed. Data are presented as the proportion of iron-positive cells (calculated by ‘number of iron+ cells/total number of cells’ per FOV).

**Analysis of ROS**

ROS generation was assessed in primary microglia treated with IFNγ+Aβ using CellRox Deep Red (Thermofisher) as per the manufacturer’s instructions. Briefly, CellRox Deep Red solution was added to the media following treatment and incubated in the dark (37°C, 5% CO₂, 30 min). Cells were washed and fixed (3.7% paraformaldehyde, 15 min) and fluorescence assessed in a fluorescent plate reader, and quantified.

**Analysis of phagocytosis and chemotaxis**

Aβ engulfment was assessed in microglia that were plated onto coverslips coated with poly-D lysine (5 µg/ml; Merck Millipore Ltd, UK). Cells were either isolated from WT and APP/PS1 mice, or from neonatal mice and microglia from neonates were incubated ± IFNγ +Aβ for 24 h as described above. Cells were fixed in 4% PFA and washed in PBT (PBS+1% Triton X-100) prior to staining. Microglia were incubated first in PBT containing 3% BSA (1 h) to block nonspecific binding and then overnight at 4°C in the presence of rabbit anti-Iba1 (1:1000; Wako, Japan) and mouse anti-Aβ (1:500; Biolegend, US). Coverslips were washed and incubated (2 h,
room temperature) with Alexa Fluor 546 donkey anti-rabbit IgG (1:1000) and Alexa Fluor 488 donkey anti-mouse IgG (1:1000), and mounted in ProLong Gold with the nuclear marker DAPI (Thermo Scientific, US). Images (8 fields/animal; 40X magnification) were acquired with a Zeiss AX10 Imager A1 microscope. Analysis of images was undertaken with ImageJ software, the number of Aβ+ microglia was assessed and the % of Iba+ cells that engulfed Aβ was calculated.

Chemotaxis was assessed in microglia from neonatal mice. Cells were plated (115,000 cells/cm²) onto the transwell insert on a Boyden chamber (pore size 8µm; Corning Inc, UK), incubated ± IFNγ+Aβ for 24 h as described above. Fresh media ± ATP (300 µM; Sigma-Aldrich, Ireland) was added to the wells in the Boyden chamber and cells were incubated for 6 h at 37°C. Non-migrating cells from the upper side of the cell culture insert were removed and migrating cells that remained on the membrane were fixed (4% PFA, 30 min), washed 3 times and mounted in ProLong Gold with the nuclear marker DAPI (Thermo Scientific, US). Images (8 fields; 20X magnification) were acquired with a Zeiss AX10 Imager A1 microscope. Analysis of images was undertaken with ImageJ software, the number of migrated microglia was assessed and chemotaxis percentage was calculated using spontaneous migration (ie the number of cells that migrated in the absence of ATP) as control.

**Analysis of Aβ**

Hippocampal tissue from WT and APP/PS1 mice was homogenised in sodium dodecyl sulphate (SDS) buffer (1% SDS, 50mM NaCl in dH2O) containing protease and inhibitors (1%; Sigma-Aldrich, UK), centrifuged (21,500 x g, 20 min, 4°C) and SDS-soluble Aβ was assessed in the supernatants. To assess insoluble Aβ, pellets were incubated in guanidine buffer (50mM Tris-HCl, 5M guanidine in dH2O) to provide sample for analysis. Aβ concentrations were determined using a MULTI-SPOT® Human/Rodent (4G8) Abeta Triplex Ultra-Sensitive Assay (Meso Scale Discovery, USA) as per the manufacturer’s instructions and as described previously (26).

**Immunohistochemistry**

Amyloid plaques were visualised by incubation of sections in saturated NaCl (80% ethanol in dH2O saturated with NaCl and supplemented with 1M NaOH) for 20 min at room temperature. Sections were incubated in filtered Congo red solution (200 ml; 0.2% Congo red dye in saturated NaCl solution, 30 min) and rinsed in dH2O.

Microglia were visualized in tissue by staining for Iba1. Tissue was washed blocked for endogenous peroxidase by incubating in hydrogen peroxide (0.75%; 20 min) and non-specific binding was blocked by incubating in normal horse serum (10%; 1 h) before incubation overnight in rabbit anti-Iba1 primary antibody (WAKO, Japan; 1:5000; at 4°C). Sections were incubated in
goat anti-rabbit biotinylated secondary antibody (Vector Labs, UK; 1:300; 90 min) and processed using the avidin-biotin immunoperoxidase method (Vectastain ABC Elite kit, Vector Laboratories, UK). Iba1 immunoreactivity was visualized by the addition of the chromogen 3’,3'-diaminobenzidine (containing 0.02% H2O2 in PBS, Sigma-Aldrich, UK). Sections were mounted and allowed to dry overnight.

Iron staining was performed using the Prussian blue reaction (Hematognost Fe, Millipore Ireland) by incubating the sections (1:1 potassium hexacyanoferrate (II; 4.78%) and HCl (4%); 20 min, 37°C). Slides were washed and dehydrated through graded alcohols and immersed in 100% xylene (VWR International UK) before mounting with DPX (Sigma) and viewed on an Olympus IX51 light microscope with a built-in camera (Olympus, Japan). The proportion of Iba1+ cells containing iron was calculated by dividing the number of Iron+ Iba1+ cells by the total number of Iba1+ cells per field of view.

PKFFB3 was assessed in floating sections which were first incubated with citrate (20 min, 85°C) and pepsin (20 min, room temperature) to promote antigen retrieval. Tissue was washed, permeabilised (10 min; PBS with 0.1% Triton-X100), blocked (1 h; 5% BSA with 0.1% Triton-X100) and incubated (overnight, 4°C) with primary antibodies (goat anti-Iba1; 1:1000 (LSBio Inc., US), rabbit anti-PKFFB3; 1:250 (Abcam, UK) and mouse anti-Aβ; 1:750 (Biolegend, US). Samples were washed and incubated (2 h, room temperature) with Alexa Fluor® 546 donkey anti-rabbit IgG (1:1000), Alexa Fluor® 633 donkey anti-goat IgG (1:1000) and Alexa Fluor® 488 donkey anti-mouse IgG (1:1000) and mounted in ProLong®Gold with the nuclear marker DAPI (ThermoScientific, US).

**Analysis of PKFFB3 in isolated microglia**

Microglia, isolated from one hemisphere of WT and APP/PSEN1 mice, were cultured as described above and, on day 5, were washed, permeabilised (10 min; PBS with 0.1% Triton-X100), blocked (1 h; 3% BSA with 0.1% Triton-X100) and incubated (overnight, 4°C in primary antibodies; rabbit anti-PKFFB3 (1:250; Abcam, UK) and goat anti-Iba1 (1:100; LSBio Inc., US)). Coverslips were incubated (2 h, room temperature) with Alexa Fluor® 546 donkey anti-goat IgG (1:1000) and Alexa Fluor® 488 donkey anti-rabbit IgG (1:1000) and mounted in ProLong®Gold with the nuclear marker DAPI (ThermoScientific, US). Images were taken using a Leica SP8 scanning confocal microscope (40X; 5 fields of view) and processed using Image J software. To analyse the localisation of PKFFB3 staining, DAPI regions of interest were generated and subtracted from total cellular fluorescence. These values were corrected for background fluorescence by
calculating the corrected total cellular fluorescence (CTCF) = integrated density - (area of selected cell x mean fluorescence of background readings).

**Statistical analysis**
Data are reported as the mean ± SEM and the number of experiments is indicated in each case. Statistical analysis was carried out using Student’s t-test for independent means. The significance level was set at p < 0.05.

**RESULTS**

**IFN\(\gamma\)+A\(\beta\) increases glycolysis in microglia from neonatal mice**
We have reported that inflammatory microglia become glycolytic (15) and to confirm this we assessed the metabolic profile of microglia under control conditions and following stimulation with IFN\(\gamma\)+A\(\beta\). We show that IFN\(\gamma\)+A\(\beta\) increased mRNA expression of TNF\(\alpha\) and iNOS (*p < 0.05; **p < 0.01; Student’s t-test for independent means; Figure 1A,B). These changes were accompanied by increased glycolysis. Thus ECAR was markedly increased in microglia incubated with IFN\(\gamma\)+A\(\beta\) compared with control (Figure 1C) and IFN\(\gamma\)+A\(\beta\) significantly increased mean glycolysis and mean glycolytic capacity (**p < 0.01; Student’s t-test for independent means; Figure 1D,E). Analysis of oxidative metabolism indicated that there was no IFN\(\gamma\)+A\(\beta\)-induced change in OCR, basal or maximal respiration (Figure 1F-H) and neither ATP production nor proton leak were altered (data not shown). To consolidate these findings, we examined expression of glycolytic enzymes. Hexokinase II and PKM2 catalyze the first and last steps in glycolysis and both are rate-limiting, and PFKFB3 is a potent activator of glycolysis because it controls production of fructose-2,6-bisphosphate which, in turn, controls the activity of phosphofructokinase (PFK)-1, a key glycolytic enzyme. Here we show that expression of these enzymes was increased in IFN\(\gamma\)+A\(\beta\)-stimulated microglia (*p < 0.05; **p < 0.01; ***p < 0.001; Student’s t-test for independent means; Figure 2A-D). Among the factors that activate glycolytic enzymes, including PFKFB3 (25) is HIF-1\(\alpha\), expression of which is modulated by ROS. Therefore we investigated both and found that HIF-1\(\alpha\) and ROS, assessed using the fluorescent probe, CellRox Deep Red Reagent, was significantly increased in IFN\(\gamma\)+A\(\beta\)-stimulated, compared with control, microglia (*p < 0.05; **p < 0.01; Student’s t-test for independent means; Figure 2E,F).

**IFN\(\gamma\)+A\(\beta\) increases iron accumulation and impairs function in microglia from neonatal mice**
Ferritin is the main iron storage protein and here we show that it was increased in IFN\(\gamma\)+A\(\beta\)-stimulated microglia, as indicated by the sample immunoblot and the mean densitometric data (**p
< 0.01; Student’s t-test for independent means; Figure 3A), and was accompanied by a significant IFNγ+Aβ-induced increase in the number of Iba1+ cells that stained positively for iron (**p < 0.01; Student’s t-test for independent means; Figure 3B,C). Together the data indicate that IFNγ+Aβ-stimulated microglia are inflammatory, glycolytic and iron retentive and produce increased ROS. The data suggest that iron loading might trigger glycolysis as indicated by the finding that incubation of microglia in the presence of FeCl₃ significantly increased PFKFB3 (*p < 0.05; student’s t-test for independent means; Figure 3D,E). To determine the impact of iron loading and glycolysis on microglial function, we evaluated phagocytosis and chemotaxis. Almost 80% of Iba1+ cells were Aβ+ in the control samples whereas this was markedly reduced in IFNγ+Aβ-treated samples (***p < 0.01; Student’s t-test for independent means; Figure 4A,B). Chemotaxis was also significantly reduced in cells incubated with IFNγ+Aβ (**p < 0.01; Student’s t-test for independent means; Figure 4C).

Iron accumulation is increased in microglia from APP/PS1 mice and iron-laden microglia cluster around Aβ-containing plaques

We asked whether these changes might also occur in vivo since IFNγ and Aβ were increased in brain tissue from APP/PS1 mice (3) and therefore first confirmed that these changes were reproducible; the data show a genotype-related increase in hippocampal IFNγ (Figure 5A), as well as soluble Aβ₁₋₄₀ and Aβ₁₋₄₂ (*p < 0.05; ***p < 0.001; student’s t-test for independent means; Figure 5B,C). Insoluble Aβ concentrations were also increased ((284.3 ng/mg (± 0.96, SEM) vs 10739 (2893) and 8.41 ng/mg (±0.19 SEM) vs 572.9 (164.0) in WT and APP/PS1 mice for Aβ₁₋₄₀ and Aβ₁₋₄₂ respectively; p< 0.05 in each case; data not shown)). We argued that the well-described inflammatory microglia that have been consistently reported in APP/PS1 mice (17, 26) would be accompanied by iron accumulation in the cells and, to investigate this in our model, we first assessed anaesthetized mice using MRI. In the hippocampus of APP/PS1 mice, there were areas of hypointensity on susceptibility-weighted images that are indicative of iron accumulation but there was little evidence of hypointensity in wildtype mice (Figure 6A). Analysis of the mean data revealed a significant genotype-related increase in the area of hypointensity and also a significant increase in average pixel intensity in these areas (**p < 0.01; student’s t-test for independent means; Figure 6B,C). To determine whether the increased iron retention was specific to microglia, tissue was stained for Iba1 and iron and, to assess whether these microglia were related to areas of Aβ accumulation, colocalization of Iba1, iron and Aβ was examined. The sample images indicate that there were several Aβ-containing plaques in hippocampal tissue prepared from APP/PS1 mice (Figure 7A), that these plaques were decorated
with iron-laden microglia (Figure 7B) and that the proportion of iron+ Iba+ cells was significantly greater in sections from APP/PS1 mice compared with WT mice (**p < 0.01; student’s t-test for independent means; Figure 7C). Further analysis indicated that the microglia surrounding the plaques were PFKFB3-positive indicating that they were glycolytic (Figure 7D). **Microglia from APP/PS1 mice are glycolytic and less phagocytic than cell from WT mice**

To confirm this, microglia were isolated from APP/PS1 mice and WT mice and stained for PFKFB3 and the data show that there was a genotype-related increase in PFKFB3 immunostaining (Figure 8A) and that mean cytosolic PFKFB3 (Figure 8B), but not nuclear PFKFB3 (Figure 8C), was significantly increased in cells from APP/PS1 mice (**p < 0.01; student’s t-test for independent means); a significant genotype-related increase in total PFKFB3 was also observed (**p < 0.01; student’s t-test for independent means; Figure 8D). **The localization of PFKFB3 to the cytosol is relevant since this is the site of glycolysis.** Consistent with the data generated in IFNγ+Aβ-treated cells, we found that microglia from APP/PS1 mice, which we show are glycolytic and iron-retentive, had a reduced capacity to engulf Aβ (Figure 9A) and therefore the number of 6e10+ Iba1+ cells was significantly decreased in cells isolated from APP/PS1 mice compared with WT mice (**p < 0.01; student’s t-test for independent means; Figure 9B).

**DISCUSSION**

We set out to determine whether microglia that are exposed to inflammatory stimuli can be identified by an inflammatory, metabolic and iron retentive signature and to determine the impact of these changes on cell function. IFNγ and Aβ increased expression of inflammatory markers TNFα and iNOS, triggered a switch towards glycolysis and stimulated iron accumulation in microglia from neonatal mice. Likewise, microglia from APP/PS1 mice, in which inflammatory changes have been consistently described, exhibited a glycolytic and iron-retentive phenotype. The evidence indicates that these cells have a reduced ability to phagocytose Aβ, perhaps because of the switch towards the metabolically-inefficient glycolysis.

In macrophages, the accepted view is that inflammatory stimuli switch the cells towards a glycolytic phenotype (12, 24), and this rapid change is considered to support the immune function of these cells (24). Here we show that IFNγ+Aβ predictably increased TNFα and iNOS in microglia and that this was paralleled by a switch towards a glycolytic phenotype indicating that glycolysis is increased in inflammatory microglia as it is in inflammatory macrophages. IFNγ alone is also sufficient to induce similar responses in microglia (15) and in BV2 cells (31, 44) but our data indicate that Aβ alone did not significantly increase glycolysis, but in combination
with LPS, it induced a significant increase (36). In the present study, despite the change in glycolysis, there was no evidence of a change in oxidative metabolism in IFNγ+Aβ-treated microglia.

Consistent with its impact on glycolysis in microglia, IFNγ+Aβ increased expression of hexokinase II and PKM2, the enzymes that catalyse the first and last reactions in glycolysis, and IFNγ+Aβ also increased PFKFB3. PFKFB3 results in the production of fructose 2,6-bisphosphate, which is an allosteric activator of phosphofructokinase (35), the enzyme responsible for the conversion of fructose-6-phosphate to fructose-1, 6-bisphosphate which is a rate-limiting, irreversible step in glycolysis. This is therefore a control point in the pathway and PFKFB3 is an acknowledged key regulator of glycolysis. Among the factors that trigger activation of PFKFB3 are inflammatory mediators; for example IL-6 increased PFKFB3 mRNA in mouse embryonic fibroblasts (1) and cancer cell lines (14) and several studies have indicated that LPS triggers PFKFB3 in macrophages (37, 40). Few studies have investigated factors that drive PFKFB3 in microglia but we have reported that IFNγ alone (15) and also LPS+Aβ (36), exert this effect. HIF-1α is recognized as a potent activator of PFKFB3 and this has been described in fibroblasts (25, 30) and macrophages (40). In turn, HIF-1α is regulated by reactive oxygen species (28, 29). It has been suggested that, in macrophages, the increase in glycolysis is necessary to facilitate maintenance of the inflammatory response and that this, at least in some circumstances, is primarily dependent on increased glucose uptake, increased hexokinase activity and increased PFKFB3 triggered by HIF-1α (18). Indeed recent findings revealed that the LPS-induced inflammatory and metabolic response in macrophages are dependent on HIF-1α and also demonstrated that inflammatory and metabolic response induced in macrophages by infection of mice with *L. monocytogenes* or *C. albicans* was compromised in HIF-1α−/− mice (18). Here we show that IFNγ+Aβ increases both HIF-1α and reactive oxygen species production providing indirect evidence that these factors play a role in modulating PFKFB3 in microglia as they do in other cells.

An important finding in macrophages is that increased glycolysis is associated with increased efferocytosis of infected cells which was accompanied by increased uptake of a glucose analogue, 2-((N-(7-nitrobenz-2-oxa-1,3-diazol-4-yl)amino)-2-deoxyglucose suggesting that an increased supply of glucose to drive PRKRB3-driven glycolysis supports efferocytosis (16). The present findings suggest that IFNγ+Aβ-induced glycolytic microglia have a reduced ability to engulf Aβ suggesting that phagocytosis, at least of Aβ, is curtailed when cells utilize this less efficient means of producing ATP. Interestingly the decrease in phagocytosis was also associated with a decrease
Iron distribution across cells is determined by a number of transport and storage proteins including ferritin, the main storage protein, and ferroportin, the protein principally responsible for exporting iron from cells. IFN$\gamma$+A$\beta$ increased ferritin in microglia while ferroportin was decreased, and consequently iron accumulated in cells. Therefore consistent with our previous results (15), we show that iron accumulation is a characteristic of glycolytic microglia. Immortalized microglial cells share this characteristic; it was shown in these cells that LPS increased non transferrin-bound iron uptake and expression of ferritin and divalent metal transporter-1, while LPS-stimulated cells also exhibited an increase in glycolysis (21). Thus although different inflammatory stimuli share the ability to trigger glycolysis in primary and immortalized microglial cells, LPS decreased oxidative metabolism in immortalized microglia (21), whereas IFN$\gamma$+A$\beta$ exerted no effect in primary microglia. The finding that FeCl$_3$ increased PFKFB3 suggests that iron accumulation drives glycolysis and this supports our earlier data (15).

The present data and our previous findings suggest that increased glycolysis in microglia, irrespectively of the stimulus, is tightly coupled to increased PFKFB3 (15, 36). Therefore as a proxy measure of glycolysis, we assessed PFKFB3 in tissue sections and isolated microglia prepared from APP/PS1 mice, in which IFN$\gamma$ and A$\beta$ are increased in vivo, compared with WT mice. The data show that PFKFB3 immunoreactivity was markedly increased in both preparations. Furthermore, analysis in hippocampal sections, which predictably showed A$\beta^+$ accumulations in APP/PS1 mice compared with WT mice, revealed that these plaques were surrounded by Iba1$^+$ microglia that stained positively for PFKFB3. There was little, if any staining for A$\beta$ in these cells perhaps suggesting that the cells have a reduced phagocytic capacity. This was confirmed when we evaluated A$\beta$ uptake in these microglia; a marked reduction in A$\beta$ staining was observed in microglia from APP/PS1 mice compared with WT mice. Therefore the coupling between PFKFB3 and glycolysis that we observed in vitro is also evident in vivo confirming previous (15). It is well documented that A$\beta$ plaques are more prevalent in female than in male APP/PS1 mice (10, 45) and, similar sex-related changes in inflammatory changes and behaviour are also observed, at least in 9-10 month-old mice (10, 11). Here, we used males and females and have not systematically assessed whether there are sex-related differences in metabolic changes.

To determine whether these changes in microglia of APP/PS1 mice were accompanied by iron accumulation, we first assessed the brain by MRI and the analysis revealed areas of hypointensity
in hippocampus that are indicative of iron accumulation. Subsequent histological analysis indicated that microglia from APP/PS1 mice were iron-loaded compared with microglia from wildtype mice. Furthermore we found that iron-laden microglia decorated Aβ deposits in APP/PS1 mice and therefore the evidence indicates that the microglia which surround Aβ plaques have a glycolytic and iron retentive phenotype and appear to have reduced phagocytic activity. A number of studies have reported that iron accumulation is a characteristic of AD and MRI analysis of tissue slabs from fixed human AD, compared with control, tissue revealed hypointensities in the subiculum extending into the hippocampus and the authors reported that the iron deposition was largely found in microglia (49). Similar areas of hypointensity were identified in slices of entorhinal cortex obtained from AD patients and these were significantly greater in areas Aβ accumulation compared with the surrounding tissue but persisted to some extent even after tissue was treated with the iron chelator deferoxamine (23). A more recent study consolidated these observations and reported that there was accumulation of iron in microglia in the frontal cortex of AD patients where the cells were often localized around Aβ plaques (43). Ferritin-labelled cells have also been found in close association with Aβ plaques in tissue from AD patients and, although these cells are also observed in APP/PS1 mice, they were not so tightly associated with plaques, highlighting an inter-species difference in distribution (22). The work of several researchers support the contention that accumulation of metal ions, including iron, is a trigger for Aβ accumulation (5) and the recent observation that iron-bearing fibrillar aggregates, as well as iron-laden plaques, are evident in the brain of APP/PS1 mice, support this view, particularly since a direct correlation between fibril density and iron was found (41).

The present data, together with our previous findings that showed FeCl₃ could increase glycolysis, suggest that iron accumulation by microglia can trigger a cascade of events, acting as a switch that pushes microglia into an inflammatory and glycolytic phenotype. The data obtained in microglia from APP/PS1 mice confirm the correlation between inflammation, iron accumulation and glycolysis and lead us to propose that driving microglia to utilize the more metabolically-inefficient glycolysis limits the ability of microglia to phagocytose Aβ. It remains to be determined whether this sequence of events contribute to the accumulation of Aβ that characterizes AD and if correcting the metabolic shift can impair progression of the disease.

**Acknowledgements:** This work was supported by PI grant to MAL from The Science Foundation Ireland (15/iA/3052), to which we are very grateful.
Declarations: The authors declare no conflict of interest

REFERENCES


FIGURE LEGENDS

Figure 1. IFNγ +Aβ increased glycolysis in microglia.
Microglia isolated from neonatal brains were cultured as described in the Methods and incubated with/without IFNγ (50ng/ml) + Aβ (10 μM) for 24h. Cells from 8 separate experiments were harvested for analysis by PCR to assess mRNA expression of TNFα and iNOS, or for metabolic
analysis using the SeaHorse Extracellular Flux (XF24) Analyser. IFNγ +Aβ increased TNFα and iNOS mRNA (*p < 0.05; **p < 0.01; ***p < 0.001; student’s t-test for independent means; Figure 1A,B; n = 8). Microglia (1x10^5 cells/well) were seeded in a Seahorse XF plate and treated with IFNγ (50ng/ml) +Aβ (10μM) for 24h. To assess ECAR, 3 baseline measures were taken and the cells were sequentially exposed to glucose (10mM), oligomycin (20μM) and 2-deoxy-D-glucose (2-DG; 500mM); the resultant bioenergetic profile is presented (C). IFNγ +Aβ significantly increased glycolysis and glycolytic capacity (D, E; **p < 0.01; student’s t-test for independent means; n=6-8) Data are presented as means ± SEM (n = 6-8). To assess OCR, 3 baseline measures were taken and subsequent measurements were taken after sequential incubation of cells in oligomycin (20μM), carbonyl cyanide-4-(trifluoromethoxy)phenylhydrazone (20μM; FCCP) and antimycin A (40μM). The OCR bioenergetics profile and analysis of separate measure indicated that IFNγ +Aβ had no significant effect on OCR (F) basal respiration (G) or maximum respiration (H). Data are presented as means ± SEM (n = 3).

Figure 2. IFNγ +Aβ increased glycolytic enzymes, as well as HIF-1α and ROS production, in microglia.

Microglia from neonatal brains, prepared as described in the Methods, were incubated with/without IFNγ (50ng/ml) + Aβ (10 μM) for 24h after which time cells were harvested, lysed, equalized for protein, separated by gel electrophoresis and transferred to nitrocellulose membrane for analysis. IFNγ +Aβ significantly increased hexokinase II (A), PKM2 (B) and inducible and constituitive PFKFB3 (C,D; **p < 0.01; student’s t-test for independent means). In addition, IFNγ+Aβ increased HIF-1α (E; *p < -0.05) and analysis of ROS by CellRox Deep Red revealed a significant IFNγ+Aβ-induced increase in ROS (F; **p < 0.01). Sample immunoblots for hexokinase II, PKM2 and PFKFB3 are shown. Data are presented as means ± SEM (n = 6).

Figure 3. IFNγ +Aβ increased iron accumulation in microglia

Primary microglia were cultured from neonatal mice as described in the Methods, incubated with/without IFNγ (50ng/ml) + Aβ (10 μM) for 24h, and harvested for analysis by western immunoblotting or for staining. A. IFNγ+Aβ significantly increased ferritin (**p < 0.01; student’s t-test for independent means; n = 6). B.C. IFNγ+Aβ markedly increased the number of cells that stained positively for iron (B, indicated by arrows) and analysis of the data indicated that the percentage of iron+ cells was significantly greater in IFNγ+Aβ-incubated cells compared with controls (**p < 0.01; student’s t-test for independent means; n = 6). D.E. Incubation of microglia in the presence of FeCl3 significantly increased PFKFB3 (*p < 0.05; student’s t-test for independent means; n = 4).

Data are presented as means ± SEM. 

Figure 4. IFNγ +Aβ decreased phagocytosis and decreased chemotaxis in microglia from neonatal mice.

Primary microglia were cultured from neonatal mice as described in the Methods, incubated with/without IFNγ (50ng/ml) + Aβ1–40/1–42 (5/10μM) for 24 h, and assessed for phagocytosis of Aβ and chemotaxis. A. The representative image shows that IFNγ+Aβ reduced the number of cells that engulfed Aβ and the mean data indicated that IFNγ +Aβ significantly decreased the percentage of 6e10+ Iba1+ microglia (B) and also significantly decreased chemotaxis, as assessed by counting migrating cells in response to ATP (C; *p < 0.05; ***p < 0.01; student’s t-test for independent means). Data are the means ± SEM for 8 estimates in 5 separate experiments.
Figure 5. Increased IFNγ and Aβ in hippocampus from APP/PS1 mice.
Hippocampal tissue prepared from WT and APP/PS1 mice was assessed for IFNγ and soluble Aβ as described in the Methods. IFNγ (A) and Aβ1-40 and Aβ1-42 (B) significantly increased in hippocampal tissue from APP/PS1 mice compared with WT mice (*p < 0.05; **p < 0.001; student’s t-test for independent means; n=5 and 7 for WT and APP/PS1 mice respectively).

Figure 6. MRI analysis reveals iron accumulation in hippocampus from APP/PS1 mice.
Iron accumulation was analysed in hippocampus as described in the Methods and areas of hypointensity in the susceptibility weighted images were quantified. A Areas of hypointensity that are indicative of iron accumulation were identified in hippocampus of APP/PS1 mice but there was little evidence of hypointensity in WT mice. B,C. Analysis of the mean data revealed significant genotype-related increases in the area of hypointensity and in average pixel intensity in these areas (C; **p < 0.01; student’s t-test for independent means; n=10 and 9 for WT and APP/PS1 mice respectively).

Figure 7. Iron+ PFKFB3+ Iba1+ microglia surround plaques in hippocampal sections from APP/PS1 mice.
Sections of hippocampus (20 μm) were assessed for Aβ, iron and PFKFB3 as described in the Methods. A. Several Aβ-containing plaques, that stain pink, were identified in sections prepared from APP/PS1 mice with none evident in sections from WT mice. B. Ramified Iba+ microglia, which are stained as light brown cells, are evident in sections prepared from WT mice. In contrast, the Iba+ microglia in sections from APP/PS1 mice are generally less ramified, and are darkly-stained due to the iron deposition (dark brown/black), for example in the inset. The Aβ deposits (shown in pink in the section from APP/PS1 mice and marked by arrows) are surrounded by darkly-stained, iron-laden Iba+ microglia. There was no evidence of iron staining in Iba+ microglia in WT mice and some lighter-stained cells, that are distant from the Aβ plaques are present in the section from APP/PS1 mice. A significant increase in iron+ Iba+ cells was observed in sections from APP/PS1, compared with WT, mice (C; **p < 0.01; student’s t-test for independent means; n=6 and 8 for WT and APP/PS1 mice respectively). D. The evidence indicates that the Iba+ cells (red) that surrounded the Aβ plaques (bright green) were PFKFB3+ (yellow) suggesting that these cells are glycolytic. The arrows indicate 2 sample cells close to the Aβ plaques, shown in the insert, that are Iba+ PFKFB3+.

Figure 8. Microglia from APP/PS1 mice exhibit increased PFKFB3.
Isolated microglia from wildtype and APP/PS1 mice, were assessed for PFKFB3+ staining as described in the Methods. A. A marked increase in PFKFB3 staining was observed in isolated microglia from APP/PS1, compared with WT, mice. B-D. Cytoplasmic PFKFB3 (B), but not nuclear PFKFB3 (C), was significantly increased in cells from APP/PS1 mice (**p < 0.01; student’s t-test for independent means) and a significant genotype-related increase in total PFKFB3 was also observed (**p < 0.01; student’s t-test for independent means). Data are means ± SEM (n=4 and 5 for APP/PS1 and WT mice respectively).

Figure 9. Phagocytosis of Aβ was decreased in microglia from APP/PS1 mice.
Isolated microglia from wildtype and APP/PS1 mice, were assessed for uptake of Aβ by staining with 6e10 as described in the Methods.
The number of $6 \times 10^6$ Iba1+ cells was decreased in cells isolated from APP/PS1 mice compared with WT mice as indicated by the representative immunohistogram and by analysis of the mean data (**p < 0.01; student’s t-test for independent means; n=4 and 5 for APP/PS1 and WT mice respectively).
Figure 2

A

Hexokinase II/actin (a.u.)

Control  |  IFNγ+Aβ

Control  |  IFNγ+Aβ

B

PKM2/actin (a.u.)

Control  |  IFNγ+Aβ

Control  |  IFNγ+Aβ

Hexokinase II
PKM2
β-Actin

C

PFKFB3 (Inducible)/actin (a.u.)

Control  |  IFNγ+Aβ

Control  |  IFNγ+Aβ

PFKFB3 (Inducible)
PFKFB3 (Constitutive)
β-Actin

D

PFKFB3 (Constitutive)/actin (a.u.)

Control  |  IFNγ+Aβ

Control  |  IFNγ+Aβ

E

HIF-1α/actin (a.u.)

Control  |  IFNγ+Aβ

Control  |  IFNγ+Aβ

HIF-1α
β-Actin

F

Mean fluorescence intensity

Control  |  IFNγ+Aβ

Control  |  IFNγ+Aβ
Figure 3

A

Ferritin/β-actin (a.u.)

Con | IFNγ+Aβ

**

B

Control | IFNγ+Aβ

C

% Iron Cells

Con | IFNγ+Aβ

**

D

Control | FeCl₃

E

PPKFB3/βα⁺ cell (%)

Con | FeCl₃

*
Figure 5

A

B

C

IFN_{\gamma} / β-Actin

Soluble Aβ_{1-40} (ng/mg)

Soluble Aβ_{1-42} (ng/mg)

WT  APP/PS1

WT  APP/PS1

WT  APP/PS1

*  ***  ***
Figure 6

A

WT

APP/PS1

B

C

Area (mm²)

Average intensity (AU)

WT

APP/PS1

**

**
Figure 8

A

WT

APP/PS1

B

Cytoplasmic PPKRB3 (Fluorescence CTCF)

0 1000 2000

WT APP/PS1

C

Nuclear PPKRB3 (Fluorescence CTCF)

0 500 1000

WT APP/PS1

D

Total PPKRB3 (Fluorescence CTCF)

0 2000 3000

WT APP/PS1

**