The influence of the R47H TREM2 variant on microglial exosome profiles

Running title: TREM2 variant microglial exosomes

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Abstract

Variants in the triggering receptor expressed on myeloid cells 2 (TREM2) gene are linked with an increased risk of dementia, in particular the R47Hhet TREM2 variant is linked to late-onset Alzheimer’s disease. Using human iPSC-derived microglia, we assessed whether variations in the dynamics of exosome secretion, including their components, from these cells might underlie some of this risk. We found exosome size was not altered between common variant controls and R47Hhet variants, but the amount and constitution of exosomes secreted were different. Exosome quantities were rescued by incubation with an ATP donor or with lipids via a phosphatidylserine TREM2 ligand. Following a lipopolysaccharide or phagocytic cell stimulus, exosomes from common variant and R47Hhet microglia were found to contain cytokines, chemokines, APOE and TREM2. Differences were observed in the expression of CCL22, IL-1β and TREM2 between common variant and R47Hhet derived exosomes. Furthermore unlike common variant-derived exosomes, R47Hhet exosomes contained additional proteins linked to negative regulation of transcription and metabolic processes. Subsequent addition of exosomes to stressed neurones showed R47Hhet-derived exosomes to be less protective. These data have ramifications for the responses of microglia in Alzheimer’s disease and may point to further targets for therapeutic intervention.

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Introduction

Alzheimer’s disease (AD) is the most common neurodegenerative disease with complex underlying disease mechanisms leading to inflammation in the brain and neuronal death (Minagar et al., 2002; Heneka et al., 2013; Zhang et al., 2013). Genome-wide association studies have linked enhanced risk of developing AD with inflammation, notably the innate immune cells of the brain, microglia (Naj et al., 2011; Guerreiro et al., 2013; Jonsson et al., 2013b). Microglia support neuronal functioning and viability through a range of processes, such as synaptic pruning, production of neurotrophic factors and reacting to threats in the environment (Minagar et al., 2002; Penzes et al., 2011; Parkhurst et al., 2013; Miyamoto et al., 2016). The triggering receptor expressed on myeloid cells-2 (TREM2) is involved in this surveillance function, as mutations in the gene for this receptor have been shown to reduce phagocytosis, migration and response to extracellular stimuli (Ulland et al., 2017; Zhong et al., 2017; Garcia-Reitboeck et al., 2018; Piers et al., 2020). Heterozygous mutations in TREM2, such as the R47Hhet mutation, are associated with AD (Guerreiro et al., 2013; Jonsson et al., 2013), whilst homozygous missense TREM2 mutations such as T66M cause Nasu-Hakola disease, a rare early-onset dementia with bone cysts (Paloneva et al., 2002; Guerreiro et al., 2013).

Exosomes are small extracellular vesicles ranging from 30-200nm in size (Hooper et al., 2012; Yang et al., 2018; Mathieu et al., 2019), created through endocytosis in multi-vesicular bodies and subsequent fusion and release (Hsu et al., 2010). Microglial exosomes have been shown to fulfill different functions, including supporting neuronal viability and functioning (Antonucci et al., 2012; Gupta and Pulliam, 2014; Asai et al., 2015). Upon activation of microglia with the toll-like receptor 4 agonist, lipopolysaccharide (LPS), subsequently secreted exosomes have been shown to contain inflammatory cytokines (Yang et al., 2018) and in turn, differentially affect neighbouring neurons. Since TREM2 variants and knock-outs have been shown to have an altered responsiveness to pathogenic stimulation (Ito and Hamerman, 2012) and display an impaired metabolism (Ulland et al., 2017; Piers et al., 2020), this could potentially affect exosomal production and release. We hypothesise that microglia with the AD-associated R47HhetTREM2 variant have an altered exosome secretion and that the exosomes have a differential effect on the viability of neurons. To study this, we used human induced pluripotent stem cells (iPSC) from patients carrying the R47Hhet mutation, differentiated into functional microglia (iPS-Mg). We analysed the composition of the exosomes and then examined their potential to influence neurones.

Methods
**iPS generation and iPS-Mg cells**

Ethical permission for this study was obtained from the National Hospital of Neurology and Neurosurgery and the Institute of Neurology joint research ethics committee (study reference 09/H0716/64). R47H<sup>het</sup> fibroblasts were acquired under a material transfer agreement between University College London and University of California Irvine Alzheimer’s Disease Research Centre (UCI ADRC; Professor M Blurton-Jones). Lines from 3 different patients carrying the R47H variant were used, with 8 clones in total being used. For one patient line, only 2 clones were included in the analysis due to chromosome abnormalities in the last clone.

Reprogramming of the R47H<sup>het</sup> fibroblasts was performed as previously described (Xiang <i>et al.</i>, 2018), with plasmids obtained from Addgene (#27077, #27078, #27080). After 7 days in vitro (DIV) nucleofected cultures were transferred to Essential 8 medium (Life Technologies) and colonies were picked after 25-30 DIV. Karyotype analysis was performed by The Doctors Laboratory (London). The iPSC were maintained and passaged in Essential 8 medium. Control lines used in this study were: CTRL1 (kindly provided by Dr Selina Wray, UCL Queen Square Institute of Neurology), BIONi010-C (obtained from EBiSC), SFC840 and SBAD03 (obtained from Stembancc) and KOLF2-C1 (obtained from Sanger Institute, available through EBiSC).

Using previously described protocols, iPSC-Mg were generated (Xiang <i>et al.</i>, 2018). Briefly, embryoid bodies (EB) were generated and maintained for 5 days in 100 ng/ml ROCK inhibitor, 50 ng/ml VEGF, 50 ng/ml BMP-4 and 20 ng/ml SCF. The EB were collected and transferred to flasks for further differentiation and kept in X-VIVO15 (Lonza) with 100 ng/ml MCSF and 25 ng/ml IL3. After 4-5 weeks of maturation, progenitor cells are collected and plated in iPSC-Mg maintenance medium, containing 100 ng/ml IL-34, 25 ng/ml MCSF and 5 ng/ml TGF-β, with the medium being changed every 3-4 days. Two weeks after plating, maturation medium containing 100 ng/ml CX3CL1 and 100 ng/ml CD200 was added to the cells in addition to IL-34, MCSF and TGF-β for another 3 days to achieve final maturation into iPSC-Mg, which was confirmed through a custom-designed gene array testing for 28 microglial genes (Xiang <i>et al.</i>, 2018).

**SH-SY5Y culture**

SH-SY5Y cells (a kind gift from Professor R De Silva, UCL Queen Square Institute of Neurology) were maintained and passaged in DMEM (ThermoFisher) containing 15% foetal bovine serum and penicillin/streptomycin (20 units/ml). The cells were differentiated using a previously established protocol (Encinas <i>et al.</i>, 2000). After 5 days of retinoic acid treatment (10 μM), the serum was removed from the medium and 50 ng/ml BDNF (Peprotech) was added for another 7 days. Exosomes were added to the SH-SY5Y neurons after differentiation for 24 h (unless stated differently), with the amount adjusted to total exosomal protein, this was 6 μg/ml.

Apoptotic neurons were generated from undifferentiated SH-SY5Y through heat-shock for 2 h at 45°C. The effectiveness of this method to induce apoptosis in SH-SY5Y was confirmed through Annexin V FITC (Miltenyi Biotech)/propidium iodide (PI) double-staining as previously shown (Garcia-Reitboeck <i>et al.</i>, 2018).
IPS-Mg Exosome collection and extraction

Medium was changed on the iPS-Mg 24 h before the experiment. Following treatment with 100 ng/ml LPS or apoptotic neurons at a ratio of 2:1 apoptotic neurons:microglia for 24 h, the supernatant (SN) from the iPS-Mg was collected and centrifuged for 15 min at 300 g. The SN was then used for extracting the exosomes.

Exosomes were extracted using an ExoQuick-TC kit (System Biosciences) by centrifugation of the supernatant for 30 min at 3 000 g before adding the ExoQuick solution overnight at 4°C. Following two further centrifugations at 1 500 g for 30 and 5 min respectively, the pellets from each step were collected. The final exosomal pellet was either resuspended in Dulbecco’s phosphate buffered saline without Ca²⁺/Mg²⁺ (DPBS), pH 7.0-7.3 or RIPA buffer (50 mM Tris-HCl, 150 mM NaCl, 1% SDS, 1% Triton X-100, 1 mM EDTA), freshly supplemented with a protease inhibitor cocktail (Halt protease and phosphatase inhibitor (Thermo Fisher Scientific), depending on downstream applications.

Nanoparticle tracking analysis (NTA)

Exosomes collected from 500 000 iPS-Mg over 24 h collection were diluted in 100 μl PBS and loaded into the recording chamber of a NanoSight LM10 (Malvern Panalytical). The content was recorded twice for 30 sec for each sample. For the human CSF standard (SBI System Biosciences, #EXOP-530A-1), 5.5 µg of exosomes were imaged. Analysis of the amount and size of particles per sample was completed with the NTA version 3.2.

Proteomic analysis

Liquid chromatography mass spectrometry (LC-MS) based proteomic analysis was performed using the TMT10-plex method for quantification (McAlister et al., 2014). Protein samples were reduced, alkylated and subjected to trypsin digestion using a filter-assisted sample preparation method, similar to a previously described protocol (Ostasiewicz et al., 2010). Three samples per condition, 12 in total, were arranged into two TMT 10-plex sets, each containing 2 samples of each condition and one reference sample, consisting of a mixture of all samples. To each 240 µl sample, 12 µl 2 M dithiothreitol were added, and the samples were then incubated for 30 min at 60°C. Urea (8 M, 250 µl) was added, and the samples were loaded on molecular-weight cut-off filters (Nanosep, 30 kDa, Pall). Ultrafiltration was performed by centrifugation at 10 000 g for 10 min at RT. The flow-through was discarded and the filters were washed twice with 200 µl 8 M Urea, passing the liquid through the filter by centrifugation as above. The filters were then washed with 200 µl 0.5 % sodium deoxycholate (SDC), 50 mM triethylammonium bicarbonate (digestion buffer). Alkylation of cysteine disulfides was performed by adding 100 µl 18 mM iodoacetamide in digestion buffer to the filters, and incubating for 20 min at RT in darkness. The liquid was removed by centrifugation and the filters were washed twice with 100 µl digestion buffer. Trypsin (Sequencing grade, modified, Promega) in digestion buffer (14 ng/µl, 70 µl) was added to the filters, followed by incubation at 37°C overnight. An additional 10 µl trypsin aliquot (0.1 µg/µl) was added and the samples were further incubated for another four h. Tryptic peptides were collected by centrifugation. TMT 10-plex reagents (8 µg dissolved in 42 µl acetonitrile) were added to the samples followed by incubation for 1 h at room temperature on a shaker. Hydroxylamine (8 µl, 5%)
were added and the samples were incubated for 15 min to inactivate excess TMT reagent, after which the individually TMT-labelled samples were pooled into two multiplex samples. The samples were acidified by addition of 10% trifluoroacetic acid to pH<2 to precipitate the SDC. The samples were centrifuged at 18 000 g for 10 min and the supernatants recovered. The TMT labelled peptides were fractionated by reversed-phase chromatography at basic pH (Batth et al. 2014). Briefly, the multiplex samples were re-dissolved in 16 µl 2 % AcN, 5 mM ammonium hydroxide and loaded on an Ultimate 3000 HPLC system (Thermo Fisher). Offline peptide separation was performed over an XBridge BEH130 C18 3.5 µm, 2.1 mm x 250 mm analytical column at a flow of 100 µl/min, using a linear gradient and collecting 1 fraction/min for 72 min. Collected fractions were then concentrated to 12 fractions, which were dried by vacuum centrifugation.

For LC-MS, samples were dissolved in 6 µl of 0.1% TFA and analyzed by LC-MS on nano-HPLC instrument (Ultimate 3000 RSLC nano, Thermo Scientific) fitted with a 75 µm x 2 cm trap column (PepMap C18, Thermo scientific) and a 75 µm x 50 cm C18 column (PepMap C18, Thermo scientific), coupled to a Fusion Trivrid mass spectrometer (Thermo scientific). Following 10 min sample loading onto the trap, separation was performed using the following gradient between 0.1% formic acid (Buffer A) and 84% acetonitrile (Buffer B): t=10 (min), 0% B; t=105, %B=40; t=112, %B=99. The mass spectrometer was operated in the positive ion mode, with the following settings: resolution 120 000; m/z range 350-1 400; max injection time 50 ms; AGC target 4e5. Data-dependent acquisition was used to record up to 10 consecutive fragment ion spectra (MS2) per full scan spectrum, selecting precursor ions with charge states 2-7 in decreasing order of intensity, and using 15 s dynamic exclusion. The isolation window was set to 1.2 m/z. The instrument settings for the MS2 scans were: resolution 50 000; fixed first mass m/z 110; max injection time 86 ms; AGC target 5e4.

The LC-MS data were processed using the software Proteome Discoverer 2.3 (Thermo). Protein identification was performed using the Mascot software (Matrix Science), searching the human subset of SwissProt, with the following settings: cleavage enzyme: trypsin; precursor mass tolerance: 15 ppm; fragment mass tolerance: 0.05 Da; static modifications: TMT10-plex; dynamic modifications: phosphorylation. Percolator scoring was used, and validation was performed using the target-decoy approach, using 1% FDR as cut-off for peptide identification. Protein identifications were accepted that had at least one matching peptide.

**Microglial gene array**

A custom gene array based on published microglial expression data (Butovsky et al., 2014; Muffat et al., 2015; Abud et al., 2017; Haenseler et al., 2017) was used to confirm a microglial signature in our iPS-Mg cultures (TaqMan Array Plate 32 plus Candidate Endogenous Control Genes; Thermo Fisher Scientific, see Table 1), as previously described (Xiang et al., 2018). The heatmap was generated through MATLAB.
Table 1 Microglial gene signature primer details:

<table>
<thead>
<tr>
<th>Gene name</th>
<th>Primer ID</th>
</tr>
</thead>
<tbody>
<tr>
<td>18 s rRNA</td>
<td>Hs99999901_s1</td>
</tr>
<tr>
<td>GAPDH</td>
<td>Hs99999905_m1</td>
</tr>
<tr>
<td>HPRT</td>
<td>Hs99999909_m1</td>
</tr>
<tr>
<td>GUSB</td>
<td>Hs99999908_m1</td>
</tr>
<tr>
<td>APOE</td>
<td>Hs00171168_m1</td>
</tr>
<tr>
<td>C1QA</td>
<td>Hs00706358_s1</td>
</tr>
<tr>
<td>C1QB</td>
<td>Hs00608019_m1</td>
</tr>
<tr>
<td>ITGAM</td>
<td>Hs00167304_m1</td>
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<tr>
<td>CSF1R</td>
<td>Hs00911250_m1</td>
</tr>
<tr>
<td>CX3CR1</td>
<td>Hs01922583_s1</td>
</tr>
<tr>
<td>GAS6</td>
<td>Hs01090305_m1</td>
</tr>
<tr>
<td>GPR34</td>
<td>Hs00271105_s1</td>
</tr>
<tr>
<td>AIF1</td>
<td>Hs00610419_g1</td>
</tr>
<tr>
<td>MERTK</td>
<td>Hs01031979_m1</td>
</tr>
<tr>
<td>OLFML3</td>
<td>Hs01113293_g1</td>
</tr>
<tr>
<td>PROS1</td>
<td>Hs00165590_m1</td>
</tr>
<tr>
<td>SALL1</td>
<td>Hs01548765_m1</td>
</tr>
<tr>
<td>SLCO2B1</td>
<td>Hs01030343_m1</td>
</tr>
<tr>
<td>TGFBR1</td>
<td>Hs00610320_m1</td>
</tr>
<tr>
<td>TMEM119</td>
<td>Hs01938722_u1</td>
</tr>
<tr>
<td>TREM2</td>
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</tr>
<tr>
<td>BIN1</td>
<td>Hs00184913_m1</td>
</tr>
<tr>
<td>CD33</td>
<td>Hs01076282_g1</td>
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<td>SPI1</td>
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<tr>
<td>HEXB</td>
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<tr>
<td>ITM2B</td>
<td>Hs00222753_m1</td>
</tr>
<tr>
<td>C3</td>
<td>Hs00163811_m1</td>
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<tr>
<td>A2M</td>
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<tr>
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<tr>
<td>RGS1</td>
<td>Hs01023772_m1</td>
</tr>
<tr>
<td>FTL</td>
<td>Hs00830226_gH</td>
</tr>
<tr>
<td>P2RY12</td>
<td>Hs01881698_s1</td>
</tr>
</tbody>
</table>
**Immunoblotting**

For the detection of exosomal CD63, exosomal lysate was deglycosylated using a PNGase kit (New England Biolabs), following manufacturer’s instructions. For all other exosomal markers, the exosomal lysates were not deglycosylated. Exosomal pellets or iPS-Mg cells were lysed in RIPA buffer as described above. Cell samples were cleared of nuclear material and cell fragments by centrifugation at 15 000 \( g \) for 15 min. The cell lysate was denatured using sample buffer and separated by SDS-PAGE. Proteins were transferred onto nitrocellulose membranes and blocked with 5% milk in phosphate buffered saline solution with 1% TWEEN-20 (PBST). For exosomal blots, positive controls consisted of purified exosomes from human serum or Jurkat T lymphocytes (System Biosciences).

Primary antibodies (see Table 2) below were incubated with the blots overnight at 4°C, with the appropriate secondary fluorescently conjugated antibody incubated at 1:5 000 for 1 h at RT.

**Table 2 List of antibodies used:**

<table>
<thead>
<tr>
<th>Antibody</th>
<th>Concentration</th>
<th>Diluted In</th>
<th>Company</th>
</tr>
</thead>
<tbody>
<tr>
<td>CD63</td>
<td>1:1000</td>
<td>5% milk</td>
<td>Abcam</td>
</tr>
<tr>
<td>ALIX</td>
<td>1:1000</td>
<td>5% milk</td>
<td>Cell Signalling Technologies</td>
</tr>
<tr>
<td>CD9</td>
<td>1:1000</td>
<td>5% milk</td>
<td>System Biosciences</td>
</tr>
<tr>
<td>CD81</td>
<td>1:1000</td>
<td>5% milk</td>
<td>System Biosciences</td>
</tr>
<tr>
<td>Calnexin</td>
<td>1:1000</td>
<td>1% BSA</td>
<td>Santa Cruz</td>
</tr>
<tr>
<td>β-actin</td>
<td>1:10,000</td>
<td>1% milk</td>
<td>Thermofisher</td>
</tr>
</tbody>
</table>

**Exosomal uptake**

Uptake of exosomes into SH-SY5Y cells was assessed with fluorescence activated cell sorting (FACS). The exosomes were labelled with cell Vybrant Dil (Thermofisher) at a dilution of 1:200 at 37°C for 1 h. The excess dye was removed using 3 kDa NMWL columns (Amicon), by centrifuging excess dye out for 20 min at 14 000 \( g \). To control for any remaining unbound Dil present after centrifugation, PBS alone was incubated with the dye as well and used as a negative control. The labelled exosomes were then added to SH-SY5Y for 2 h, before the cells were detached using PBS and analysed by FACS. As a negative control, some SH-SY5Y were pre-treated with cytochalasin D (CytoD, 10 \( \mu \)M), chlorpromazine (CPZ, 5 \( \mu \)g/ml) or genistein (200 \( \mu \)M) to inhibit phagocytosis, for 30 min before exposure to labelled exosomes. The data were analysed using Flowing Software 2.

To further visualize exosomal uptake, exosomes were stained with cell Vybrant DiO (Thermofisher) at a dilution of 1:200 at 37°C for 1 h, before excess dye was removed using 3 kDa NMWL columns (Amicon), as described above. The SH-SY5Y were labelled with BioTracker 555 Orange Cytoplasmic Membrane Dye (Merck Millipore) at 2 \( \mu \)g/ml for 10 min. The cytoplasmic membrane labelling was carried out either before exosome incubation in the 1 h condition or after exosome incubation in the 24 h condition. Nuclei were stained with DAPI before being mounted and imaged with a Zeiss LSM710 confocal microscope.
Cell death FACs

To analyse the ability of exosomes to modify neuronal death, we treated SH-SY5Y cells with hydrogen peroxide (H$_2$O$_2$, 100 μM) for 24 h with or without iPS-Mg exosomes. Subsequently SH-SY5Y cells were labelled with propidium iodide (PI, 5 μg/ml) for 15 min before cell death was analysed by FACs. Controls consisted of unstained cells or cells boiled at 95°C for 1 h prior to PI labelling. Data were analysed using Flowing Software 2 and plotted as % of cells in H2 region/total number of cells.

ATP analysis

ATP levels were determined in the iPS-Mg using a commercially available ATP kit (#A22066, Thermo Fisher Scientific). The cells were plated in black walled, glass-bottomed 96-well plates. The iPS-Mg were treated with either apoptotic neurons (2:1 apoptotic neurons: iPS-Mg) or cyclocreatine (10 mM) for 24 h before they were lysed in cell lysis reagent (#1715747, Boehringer Mannheim) and centrifuged at 1,000g for 1 min. Then, 10 μl of sample supernatant or ATP standard was added to 100 μl of the ATP reaction mix, after the background luminescence was read. Luminescence was recorded on a Tecan 10M plate reader. Data were normalised to ATP standard curves and then to the protein concentration of the sample determined by BCA assay.

Statistical Analysis

The results were represented as mean of at least three separate experiments with internal replicates of at least three ± SEM with a $p$ value of 0.05 or below considered statistical significant. The results were analysed using Prism Software version 5. Analysis was performed on pooled control lines and pooled R47H$^{het}$ cell lines.

With regard to the LC-MS data, the relative abundances, normalised to the reference sample, were analysed using MATLAB and tested for normality and the potential influence of the different experimental runs. Following from this quality control, the abundances were normalised to the individual reference sample for each experimental run, accounting for the overall different number of proteins detected in the samples. In addition to this, for further analysis, the abundance ratios were log$_{10}$-transformed to generate a normal distribution, allowing for standard statistical tests to be performed on the data. The results from the data base search were analysed using MATLAB. Analysis of the functional effect of the different proteins was conducted using FunRich and STRING.

Power calculations for exosomal size differences as measured through NTA (Fig 1D) were performed in line with Yang et al., 2018 and statistics were performed using the D90 measurement (calculated = 9). We include n=10 for the Cv and n=11 for the TREM2 R47H$^{het}$ and therefore achieve the necessary power required to state that based on the D90 measurement no difference is observed.

Data availability

All data are available upon request. All data points are shown on graphs.
Results

Exosome characterisation

To confirm that the particles in the exosomal fraction were indeed exosomes, we assessed whether they displayed common exosomal characteristics. Classical exosomal markers, such as CD63, CD9, ALIX, CD81, HSP70, TSG101 and Flotillin-1 (Hooper et al., 2012; Witwer et al., 2013; Frohlich et al., 2014; Mehdiani et al., 2015; Schiera et al., 2015; Horibe et al., 2018; Chen et al., 2019) were all found to be expressed in the iPS-Mg exosomal fraction of both common variant (Cv) and R47H\textsuperscript{het} derived exosomes (Fig 1A and Supplementary Fig 1C). Calnexin, an endoplasmic reticulum (ER) marker, was absent from the exosomal fraction, but shown to be present in the cell lysate (Fig 1B) confirming the absence of apoptotic bodies from the exosomal fraction (Lötvall et al., 2014).

The size distribution of the particles as measured through NTA was similar in both Cv and R47H\textsuperscript{het} variants and comparative in size with a human CSF exosome standard (Fig 1C and 1D). The average size of the particles was not significantly different from each other (Fig 1D). This was further confirmed by the D90 measurement, again showing no statistical size difference between the Cv and R47H\textsuperscript{het} exosomes (Supplementary Fig 1B). These results confirm that the particles found in the iPS-Mg exosomal fraction are indeed exosomal and that they can be extracted reliably from iPS-Mg with \textit{TREM2} variants.

Exosome secretion

As exosome secretion is dependent on interactions with the cytoskeleton and energy availability within the cells (Hoshino et al., 2013; Zou et al., 2019), we investigated the exosome secretion concentration in iPS-Mg carrying the R47H\textsuperscript{het} \textit{TREM2} variant, as it has been shown that these cells have deficits in both their cytoskeleton and metabolic fitness (Takahashi et al., 2005; Ulland et al., 2017; Piers et al., 2020). Measurements by NTA indicated that exosomes were enriched following purification (Video 1) but not present in cell medium alone (Video 2). Further NTA measurements indicated that the concentration of exosomal particles secreted over 24 h from the R47H\textsuperscript{het} expressing iPS-Mg compared with Cv was significantly different (Fig 1E). Subsequent analysis of exosomal CD63, a classical marker of exosomes and an approximation of exosomal number, was significantly decreased (Fig 1F).

As extracellular stimuli, such as inflammatory signals, can influence the profile of secreted exosomes, both in their size and amount (Qu et al., 2007; Yang et al., 2018), we investigated the amount of exosome secretion following activation of iPS-Mg with 100 ng/ml LPS for 24 h. As we found deficits in exosome secretion in the \textit{TREM2} variant (Fig 1F), we also activated the TREM2-DAP12 pathway through the addition of apoptotic neurons, which express the TREM2 ligand phosphatidylserine (PS\textsuperscript{*}) on their surface (Wang et al., 2015; Garcia-Reitboeck et al., 2018; Shirotani et al., 2019) and used exosomal CD63 as a readout of exosomal production.

CD63 expression was unchanged in exosomes from Cv or R47H\textsuperscript{het} iPS-Mg with LPS or apoptotic neuron treatment (but was significantly lower in unstimulated and LPS stimulated R47H\textsuperscript{het}-derived exosomes Fig 2A and B). Apoptotic neuron-treatment increased exosome secretion from R47H\textsuperscript{het} iPS-Mg (Fig 2A and B), whilst Cv exosome secretion was not affected. Apoptotic neurons have been shown...
to supply iPS-Mg with additional energy (Cosker et al., 2020), possibly by providing additional energy via lipid catabolism (Nadjar, 2018), rescuing the energy deficit of R47Hhet iPS-Mg previously reported (Piers et al., 2020). Based on this, we tested whether the increase in exosome secretion from R47Hhet following exposure to apoptotic neurons was due to an increase in energy supply. Thus, iPS-Mg were treated with 10 mM cyclocreatine for 24 h, which acts as an ATP supply fuelling oxidative phosphorylation rather than glycolysis, which we have shown to be deficient in TREM2 variant iPS-Mg (Piers et al., 2020). Again, an increase in exosomal CD63 expression was observed in the R47Hhet exosomes (Fig 2C and D), whilst there was no significant difference in Cv, suggesting that the reported energy deficit also underlies the reduced secretion of exosomes in the R47Hhet iPS-Mg. Furthermore analyses of ATP content revealed that there was a significantly lower level of ATP in R47Hhet cells relative to Cv, and this was increased to Cv levels by prior exposure to PS+ cells or cyclocreatine (Supplementary Fig 2A).

**Exosome content by MS analysis**

LC-MS performed on exosomes extracted from iPS-Mg revealed over 70% of the proteins detected have been previously linked to exosomal proteins (Fig 3A) and furthermore, by virtue of their cellular compartment of origin, these proteins were found in exosomes, (Fig 3B), providing additional support that the extracted particles from iPS-Mg were exosomes.

Analysis of baseline differences between exosomes secreted from Cv and R47Hhet iPS-Mg by volcano plot revealed nine proteins with higher abundance in R47Hhet iPS-Mg (ratio > 2, p < 0.05). (Fig 3C, and Supplementary Fig 2C). STRING analyses revealed that some of these proteins are involved in the negative regulation of transcription and metabolic processes (Fig 3D).

Analysis of microglial gene signature was assessed in all lines studied using a custom gene array based on previously published expression data (Xiang et al., 2018). Lines from both TREM2 genotypes used in this study cluster strongly together with only one outlier observed clustering more closely with human macrophages (Fig 3E). These data suggest differences observed in exosomal content between genotypes is not due to gross microglial gene signature differences at basal.

**Exosome content following LPS or apoptotic cell stimulation**

We investigated changes in the relative expression of exosomal proteins following treatment of iPS-Mg with LPS, as previous research has suggested changes in exosomal proteins occur following activation of microglia (Yang et al., 2018). Volcano plots of proteomic data showed that several proteins differ in abundance in exosomes derived from Cv iPS-Mg and R47Hhet iPS-Mg with and without stimulation with LPS (Fig 4A and 4B respectively). Following activation with LPS, the content of the iPS-Mg exosomes changed significantly to include increased secretion of cytokines in both Cv and R47Hhet TREM2 variants, such as TRAF1, (Fig 4C), whilst others such as C1QA and APOE showed a significant reduction in exosomes from LPS-treated Cv and R47Hhet iPS-Mg. TREM2 was not significantly reduced in exosomes from Cv but was from R47Hhet exosomes (Fig 4C).

Volcano plots of proteomic data from Cv or R47Hhet derived exosomes exposed to apoptotic neurons revealed differences in abundance (Fig 5A and 5B). Furthermore CCL22 was significantly reduced in
R47H\textsuperscript{het} exosomes compared with Cv following stimulation with apoptotic neurons (Fig 5C). Furthermore a significant number of proteins were not increased in R47H\textsuperscript{het} derived exosomes compared with Cv following exposure of iPS-Mg to apoptotic neurons (Fig 6).

**Uptake of iPS-Mg exosomes into neurons**

As microglia exosomes have been shown to affect neuronal functioning (Antonucci et al., 2012; Chen et al., 2019) we determined whether iPS-Mg-derived exosomes from Cv or R47H\textsuperscript{het} cells were taken up into neurons (Fig 7). Uptake of the exosomes was verified both through confocal microscopy, showing DiO labelled exosomal particles inside the neurons within 1 h of incubation and 24 h (Fig 7B). This was also quantified by FACs, showing a significant uptake of Dil labelled exosomes into the neurons after 2 h in comparison with unstained cells (Fig 7C and D). No significant difference was observed in the level of uptake of exosomes from Cv or R47H\textsuperscript{het} iPS-Mg, albeit that the R47H\textsuperscript{het} exosomes appeared to be taken up at slightly lower levels. As a negative control, exosome-free saline solution was incubated with label to control for any carry-over of Vybrant dye, however this appeared not to be a confounding factor (Fig 7D). Uptake could be inhibited by CPZ or genistein, but not cytochalasin D (Fig 6Cii).

Based on our finding of changes in exosome content in Cv and R47H\textsuperscript{het} exosomes (Fig 4, 5 and 6), the effect of exosomes from either Cv or R47H\textsuperscript{het} iPS-Mg on neuronal viability was assessed. Following incubation of exosomes with differentiated SH-SY5Y (see Supplementary Fig 3 for SH-SY5Y differentiation), no significant differences in the levels of neuronal death were observed after addition of Cv or R47H\textsuperscript{het} exosomes derived from non-treated iPS-Mg (Fig 8A and 8B).

Previous research has shown that the protective functions of glial exosomes are more easily revealed when the SH-SY5Y were subjected to H\textsubscript{2}O\textsubscript{2}-induced cell stress (Pascua-Maestro et al., 2019). Based on this, the ability of exosomes from iPS-Mg to rescue SH-SY5Y neurons from H\textsubscript{2}O\textsubscript{2}-induced cell death was investigated (Fig 8C and D). H\textsubscript{2}O\textsubscript{2} increased neuronal death significantly (Fig 8D). However, the addition of Cv iPS-Mg-derived exosomes protected SH-SY5Y neurons from H\textsubscript{2}O\textsubscript{2}-induced death as there was no significant difference between the levels of death in non-treated neurons compared with those treated with Cv exosomes and H\textsubscript{2}O\textsubscript{2} indicating a protective effect of Cv-derived exosomes (Fig 8D). Conversely, the level of death in neurons treated with H\textsubscript{2}O\textsubscript{2} plus exosomes from R47H\textsuperscript{het} iPS-Mg was no different from neurons exposed solely to H\textsubscript{2}O\textsubscript{2}, indicating that these exosomes were not protective (Fig 8D).

**Discussion**

We were able to reliably extract exosomes from iPS-Mg, the exosomes showed a range of common exosomal markers, and were also devoid of markers which might indicate contamination from different cellular components, such as ER (Fig 1B). The average size of the exosomes from cells with or without the R47H\textsuperscript{het} mutation was approximately 150 nm, and a comparison of the size of exosomes from Cv with R47H\textsuperscript{het} indicated that they were not significantly different (Fig 1D and Supplementary Fig 1B), suggesting that the R47H\textsuperscript{het} variant does not affect exosome size. Our findings also suggest that the particles as analysed by NTA were potentially larger than those reported elsewhere (Witwer et al., 2013), but in part this can be explained by the measuring technique we employed, as even the CSF
Exosome standard was measured to be around the same size as the iPS-Mg derived exosomes (Fig 1D). Due to the size of the NTA chamber, during imaging, many particles are out of focus and could skew the average size measured. The R47H<sup>het</sup> variant appeared to influence exosome number (Fig 1E), which was also reflected in the decrease in exosomal CD63 (Fig 1F). This could be due to the metabolic deficit in microglia with this variant (Ulland et al., 2017; Piers et al., 2020), as supplying the cells with additional lipids in the form of apoptotic cells was able to alter this to control levels for the protein we tested as indeed did cyclocreatine, generating ATP in the cells (Kurosawa et al., 2012).

In addition to influencing exosome secretion, the R47H<sup>het</sup> variant altered the packaging of proteins into exosomes, as shown through proteomic analysis with nine proteins increasingly packaged into exosomes from R47H<sup>het</sup> iPS-Mg which were found to be involved in negative regulation of metabolic processes (Fig 3C, Fig 3D and Supplementary Fig 2B). In a recent study, one of the most altered AD protein co-expression modules was linked to sugar metabolism in glia such as microglia (Johnson et al., 2020). Our data suggest that exosome signalling for metabolism is thus likely to be decreased in R47H<sup>het</sup> iPS-Mg.

Whether these proteins are present in R47H<sup>het</sup> exosomes because of the metabolic deficits present in the microglia (Ulland et al., 2017; Piers et al., 2020) is not yet known. However, our study confirms previous research that exosomes from LPS-activated microglia contain more inflammatory cytokines (Yang et al., 2018), although we also found that TREM2 was present in exosomes but also decreased following LPS stimulation, and significantly so in R47H<sup>het</sup> derived exosomes (Fig 4C). There were also differences in the proteins present in exosomes from Cv versus R47H<sup>het</sup> iPS-Mg, and in particular, a number of proteins were not increased in the latter after exposure to apoptotic cells. This suggests that whilst some proteins can be increased by a TREM2 ligand in Cv, in the R47H<sup>het</sup> iPS-Mg, this is not the case.

Whilst full transcriptomic analysis of the iPS-Mg lines used in this study may go further to identify if there were significant differences in the underlying TREM2 genotypes that could contribute to exosomal content variance, the microglial gene signature array performed here suggests limited differences between the genotypes (Fig 3E). Strong clustering between all the lines was observed (with the exception of one potential outlier repeat), suggesting that the TREM2 genotypes do not differ significantly at the microglial signature level. This observation means differences in exosomal content were not likely to be due to differences in the underlying basal microglia signature of the TREM2 variants.

Exosomes can be taken up through a range of different routes into recipient cells (Mulcahy et al., 2014; Schneider et al., 2017; Horibe et al., 2018). We did not find exosome uptake into SH-SY5Y neurons to be inhibited by CytoD, which mainly inhibits phagocytosis and macropinocytosis (Kuhn et al., 2014) through inhibition of actin polymerization, but other uptake mechanisms into SH-SY5Y cells such as membrane fusion and lipid raft-mediated endocytosis are still available for the exosomal cell entry, possibly explaining why exosomal uptake into SH-SY5Y was not be blocked. CPZ, an inhibitor of clathrin-mediated endocytosis (Liu et al., 2007; Kuhn et al., 2014; Wesén et al., 2017) or genistein, an inhibitor of caveolin-mediated endocytosis (Horibe et al., 2018), did inhibit uptake however.
Since the same amount of exosomal protein from Cv or R47Hhet iPS-Mg was added to the neurons and no significant differences were observed in uptake, this suggests that the actual composition and content of R47Hhet exosomes influences neuronal functioning. When we compared whether the exosomes from either Cv or R47Hhet affected baseline neuronal viability (Fig 8A and B), there was no difference, suggesting that the composition of exosomes from unstimulated iPS-Mg does not in itself influence neuronal viability over the time course of our experiments. However exosomes from Cv were protective against an H₂O₂ stress whilst those from R47Hhet iPS-Mg were not (Fig 8C and D); this may be due to our finding of negative regulators of metabolism being preferentially packaged into R47Hhet exosomes. It was interesting that exosomes contained TREM2, present in exosomes from Cv and R47H iPS-Mg, although we do not know as yet if this is the full length protein. It has recently been reported that TREM2 is present in exosomes in serum and that these exosomes may transport TREM2 into the brain (Raha-Chowdhury et al., 2018). We show here that exosomes were taken up in neurons, the ramifications of which would need to be further investigated.

In this study, we have shown that exosomes can be reliably extracted from human iPS-Mg expressing TREM2 variants and that disease-associated TREM2 variants secrete less exosomes compared with Cv, potentially due to the reported metabolic deficits. Differences in exosomal protein content between the Cv and R47Hhet iPS-Mg may influence the ability of neurons to protect themselves against H₂O₂ induced stress or other stresses. These differences may influence the ability of human microglia to protect neurons in Alzheimer’s disease if the microglia express the R47Hhet variant.

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Competing interests
H Zetterberg has served at scientific advisory boards for Denali, Roche Diagnostics, Wave, Samumed and CogRx, has given lectures in symposia sponsored by Fujirebio, Alzecure and Biogen, and is a co-founder of Brain Biomarker Solutions in Gothenburg AB (BBS), which is a part of the GU Ventures Incubator Program (outside submitted work). The other authors declare that they have no conflict of interest.

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Figure legends

Figure 1 – Exosome characterisation

Western blot of exosomes extracted from common variant (Cv) and R47Hhet TREM2 expressing iPS-Mg express the exosomal markers CD63, and CD9, with exosomes extracted from human serum used as a positive control (Exo standard) (A). The extracted exosomal fraction did not contain calnexin, indicating that the fraction is clear of ER contamination, whilst the exosome fraction contained another exosomal marker CD81 (B). Size distribution curves of exosomes from Cv and R47Hhet iPS-Mg, against a human CSF standard assessed by nanoparticle tracking analysis (NTA) (C, D). Quantification of the exosome concentration of Cv or R47Hhet iPS-Mg derived exosomes following NTA (E) or by western blot of CD63 expression (F). The full blots for A and B can be found in Supplementary Fig 5 and expanded datasets for D-F in Supplementary Fig 4. All data are the mean ± SEM, n ≥ 4 with one-way ANOVA (D) and Students t-test (E and F) * ρ < 0.05, *** ρ < 0.001.

Figure 2 – Exosome secretion can be rescued

Western blot of exosomal CD63 levels, (used to approximate exosome secretion), following stimulation of iPS-Mg with 100 ng/ml LPS or apoptotic neurons (ratio of 2 apoptotic neurons: 1 iPS-Mg) for 24 h (A) and quantification of CD63 expression (B). Western blot of exosomal CD63 (C) and CD63 quantification (D) following exposure of iPS-Mg to 10 mM cyclocreatine for 24 h. The full blots for A and C can be found in Supplementary Fig 5 and expanded datasets for B and D in Supplementary Fig 4. Data are the mean ± SEM, n ≥ 3 following two-way ANOVA followed by Tukey’s post-hoc tests, n.s not significant, * ρ < 0.05, ** ρ < 0.01.

Figure 3 – Proteomics of exosomes

Mass spectrometry was performed on exosomes extracted from iPS-Mg. Over 70% of the proteins detected by mass spectrometry (blue) have been previously linked to exosomal proteins (A). 2147 proteins have been previously published in a meta-analysis of high-throughput proteomics of exosomes (Choi et al., 2013), whilst 475 proteins were shared with the online repository of exosomal proteins (ExoCarta-Accession, red). This was also verified following analysis of the proteins for their cellular
compartment of origin (B). The proteins are significantly associated with exosomes, followed by the lysosomes and cytoplasm relative to the annotated control genome. This confirms the extracted particles to be exosomes. The difference between exosomes secreted from Cv and R47H\textsuperscript{het} iPS-Mg was compared by volcano plot (C). The ratio in abundance of proteins in R47H exosomes, compared with Cv exosomes, was plotted in the x-axis whilst the significance of the change, in form of the \( \rho \) value, was plotted in the y-axis, meaning that any proteins in the top left and right section had a significant 2-fold change. Nine proteins were found to be significantly packaged into exosomes originating from R47H\textsuperscript{het} iPS-Mg compared with exosomes from Cv iPS-Mg (C). The function of these proteins was analysed through STRING analysis (D), which indicated that some of the proteins increasingly packaged into R47H\textsuperscript{het} exosomes are involved in the negative regulation of transcription and metabolic processes, \( n = 3 \) independent experiments. Heatmap of the analysis of microglial gene signature in all iPS-Mg lines and clones used in this study using a custom gene array (E).

**Figure 4 – Stimulation-induced differential changes in exosomal proteins following TLR4 stimulation**

Volcano plots of the effect of LPS stimulation on exosomal proteins in Cv iPS-Mg (A) and R47H\textsuperscript{het} iPS-Mg (B). Proteins of interest were circled and further plotted (C). Data are mean ± SEM with \( n = 3 \) independent experiments. Two-way ANOVA followed by Tukey’s post-hoc test with * \( \rho < 0.05 \), ** \( \rho < 0.01 \).

**Figure 5 – Changes in exosomal protein following TREM2 ligand exposure**

Volcano plots for the effect of apoptotic neuron treatment on exosomal proteins in Cv iPS-Mg (A) and R47H\textsuperscript{het} iPS-Mg (B). Proteins of interest were further plotted (C). Data are mean ± SEM with \( n = 3 \) independent experiments. Two-way ANOVA followed by Tukey’s post-hoc test with * \( \rho < 0.05 \), ** \( \rho < 0.01 \).

**Figure 6 – Comparison of exosomal proteins increased following exposure to a TREM2 ligand**

Volcano plots comparing the increased abundance levels of exosomal proteins from Cv or R47H\textsuperscript{het} iPS-Mg treated with apoptotic neurons to stimulate TREM2-mediated pathways. (A). Abundance levels of some of the proteins observed to change following this stimulation (B). Data are mean ± SEM with \( n = 3 \). Two-way ANOVA followed by Tukey’s post-hoc test with * \( \rho < 0.05 \), ** \( \rho < 0.01 \).

**Figure 7 – Uptake of iPS-Mg exosomes into SH-SY5Y**

IPS-Mg derived exosomes were added to differentiated SH-SY5Y (workflow: A). Uptake of exosomes into SH-SY5Y was determined by imaging DiO-labelled exosomes inside SH-SY5Y, (the latter were stained with BioTracker 555 Orange Cytoplasmic Membrane Dye, 2 \( \mu \)g/ml) after 1h and 24 h of incubation with the exosomes (B) Scale bar = 10 \( \mu \)m. Uptake was quantified by FACs analysis of Dil-labelled exosomes and compared with unstained cells (Unst), (C) or with Dil-labelled PBS as a negative control (PBS) or following 30 min preincubation with cytochalasin D (CytoD, 10 \( \mu \)M), chlorpromazine (CPZ, 5 \( \mu \)g/ml) or genistein (200 \( \mu \)M) (D). Expanded dataset for D can be found in Supplementary Fig...
4. Data are the mean ±SEM with one-way ANOVA followed by Tukey’s post-hoc test, n ≥ 4. ** p < 0.01, * p < 0.05, ns, not significant.

**Figure 8 – SH-SY5Y viability following exosome incubation**

The effect of exosomes on the viability of SH-SY5Y assessed with prodidium iodide (PI) staining and FACS analysis. Baseline cell death following addition of exosomes from unstimulated iPS-Mg from Cv or R47Hhet to SH-SY5Y (A and B). Cell death in SH-SY5Y challenged with H₂O₂ (100 μM) for 24 h in the presence of exosomes from Cv or R47Hhet iPS-Mg or (C and D). Expanded datasets for B and D can be found in Supplementary Fig 4. One-way ANOVA followed by Tukey’s post-hoc test, with n ≥ 5 and n ≥ 4 for B and D respectively, **p < 0.01.

**Video 1.** Representative video of NTA showing particles in the exosomal fraction of iPS-Mg.

**Video 2.** Representative video of NTA showing no particles in the cell medium itself.
Figure 1

190x275mm (300 x 300 DPI)
Figure 2
Figure 3

190x275mm (300 x 300 DPI)
Figure 4

Cv exosomes

Increased in NT

Increased with LPS

R47H exosomes

Increased in NT

Increased with LPS

C

CXCL5

NT

LPS

CCL22

C1QA

IL-1β

APOE

TREM2

190x275mm (300 x 300 DPI)
Figure 5

190x275mm (300 x 300 DPI)
Figure 6

Exosomal content after iPS-Mg were treated with apoptotic neurons

A

B

MAP1A

TNFAIP8L2

TMPO α

UBQLN2

TMPO β/γ

Cv  R47H***

190x275mm (300 x 300 DPI)
Figure 7

190x275mm (300 x 300 DPI)
Figure 8

190x275mm (300 x 300 DPI)
50 word summary

Mallach et al. investigated the effect of the R47H TREM2 variant in human induced pluripotent stem cell microglia-derived cells on exosome secretion and content. They found decreased exosome secretion rate from TREM2 variants, linked to decreased energy availability, and differences in exosomal protein content, which influenced the viability of neuron-like cells.
Induced pluripotent stem cell-derived microglia

Neuron-like cells

Exosomes

Reduced secretion rate

Altered content

R47H<sup>het</sup> TREM2 variant

Not protective

Reduce cell death