Research Paper

Oxysterols present in Alzheimer’s disease brain induce synaptotoxicity by activating astrocytes: A major role for lipocalin-2

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ABSTRACT

Among Alzheimer’s disease (AD) brain hallmarks, the presence of reactive astrocytes was demonstrated to correlate with neuronal loss and cognitive deficits. Evidence indeed supports the role of reactive astrocytes as mediators of changes in neurons, including synapses. However, the complexity and the outcomes of astrocyte reactivity are far from being completely elucidated. Another key role in AD pathogenesis is played by alterations in brain cholesterol metabolism. Oxysterols (cholesterol oxidation products) are crucial for brain cholesterol homeostasis, and we previously demonstrated that changes in the brain levels of various oxysterols correlate with AD progression. Moreover, oxysterols have been shown to contribute to various pathological mechanisms involved in AD pathogenesis. In order to deepen the role of oxysterols in AD, we investigated whether they could contribute to astrocyte reactivity, and consequently impact on neuronal health. Results showed that oxysterols present in mild or severe AD brains induce a clear morphological change in mouse primary astrocytes, accompanied by the upregulation of some reactive astrocyte markers, including lipocalin-2 (Lcn2). Moreover, astrocyte-conditioned media analysis revealed a significant increase in the release of Lcn2, cytokines, and chemokines in response to oxysterols. A significant reduction of postsynaptic density protein 95 (PSD95) and a concurrent increase in cleaved caspase-3 protein levels have been demonstrated in neurons co-cultured with oxysterol-treated astrocytes, pointing out that mediators released by astrocytes have an impact on neurons. Among these mediators, Lcn2 has been demonstrated to play a major role on synapses, affecting neurite morphology and decreasing dendritic spine density. These data demonstrated that oxysterols present in the AD brain promote astrocyte reactivity, determining the release of several mediators that affect neuronal health and synapses. Lcn2 has been shown to exert a key role in mediating the synaptotoxic effect of oxysterol-treated astrocytes.

Abbreviations: a-EPOX, 5α,6α-epoxycholesterol; β-EPOX, 5β,6β-epoxycholesterol; 24-OHC, 24-hydroxycholesterol; 27-OHC, 27-hydroxycholesterol; 7-KC, 7-Ketocholesterol; 7α-OHC, 7α-hydroxycholesterol; 7β-OHC, 7β-hydroxycholesterol; ApoE, Apolipoprotein E; CCL, C–C motif chemokine; CXCL, C–X–C motif chemokine; DIV, Days in vitro; GC–MS, Gas chromatography-mass spectrometry; G-CSF, Granulocyte colony-stimulating factor; GFAP, Glial fibrillary acidic protein; Il1β, Interleukin 1β; Lcn2, Lipocalin-2; LDH, Lactate dehydrogenase; LPS, Lipopolysaccharide; NTs, Neurofilament tangles; NGAL, Neutrophil gelatinase-associated lipocalin; NMDAR, N-methyl-D-aspartate receptor; PSD, Postsynaptic density; PSD95, Postsynaptic density protein 95; SerpinA3N, Serine protease inhibitor A3N; sICAM-1, Soluble intercellular adhesion molecule-1; sIL-1, Small interfering RNA; TNF-α, Tumor necrosis factor-α.

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1. Introduction

Alzheimer’s disease (AD) is a neurodegenerative disorder, that represents the most common form of dementia and afflicts millions of people worldwide [1]. The pathogenic hallmarks of AD are extracellular deposits of amyloid-β (Aβ) peptides in the form of senile plaques and intracellular neurofibrillary tangles (NFTs) made of hyperphosphorylated tau protein. These lesions are typically accompanied by gliosis, especially surrounding senile plaques [2]. Neuronal death and synapse loss are also observed; in particular, synaptic loss exceeds neuronal loss, meaning that remaining neurons also lose synapses. Synapse loss is indeed the best correlate of cognitive decline [3].

Astrocytes play a crucial role in maintaining brain homeostasis and their reaction to different kinds of insults leads to a heterogeneous range of changes, known as “astrocyte reactivity” [4]. An increase in the number of reactive astrocytes is a typical histopathological feature of AD brain, and it correlates with cognitive decline and neuronal loss also in transgenic mouse models [5,6]. Besides morphological changes and the upregulation of common markers of reactivity (e.g. the glial fibrillary acidic protein, GFAP), reactive astrocytes show significant alterations in gene expression and functions, depending on the specific stimulus [7,8]. However, the outcome of astrocyte reactivity is still somewhat controversial. Several studies highlighted that reactive astrocytes lose neuroprotective functions, including their ability to promote neurite growth, neuronal survival, and synapse formation [9,10]. One of the mechanisms through which astrocytes could have an impact on neurons is by the altered release of different kinds of molecules including cytokines, chemokines, growth factors, and neurotransmitters. For instance, an increase in the release of inflammatory mediators by reactive astrocytes has been shown to affect neuronal viability, tau phosphorylation [9,11], and synaptic function [12,13]. Another mediator released by reactive astrocytes is lipocalin-2 (Lcn2), a protein identified as a pan-reactive inducible factor by activated microglia, reactive astrocytes, neurons, and endothelial cells in response to inflammatory stimuli, infections or other insults [16,17].

In addition to amyloid plaques and NFTs, Alois Alzheimer originally described the presence of “adipose inclusions” in the glial cells of AD brains, suggesting a malfunctioning of lipid metabolism [18]. A clear link between AD and lipid metabolism was established later by the identification of the ε4 allele of apolipoprotein E (ApoE) as a strong genetic risk factor for AD [19]. Moreover, in the last few decades, much other evidence supports a role for lipids, in particular cholesterol, in AD pathogenesis [20–22]. As the main lipid component of neuronal and glial membranes, as well as a key constituent of myelin, cholesterol plays an essential role in synapse formation, maintenance, and function [23,24]. Astrocytes are the main producers of cholesterol in the brain, that is delivered to neurons loaded into ApoE-containing lipoproteins. Brain cholesterol homeostasis is closely controlled by pathways regulating cholesterol biosynthesis, storage, and elimination; in particular, the main process responsible for cholesterol elimination is enzymatic oxidation to oxysterols that are able to cross the blood-brain barrier. Enzymatically produced oxysterols are synthesized by both neurons and glial cells, whereas cholesterol is indeed essentially oxidized to 24-hydroxycholesterol (24-OHC) mainly by neurons. Another enzymatically produced oxidoster, 27-hydroxycholesterol (27-OHC), is mainly produced by glial cells [25] and various oxysterols deriving from cholesterol non-enzymatic oxidation are also present [26,27]. Oxysterols are not only cholesterol metabolites but they also play many regulatory functions, such as modulating cholesterol biosynthesis, inflammatory pathways, and the immune response [28]. We have previously demonstrated a correlation between changes in brain levels of various oxysterols and AD progression. Through analysis of the oxysterol composition in AD and aged-matched control cortex, we observed a significant increase in the levels of various enzymatic (e.g. 27-OHC) and non-enzymatic oxysterols (e.g. 7α-hydroxycholesterol, 7α-OHC; 7β-hydroxycholesterol, 7-β-OHC; 7-ketocholesterol, 7-KC; 5α,6α-epoxycholesterol, α-EPOX; 5β,6β-epoxycholesterol, β-EPOX). In contrast, 24-OHC levels were markedly decreased compared to control brains, probably as a result of neuronal loss [27]. The effects of oxysterols are still controversial but growing evidence suggests that some of them (e.g. 27-OHC, 7-KC, 7α- and 7β-OHC) may play a role in AD pathogenesis [29,30] by inducing oxidative stress, inflammation [31], Aβ accumulation and formation [32,33], tau hyperphosphorylation [34], synaptic dysfunction [35], and cell death [36,37].

At present, data regarding the impact of oxysterols on astrocytes are limited and mostly related to the ability of some of them to affect brain cholesterol synthesis and transport in various astrocytic cell lines [38,39]. 27-OHC was also observed to induce oxidative stress and to downregulate the antioxidant response in C6 glioma cells, leading to cell toxicity [40]. 24-OHC has been shown to affect redox homeostasis in human glial cells, although its impact may depend on the concentration [41], and 7-KC and 7β-OHC exhibited cytotoxic effects in mixed glial murine primary cultures [37]. High concentrations of 7β-OHC have also been observed to induce toxic effects and morphological changes in an in vitro model of reactive astrocytes [42].

Since growing data support the involvement of oxysterols in several aspects of AD pathology and given the presence of reactive astrocytes in the disease, we wanted to investigate whether these compounds could impact on astrocyte reactivity, potentially compromising neuronal health. With the aim of mimicking the oxysterol composition in human AD brain, we used two oxysterol mixtures, both including the main seven oxysterols previously quantified in cortical AD brain samples and representative of early or late stages of the disease [27], to investigate the effect of oxysterols in mouse cortical astrocytes and neuronal cultures.

2. Materials and methods

2.1. Composition of oxysterol mixtures

Cell cultures were treated with two oxysterol mixtures, whose compositions represent oxysterol amounts previously quantified in mild (Early AD mixture) or severe (Late AD mixture) AD brain samples [27]. Both oxysterol mixtures consisted of the same seven oxysterols but in different proportions. The Early AD mixture composition was: 24-OHC (52.9%), 27-OHC (3%), 7-KC (9.2%), 7α-OHC (4.5%), 7β-OHC (19.2%), α-EPOX (3%), and β-EPOX (8.2%). The Late AD mixture composition was: 24-OHC (33.4%), 27-OHC (5.8%), 7-KC (12.7%), 7α-OHC (5.4%), 7β-OHC (23.8%), α-EPOX (4.9%), and β-EPOX (14%). Oxysterols were dissolved in absolute ethanol.

Oxysterols were provided as described: 24-OHC (5275, Medical Isotopes, Pehlam, NH, USA), 27-OHC (700061P), 7-KC (700015P), 7α-OHC (700034P), 7β-OHC (700035P) (Avanti Polar Lipids, Alabaster, AL, USA), α-EPOX (C4130-000), and β-EPOX (C5030-000) (Steraloids, Newport, RI, USA).

2.2. Primary cultures and co-cultures

All animal procedures were in accordance with the European Communities Council Directive (86/609/EEC and 2010/63/EU), the UK Animals (Scientific Procedures) Act 1986, and the Italian Law for Care and Use of Experimental Animals (26/2016), with agreement from both the King’s College London (Denmark Hill) Animal Welfare and Ethical Review Board and the Ethical Committee of the University of Turin.

Primary astrocyte cultures were obtained from cerebral cortex of wild type CD1 mice on postnatal day 1–3 as previously described [43]. Briefly, after dissociation of the cortices, cells were re-suspended in...
 growing medium (DMEM high glucose with glutaMAX, sodium pyruvate, 10% fetal bovine serum, 100 U/mL penicillin, 100 μg/mL streptomycin) and seeded into T75 flasks, previously coated with poly-L-lysine (10 μg/mL) for at least 1 h at 37 °C. Astrocytes were cultured for 7–14 days in a humidified 5% CO₂ incubator at 37 °C, shaking them at 200 rpm overnight on days 3 and 7 to remove microglia and oligodendrocytes. The absence of microglial contamination was assessed by immunochemistry and Western blotting using antibodies that recognize the ionized calcium binding adaptor molecule 1 (Iba1, microglial marker) (Supplemental Fig. 1). Astrocyte-enriched cultures were then re-plated into 12-well plates using trypsin and the medium was changed to Neurobasal serum-free medium (supplemented with 2% B-27, 2 mM glutaMAX, 100 U/mL penicillin, 100 μg/mL streptomycin) 24 h before treatment. Astrocyte cultures were treated with the oxysterol mixtures (Early or Late at different concentrations) up to 24 h. Vehicle (ethanol, 34.2 mM) was added to control astrocytes. 

Primary neurons were obtained from cerebral cortex of wild type CD1 mouse embryos at embryonic day 15, according to a previously published protocol [44]. Briefly, after dissociation, cortices were transfected at 5 × 10⁵ cells per well of 6-well plates previously coated with poly-D-lysine (10 μg/mL) for at least 1 h at 37 °C. After 1 h, neuron adhesion was checked and the medium was changed to Neurobasal serum-free medium. Neuronal cultures were treated at 5–13 DIV with the plasmid peGFP-N1 (Clontech, Mountain View, CA, USA) and imaged live 24 h after treatment. Astrocyte and Western blotting using antibodies that recognize the ionized calcium binding adaptor molecule 1 (Iba1, microglial marker) (Supplemental Fig. 1). Astrocyte-enriched cultures were then re-plated into 12-well plates using trypsin and the medium was changed to Neurobasal serum-free medium (supplemented with 2% B-27, 2 mM glutaMAX, 100 U/mL penicillin, 100 μg/mL streptomycin). The suspension was then filtered using a 40 μm cell strainer, live cells were counted and neurons were plated to a density of 5 × 10⁵ viable cells per well of 6-well plates previously coated with poly-D-lysine (10 μg/mL) for at least 1 h at 37 °C. After 1 h, neuron adhesion was checked and the medium was changed. Cultures were maintained at 37 °C with 5% CO₂ in Neurobasal serum-free medium. Neuronal cultures were checked at 13–14 days in vitro (DIV). For neurite and spine analysis, neurons were transfected at 5–7 DIV with the plasmid pcGFP-N1 (Clontech, Mountain View, CA, USA) using Lipofectamine 2000 (Invitrogen, Thermo Fisher Scientific, Waltham, MA, USA) and imaged live 24 h after treatment at 14 DIV using an Opera Phenix microscope (PerkinElmer, Waltham, MA, USA).

For co-culture experiments, primary astrocytes were plated into cell culture inserts (0.4 μm pore membrane, Falcon, Corning, Corning, NY, USA), that allow the passage of small molecules in culture medium. Astrocytes were pre-treated with the Late AD mixture (10 μM) or vehicle (ethanol) for 12 h, medium was changed to remove oxysterols, and inserts on which astrocytes were grown were added to neuron cultures for a further 24 h.

2.3. Cytotoxicity assay

Cytotoxicity was assessed by measuring levels of lactate dehydrogenase (LDH) enzyme release into the media using Pierce LDH Cytotoxicity Assay Kit (Thermo Scientific, Thermo Fisher Scientific) according to the manufacturers’ instructions. Some cells were lysed with 0.5% Triton X-100 and LDH content in the medium was measured in order to evaluate the maximum LDH amount released from dead cells. LDH release was calculated as a percentage of total LDH released by lysed cells.

2.4. Gel electrophoresis and Western blotting

After treatments, the cell culture media were collected, cells were washed with PBS and directly lysed with PBS containing sample buffer (NuPAGE LDS Sample Buffer 4X, Invitrogen), reducing agent (NuPAGE Sample Reducing Agent 10X, Invitrogen), protease inhibitor (complete Mini EDTA-Free Protease Inhibitor Cocktail, Roche, Basel, CH) and phosphatase inhibitor (PhosSTOP, Roche) cocktails.

Equal amounts of protein samples (20 μg) were boiled, separated by electrophoresis using 10% precast gels (NuPAGE 10% Bis-Tris Protein Gels, Invitrogen) and then transferred to nitrocellulose membranes (Amersham Protran, GE Healthcare, Chicago, IL, USA). After blocking with Odyssey Blocking Buffer (LI-COR Biosciences, Lincoln, NE, USA) for 1 h at room temperature, membranes were incubated with primary antibodies overnight at 4 °C. The following primary antibodies were used: GFAP (Z0334, Dako, Agilent Technologies, Santa Clara, CA, USA), Lipocalin-2/NGAL (AF1857, R&D Systems, Minneapolis, MN, USA), SerpinA3N (AF4709, R&D Systems), cleaved caspase-3 (Asp175, 9661, Cell Signaling, Danvers, MA, USA), PSD95 (D74D3, Cell Signaling), Synapsin (6008-30, BioVision, Milpitas, CA, USA), Iba1 (019-19741, Wako Chemicals, Richmond, VA, USA) and β-actin (ab2286, Abcam, Cambridge, UK). After washing to remove unbound antibody with TBS-Tween20 0.05%, the appropriate fluorophore-coupled secondary antibody (1:10,000, LI-COR Biosciences) was added for 1 h at room temperature. Membranes were washed with TBS-Tween20 0.05% and scanned using an Odyssey infrared imaging system (LI-COR Biosciences). Band intensities were quantified using the Image Studio Software (LI-COR Biosciences) and normalized to the corresponding β-actin bands.

Similarly, to detect secreted Lcn2, 250 μL of culture media were concentrated using centrifugal filters (Amicon Ultra, Millipore, Merck, Darmstadt, DE); the protein concentration was calculated using the Bradford Assay (Protein Assay Dye Reagent Concentrate, Bio-Rad Laboratories, Hercules, CA, USA) and samples were prepared as above to perform Western blotting of cell culture media.

2.5. Immunochemistry

Astrocytes were plated on coverslips (12 or 18 mm diameter, No. 1.5) for 12- or 24-well plates. After treatment, cells were washed with PBS, fixed in 4% paraformaldehyde in PBS for 10 min at room temperature, and then washed again twice with PBS. Cells were permeabilized and blocked (4% goat serum in PBS, Sigma-Aldrich, St. Louis, MO, USA). Cells were images using an Eclipse Ti-E inverted Microscope (Nikon, Tokyo, JP) or LSM800 confocal microscope (Carl Zeiss, Oberkochen, DE).

2.6. Cytokine array

Astrocyte culture media were collected and stored at −20 °C prior to analysis of cytokine and chemokine content using Mouse Proteome Profiler arrays (Mouse Cytokine Array Panel A, R&D Systems), according to the manufacturers’ instructions. Briefly, array membranes were incubated in blocking buffer for 1 h at room temperature. Each sample of medium was incubated with the Detection Antibody Cocktail and this mix was then placed onto blocked membranes overnight at 4 °C. After washing, membranes were incubated with IRDye 800CW Streptavidin (1:2000, LI-COR Biosciences) for 30 min at room temperature. Membranes were then scanned using an Odyssey infrared imaging system and the spot intensities were quantified using the Image Studio Software (LI-COR Biosciences). Positive and negative control spots included in each membrane allowed quantitative analysis by densitometry and results were expressed as percentage change compared to control cultures.

2.7. Oxysterol quantification in astrocyte culture medium

Astrocyte culture medium (100 μL) was added to a screw-capped vial sealed with a Teflon septum together with D7-7α-hydroxycholesterol (50 ng), D7-7β-hydroxycholesterol (50 ng), D7-7α-ketocholesterol (50 ng), D6-5α,6α-epoxycholesterol (50 ng), D6-5β,6β-epoxycholesterol, D6-24-hydroxycholesterol (250 ng), D6-27-hydroxycholesterol (50 ng) as internal standards, as well as 50 μL of butylated hydroxytoluene (5 g/L) and 50 μL of K3-EDTA (10 g/L) to prevent auto-oxidation. Each vial was
then flushed with argon for 5 min to remove air. Alkaline hydrolysis, sterol extraction, and gas chromatography-mass spectrometry (GC-MS) analysis were performed as previously described [45].

2.8. RNA extraction and real-time RT-PCR

Total RNA was extracted using TriFast reagent (Eurogold TriFast, EuroClone, Pero, IT) following the manufacturers’ instructions. RNA was dissolved in RNase-free water with RNase inhibitors (SUPERase-In RNase inhibitor, Invitrogen). The amount and purity of the extracted RNA were assessed by using a NanoDrop ND-1000 spectrophotometer (Thermo Fisher Scientific). cDNA was synthesized by reverse transcription (Thermo Fisher Scientific) following the manufacturers’ instructions.

Singleplex real-time RT-PCR was performed on 30 ng of cDNA by using TaqMan Gene Expression Assays for mouse Lcn2 (Mm01324470_m1) and β-actin (Mm02619580_g1), TaqMan Fast Universal PCR Master Mix, and a 7500 Fast Real-Time PCR System (Applied Biosystems). The PCR cycling parameters were set up as previously described [33]. The fractional cycle number (Ct) was determined for each gene considered and results were then normalized to β-actin expression. Relative quantification of target gene expression was achieved with a mathematical method [46].

2.9. Transient gene knockdown

The transient knockdown of Lcn2 gene was performed by using a Small Interfering RNA (siRNA) (Silencer Select Pre-designed siRNA S69122, Ambion, Thermo Fisher Scientific) following the manufacturers’ instructions. A non-targeting siRNA was used as a negative control (Silencer Select Negative Control #2 siRNA, Ambion). Briefly, astrocytes were plated into 12- or 24-well plates and respectively 50 μl or 100 μl of a mix containing siRNA and transfection agent (Lipofectamine RNAiMAX Reagent, Invitrogen) in a 1:1 ratio was added to each well, in order to reach a final siRNA concentration of 100 nM and 3 μl/ml of transfection agent. After 6 h transfection, the medium was changed and astrocytes were treated with the Late AD mixture (10 μM) for 12 h. After treatment, cells were washed with PBS and fresh medium was added. Astrocyte conditioned media were then collected after a further 24 h. Protein and RNA extraction, as well as real-time RT-PCR, were performed as described above. The silencing efficiency, validated by real-time RT-PCR, was approximately 85% (Supplemental Fig. 2).

2.10. Analysis of neuronal morphology

The neuronal morphology of 14 DIV neurons previously transfected with the plasmid pEGFP-N1 (Clontech) on 5–7 DIV and treated on 13 DIV was assessed using high resolution digital images of live neurons taken using an Opera-Phenix microscope (PerkinElmer). Neurite complexity was analysed using Harmony software: total and maximum neurite length, and number of nodes and extremities were quantified and compared among groups. NeuronStudio software (CNIC, Mount Sinai School of Medicine) was used for dendritic spine analysis. Spine density was defined as number of spines per micrometer of dendrite length. Dendritic spine densities were calculated from 20 neurons/condition.

2.11. Statistical analysis

After performing a Shapiro-Wilk normality test, data were analysed using one-way ANOVA followed by Bonferroni post hoc test or Student’s t-test (GraphPad Prism 7 Software, Graphpad Software, La Jolla, CA, USA). Results were considered statistically significant when P < 0.05. Data are represented as means ± standard deviation (SD).

3. Results

3.1. Oxysterol mixtures induce a morphological change in astrocytes

Astrocyte reaction to certain stimuli is defined by a wide range of heterogeneous responses, including changes in morphology [7,47]. To analyse the effect of oxysterols on astrocyte morphology and survival, cells were exposed to increasing concentrations of the Early and Late mixtures (1, 5 or 10 μM). Exposure of primary astrocytes to 5 or 10 μM of both oxysterol mixtures for 24 h induced a clear change in their morphology with astrocytes adopting a more stellate “brain-like” and reactive appearance rather than the large, flat fibroblast-like morphology typical of unstimulated dissociated astrocytes in culture (Fig. 1B). Importantly, this was not accompanied by cell death since there were no significant increases in the abundance of LDH released into culture medium following treatment when compared to control cultures (Fig. 1A). Time course treatments showed that astrocytes started to display the typical signs of reactivity (increase in the number of GFAP positive processes and stellate appearance) after 12 h of exposure to 10 μM of both Early and Late mixtures, with these changes becoming more evident 24 h after treatment (Fig. 2).

3.2. Oxysterol mixtures increase the synthesis of the pan-reactive astrocyte markers Lcn2 and SerpinA3N

To validate the induction of astrocytic reactivity by oxysterols and to confirm that the morphological alteration is accompanied by functional changes, we determined whether treatment with the Early or Late AD mixtures led to an increase in the levels of some pan-reactive astrocyte markers [8]. Exposure to both oxysterol mixtures (10 μM) resulted in a marked increase in Lcn2 and serum protease inhibitor A3N (SerpinA3N) protein levels 12 h (P < 0.0001 and P < 0.001) and 24 h (P < 0.0001) after treatment (Fig. 3), contemporaneous with morphological changes. These molecules are involved in the acute-phase response and were both identified as new reactive astrocyte markers [8,48]. Total levels of GFAP were only slightly increased (P < 0.05) or remained unchanged in response to the mixtures (Fig. 3), suggesting that the observed morphological changes are not only dependent on the expression of GFAP.

3.3. Oxysterol-induced reactivity triggers the release of Lcn2, cytokines, and chemokines by astrocytes

Astrocyte reactivity is typically characterized by the release of cytokines, chemokines, and other mediators, that could have several effects on the surrounding cells [49]. Therefore, we investigated whether oxysterol-mediated astrocyte reactivity was accompanied by altered release of mediators into the cell culture medium.

Since Lcn2 is secreted by reactive astrocytes [17,50], we analysed its protein levels in astrocyte conditioned media (ACM) by Western blotting. We saw that the robust increase in intracellular levels of Lcn2 is accompanied by a significant increase in the secretion of Lcn2 after 24 h treatment with 10 μM of both Early and Late oxysterol mixtures (P < 0.0001) (Fig. 4A).

The amounts of 40 cytokines and chemokines in ACM samples were next evaluated using Proteome Profiler antibody arrays. The analysis showed a significant increase in the release of several mediators by astrocytes treated for 24 h, especially with the Late AD oxysterol mixture (10 μM). Some of the cytokines found to be increased in the medium of astrocytes treated with this oxysterol mixture were interleukin-1β (IL-1β), soluble intercellular adhesion molecule-1 (sICAM-1) (P < 0.001), IL-1α, IL-5, granulocyte colony-stimulating factor (G-CSF), and C–C motif chemokine 12 (CCL12) (P < 0.01). In addition, the levels of CCL1, IL-7, IL-10, IL-13, IL-16, IL-17, and C–C–X–C motif chemokine 9 (CXCL9) were slightly but significantly increased compared to levels in control media (P < 0.05) (Fig. 4B and C). In contrast, when astrocytes were...
exposed to the Early AD oxysterol mixture (10 μM), only sICAM-1 ($P < 0.01$), IL-1α, and IL-1β ($P < 0.05$) were found to be significantly increased relative to levels in control ACM, highlighting a different impact of the two oxysterol mixtures on release of cytokines and chemokines by astrocytes (Fig. 4C).

3.4. Oxysterol-stimulated astrocytes compromise neuronal health

It has previously been shown that astrocyte reactivity depends on the specific stimulus and that heterogeneous responses of reactive astrocytes involve different signaling pathways. The consequences of these alterations for astrocytic support of neuronal health require further investigation [7,48].

To test the effect of oxysterol-stimulated astrocytes on neuronal health, we performed co-culture experiments. GC-MS analysis of oxysterols in ACM showed that, although variable, a substantial proportion (8–54%) of some oxysterols remain in ACM 24 h after treatment (Supplemental Fig. 3). Since these oxysterols are known to directly affect neuron viability [33,35,51], we co-cultured neurons with astrocytes that had been pre-treated with oxysterols. As shown in Fig. 5A, astrocytes grown on cell culture inserts were pre-treated for 12 h with the Late AD oxysterol mixture (10 μM) and then transferred to culture plates containing neurons for a further 24 h; this allowed us to investigate the effect of mediators released from stimulated astrocyte on neurons, without transferring oxysterols that remained in ACM. In parallel, neuronal cultures were treated with the Late AD mixture or with ethanol.
for 24 h in the absence of astrocytes, in order to assess the direct effect of the same oxysterol mixture on neurons.

Interestingly, Western blotting analysis showed a significant reduction of postsynaptic density protein 95 (PSD95) levels (P < 0.05), a scaffold protein important for postsynaptic density structure and function, in neurons cultured with astrocytes that had been previously exposed to oxysterols (Fig. 5B). This reduction was accompanied by a significant increase in the amount of cleaved (active) caspase-3 (P < 0.01) (Fig. 5B), that was demonstrated to play also various non-apoptotic roles in neurons, including to affect synaptic functions [52]. A mild but not significant reduction of the pre-synaptic marker synapsin was also observed. Moreover, when neurons were directly treated with oxysterols, PSD95 (P < 0.05), cleaved caspase-3 (P < 0.01), and synapsin levels (P < 0.05) were all significantly altered (Fig. 5B), confirming previous results showing that oxysterols directly compromise neuron health [33,35,53]. Importantly, all these results were obtained in the absence of significant neuronal death, as assessed by LDH release (< 0.05), cleaved caspase-3 (P < 0.01), and synapsin levels (P < 0.05) were all significantly altered (Fig. 5B), confirming previous results showing that oxysterols directly compromise neuron health [33,35,53]. Importantly, all these results were obtained in the absence of significant neuronal death, as assessed by LDH release (Fig. 5B). Overall, these data suggest that factors released by astrocytes in response to the oxysterol mixture, whose composition is similar to AD brain oxysterol content, compromise synaptic and neuronal health without causing overt neurotoxicity.

3.5. Lcn2 secreted by oxysterol-stimulated astrocytes affects neurite complexity and decreases dendritic spine density

Lcn2 has previously been shown to affect synapses, neuronal health and morphology [17,54,55]. To investigate whether Lcn2 could play a role in mediating the effect of oxysterol-stimulated astrocytes on neurons, Lcn2 expression was silenced in astrocytes prior to determining the impact of ACM on neuron morphology. To further investigate the effects of ACM on synapses under these conditions, dendritic spine density was also evaluated.

For this purpose, astrocytes were transfected for 6 h with Lcn2 or scrambled siRNA, and the medium removed prior to treatment with the Late AD mixture (10 μM) for 12 h. The medium was removed and fresh medium was added for a further 24 h, to obtain ACM without oxysterols. Both real-time RT-PCR (Fig. 6A) and Western blotting results (Fig. 6B) confirmed that siRNA-mediated Lcn2 gene silencing prevents the increase in Lcn2 expression, synthesis, and release into the medium induced by the Late AD oxysterol mixture, compared to control conditions (scrambled siRNA). Interestingly, oxysterol-induced morphological changes were not prevented by Lcn2 gene silencing, suggesting that the pathway by which oxysterols induce astrocyte alterations is not dependent upon the presence of Lcn2 (Fig. 6C).

Next, ACM was added to cultured neurons to determine the impact of secreted Lcn2 on neuron morphology and dendritic spines. Medium from oxysterol-activated astrocytes transfected with the scrambled siRNA compromises neurite complexity, as shown by reduction of total and maximum neurite length, and the number of nodes, automatically identified by Harmony software (P < 0.05) (Fig. 7A). Importantly, the addition of ACM was also found to substantially reduce dendritic spine density (P < 0.001) (Fig. 7B), further indicating a synaptotoxic effect of proteins secreted by oxysterol-stimulated astrocytes. All of these effects were prevented by silencing Lcn2, confirming a key role of Lcn2 in mediating the impact of oxysterol-stimulated astrocytes on neurite complexity and synaptic health (Fig. 7).

4. Discussion

The pathophysiology of AD is still not fully understood, but evidence supports the involvement of many factors in sporadic AD onset, including inflammation and impaired cholesterol metabolism [56]. Microglia and astrocytes play a role in the neuroinflammation observed in AD and they are found in their activated state in affected brain regions, particularly surrounding amyloid plaques. Astrocyte reactivity is not an exclusive feature of AD brain but the association of reactive astrocytes with amyloid plaques is considered functionally significant. Indeed, post mortem neuropathological studies have shown that the number of reactive astrocytes increases with disease progression [6,57], and activated astrocytes and microglia correlate with dementia in AD [5].

Astrocyte reactivity leads to a heterogeneous range of phenotypic, transcriptomic, and functional changes, depending on the specific environmental stimulus [7,8], including those related to ageing and...
Fig. 3. Oxysterol mixtures increase the synthesis of pan-reactive astrocyte markers. The glial fibrillary acidic protein (GFAP), serine protease inhibitor A3N (SerpinA3N), and lipocalin-2 (Lcn2) protein levels were determined by Western blotting of lysates from primary astrocytes treated with 10 μM of the Early or Late AD oxysterol mixture for up to 24 h. The amounts of proteins of interest were normalized to β-actin levels in the same sample and are represented as percentage of average control values. Data are expressed as mean values ± SD of three different experiments (one-way ANOVA). ****P < 0.0001, ***P < 0.001, *P < 0.05 vs. control.
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Fig. 4. Oxysterol treatment increases the release of lipocalin-2 (Lcn2), cytokines and chemokines by astrocytes. Astrocyte cultures were treated with the Early or Late AD oxysterol mixture (10 μM) for 24 h. (A) Lcn2 protein levels in astrocyte conditioned media (ACM) were examined by Western blotting. Data are expressed as mean values ± SD from three different experiments and are shown as percentage change from average control values (one-way ANOVA). ****P < 0.0001 vs control. (B) Mouse cytokine profiler antibody arrays were used to detect the amounts of 40 mediators in ACM. Representative images of array membranes are shown, highlighted some of the cytokines found to be significantly increased in ACM from treated cells. The bar chart shows the cytokines significantly affected by oxysterol mixture treatments as percentage change from control. Data are expressed as mean values ± SD from three different experiments (one-way ANOVA). ***P < 0.001, **P < 0.01, and *P < 0.05 vs control.

disease [10,58]. Here, we observed that application of various oxysterols, in the same proportions as those identified in early and late stage AD brain, distinctly altered astrocyte morphology after 24 h treatment. In particular, astrocytes adopted a more stellate appearance with many long and branched processes evident. In support of these data, 7β-OHC has previously been shown to induce morphological changes, characterized by process elongation, in cultured reactive astrocytes [42]. An increase in the number of processes is considered a reliable feature of reactivity, demonstrated in several cell culture models of reactive astrocytes [59,60].

Reactive astrocytes were conventionally identified by GFAP immunoreactivity; however, it is noteworthy that different stimuli can lead to similar degrees of GFAP upregulation while causing substantially different changes in transcriptome profile and cell functions [7].
Furthermore, unlike brain astrocytes, most astrocytes in culture are GFAP reactive as a result of the in vitro conditions, mainly the absence of their usual environment and the loss of inhibitory influences of neurons [42]. Astrocytes treated with Early or Late AD oxysterol mixture up to 24 h showed a significant but mild increase in GFAP protein levels despite the extensive morphological changes observed. The morphological changes we observed in response to oxysterol mixtures were, however, accompanied by a marked upregulation of Lcn2 and SerpinA3N, both proteins identified as “pan-reactive” astrocytes markers [8], as well as by increased release of Lcn2 into culture medium. Lcn2, also known as neutrophil gelatinase-associated lipocalin (NGAL), is an acute phase protein with several roles [14] that is secreted by different kinds of cells, including reactive astrocytes [16,17]. For instance, it has recently been shown that Aβ induces Lcn2 production in astrocytes [61]. It can act as an autocrine mediator of reactive astrogliosis, since it is able to induce morphological changes in primary astrocytes [59]. In addition, several studies showed that Lcn2 exerts synaptotoxic and neurotoxic effects, affecting neuronal viability, dendritic spine density and...
morphology, and ultimately cognitive functions [17,50,54,55]; for this reason, we decided to focus mainly on the effects mediated by astrocyte-produced Lcn2 on neuronal health. SerpinA3N, also known as alpha1-antichymotrypsin, is a protease inhibitor involved in the acute phase response and reactive and ageing astrocytes are its main producers in the brain [8,58]. Immunohistochemical analysis of AD brain samples showed that SerpinA3N co-localizes with amyloid plaques, reactive microglia and astrocytes [62].

It is well known that in many cases astrocyte activation is accompanied by an increase in the release of several soluble factors, including reactive oxygen species and cytokines [11,49]. Astrocytes are both targets and effectors of cytokines and many other mediators that affect not only their immune and inflammatory functions but also their synapse-directed and neuronal functions [49]. Interestingly, 7α, 25-dihydroxycholesterol has been shown to inhibit the lipopolysaccharide (LPS)-induced release of pro-inflammatory cytokines, including IL-17/tumor necrosis factor-α (TNF-α), in cultured human astrocytes [63]. Other oxysterols (e.g. 27-OHC) have been demonstrated to decrease LPS-induced expression of IL-6 and TNF-α mRNA in mouse primary glial cells [64]. However, there are no data on how oxysterols directly affect astrocyte secretion profile. Our cytokine array analysis showed that oxysterol treatment significantly increases the release of a wide range of cytokines and chemokines by primary astrocytes after 24 h. Interestingly, this analysis highlighted a differential impact of the two oxysterol mixtures; the majority of the significant increases were indeed observed in ACM from astrocytes treated with the Late AD oxysterol mixture, likely because of the different composition of this mixture. Moreover, even if both mixtures showed a similar effect on astrocyte

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**Fig. 6.** Lipocalin-2 (Lcn2) gene silencing prevents Lcn2 upregulation and release into the medium. Astrocyte cultures were transfected for 6 h with Lcn2 or scrambled siRNA and then treated with the Late AD oxysterol mixture 10 μM (Mix) for 12 h. After treatment, the medium was changed and astrocytes were incubated with fresh medium for 24 h. Transient Lcn2 gene knockdown was evaluated by (A) real-time RT-PCR and (B) Western blotting of both lysates and astrocyte conditioned media (ACM) samples. Data were normalized to the corresponding β-actin levels. Data are expressed as mean values ± SD from three different experiments as percentage change from control (one-way ANOVA). ****P < 0.0001, ***P < 0.001, and *P < 0.05 vs control; ####P < 0.0001 vs oxysterol treated. (C) Astrocyte morphology was examined by immunocytochemistry using a glial fibrillary acidic protein (GFAP) antibody (red) and nuclei were stained with Hoechst 33258 (blue). Representative images from three experiments are shown. Cells were imaged using an LSM800 confocal microscope (Zeiss, 40× objective; scale bar: 100 μm).
Fig. 7. Lipocalin-2 (Lcn2) secreted by oxysterol-stimulated astrocytes affects neurite complexity and decreases dendritic spine density. Primary neurons were incubated for 24 h with conditioned media from astrocytes transfected for 6 h with Lcn2 or scrambled siRNA and then treated with the Late AD oxysterol mixture 10 μM for 12 h. After treatment, the medium was changed and astrocytes were incubated with fresh medium for 24 h. High resolution digital images of live neurons were obtained using an Opera-Phenix microscope (PerkinElmer). (A) Neurite complexity was analysed using Harmony software. Total and maximum neurite length, number of nodes and extremities were quantified and compared among groups. Representative images are shown (scale bar: 100 μm). Data are expressed as mean values ± SD from three different experiments and were obtained from analysis of 20 neurons/condition (one-way ANOVA). *P < 0.05 vs control. (B) NeuronStudio software was used for dendritic spine analysis. Spine density was defined as number of spines per micrometer of dendrite length and was obtained from analysis of 20 neurons/condition. Representative images are shown (scale bar: 25 μm). Data are expressed as mean values ± SD from three different experiments (one-way ANOVA). ***P < 0.001 vs control.
Among the released mediators, inflammatory cytokines (IL-1α, IL-1β, and IL-17), adhesion molecules (sICAM-1), growth factors and chemokines (IL-5, IL-7, IL-16, G-CSF, CCL1, CCL12, and CXCL9), immunoregulatory cytokines (IL-5 and IL-13), and anti-inflammatory cytokines (IL-10) were elevated. In particular, IL-1β and sICAM-1 were the most significantly increased. IL-1β has been previously shown to induce reactive astrogliosis [10], to be released by reactive astrocytes and astrocytes from 5xFAD AD mouse model [65,66], and to contribute to astrocyte-mediated neuronal death [67]; moreover, its levels are increased in AD brains [27,68]. ICAM-1 is a transmembrane glycoprotein belonging to the immunoglobulin family of adhesion molecules, expressed not only by endothelial and immune system cells [69] but also by astrocytes [8]. sICAM-1 is released by proteolytic cleavage of the transmembrane protein [66,70]; its levels are increased in AD cerebrospinal fluid and correlate with various AD markers including total tau, phosphorylated tau, and cortical thinning [71,72]. The data presented here provide the first evidence for a direct impact of oxysterols on astrocyte morphology, markers of reactivity, and secretion profile, and strongly suggest that oxysterol mixtures mimicking oxysterol composition in AD brain induce astrocyte reactivity.

The functional implications of astrocyte reactivity are various and very context-dependent. For instance, astrocytes have been shown to mediate Aβ-induced neurotoxicity and tau phosphorylation, as well as to induce a decrease of synaptic markers; interestingly, the release of pro-inflammatory cytokines from astrocytes seems to play a key role in some of these processes [11,73]. To investigate the consequences of oxysterol-mediated astrocytic reactivity on neurons, we exposed neurons to astrocytes that had previously been challenged with oxysterols. Our data indicated that neurons co-cultured with pre-treated astrocytes are characterized by a significant reduction in PSD95 protein levels and by a clear increase in cleaved caspase-3 protein levels, suggesting that oxysterol-induced reactivity is detrimental for neuronal health. Importantly, similar changes were observed when neuronal cultures were directly treated with the Late AD mixture in the absence of astrocytes, confirming that oxysterols have a direct negative effect on neuronal health. Then, these results suggest that brain oxysterols could affect neuronal health both directly and by inducing astrocyte reactivity. PSD95 is a scaffold protein abundant in the postsynaptic density (PSD), that anchors N-methyl-D-aspartate receptor (NMDAR) as well as other proteins (e.g. potassium channels, cell adhesion molecules) to the PSD. It is also involved in the assembly of specific signaling proteins which act downstream of NMDAR [74]. PSD95 is recognized to play an important role in synaptic plasticity and memory, and in keeping with this its levels decrease in AD [5]. To the best of our knowledge, no studies have yet assessed the specific actions of oxysterol-treated astrocytes on PSD95 levels, however various mediators released by oxysterol-treated astrocytes have previously been shown to affect synapses, including Lcn2 and IL-1α/β [54,75,76]. Indeed, the binding of cytokines and chemokines to their receptors leads to the activation of diverse signaling pathways [77], some of which are involved in PSD95 regulation [78]. Concerning oxysterols, it has been demonstrated that 27-OHC reduces dendritic spine density and PSD95 levels in primary hippocampal neurons [35], whereas 24-OHC affects synaptic plasticity via modulation of NMDAR [79]. Since oxysterols are able to directly modulate several pathways involved in PSD95 regulation [31], other oxysterols present into the mixture could be involved in the observed PSD95 reduction, even if their role needs further investigation. Moreover, the ability of oxysterols to alter membrane structure could affect transmembrane protein localization and function, thus modifying the integrity of the PSD structure [80,81]. The increase in the levels of cleaved caspase-3 fragment further indicates that oxysterol-activated astrocytes are detrimental to neurons. Besides being an effector of apoptosis, caspase-3 was demonstrated to play various non-apoptotic roles in neurons involved in both physiological processes and neurodegenerative diseases [52]. For instance, it is involved in the pathological cleavage of tau protein [82] and its activation correlates with dendritic spine loss and cognitive decline in the absence of neuronal cell death [83]. Interestingly, it has previously been shown that increase in neuronal cleaved caspase-3 levels in response to Aβ is exacerbated in the presence of reactive astrocytes, and that increased caspase-3 activity is reduced upon pre-treatment with the anti-inflammatory molecule minocycline in vitro [11] and in vivo [84]. These and other data highlight that neuronal caspase-3 can be activated by extracellular inflammatory mediators. Caspase-3 can also be activated by oxysterols, mainly those oxidized at C7 position but also others [35,51,85–87].

In AD, synaptic loss is the parameter that best correlates with dementia [88,89]; therefore, elucidating how synapses are lost is of great importance. This can be examined in vitro by monitoring dendritic spine density and/or neurite complexity. The direct effect of oxysterols on synaptic health has been partially investigated as stated above, but we were interested in assessing how oxysterol-activated astrocytes affect synapses. Our results indicate that ACM from oxysterol-activated astrocytes is synaptotoxic, as shown by significant decreases in the number of dendritic spines and reduced complexity of the neurites in primary neurons, as well as by PSD95 reduction as described above. Since Lcn2 is secreted upon oxysterol-induced astrocyte activation and has proven to be neurotoxic [17,55], we tested whether we could revert the synaptotoxic effect of reactive astrocytes by suppressing Lcn2 expression and therefore reducing its secretion. Our results clearly demonstrate that dendritic spine density and neurites architecture are better preserved when Lcn2 expression is silenced prior to treating astrocytes with oxysterols. These data reveal a major role for Lcn2 in mediating the synaptotoxic effect of oxysterol-treated astrocytes. These findings extend previous studies showing that Lcn2 affects dendritic spine density and morphology [54], as well as contributes to hippocampal damage and cognitive impairment in a mouse model of vascular dementia [50].

5. Conclusion

Overall, these results describe the complexity of oxysterol effects on astrocytes and neurons. Of interest was our observation that astrocytes respond most strongly to the oxysterol mixture representative of late AD brain in comparison to early AD-mimicking mixture. These data add to an emerging consensus that astrocyte responses are differentially altered at specific disease stages in response to various changes in the local environment. Indeed, we demonstrated that oxysterols induce a clear morphological change in astrocytes, that is accompanied by an increase in some reactive astrocyte markers and the release of several mediators. We also demonstrated that oxysterols are detrimental to neuronal health but do not cause overt neurotoxicity. Importantly, we have shown that oxysterol-activated astrocytes induce synaptotoxicity that is mediated by Lcn2. This study thus reveals new aspects of brain oxysterol effects on astrocytes and neurons, confirming their potential ability to contribute to AD pathogenesis, and providing support for further investigations into a potential role for Lcn2 as a novel therapeutic target in AD.

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Declaration of competing interest

The authors declare that they have no competing interests.

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Appendix A. Supplementary data

Supplementary data to this article can be found at https://doi.org/10.1016/j.redox.2020.101837.

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