

Pathogenesis of Alzheimer's disease: Involvement of the choroid plexus

Maria Čarna¹ | Isaac G. Onyango¹ | Stanislav Katina^{1,2} | Dušan Holub³ |
Jan Sebastian Novotny¹ | Marketa Nezvedova⁴ | Durga Jha⁴ | Zuzana Nedelska^{1,5} |
Valentina Lacovich¹ | Thijs Vande Vyvere⁶ | Ruben Houbrechts⁶ |
Krystine Garcia-Mansfield⁷ | Ritin Sharma⁷ | Victoria David-Dirgo⁷ |
Martin Vyhnaek^{1,5} | Kateřina Texlova¹ | Hernan Chaves⁸ | Nadine Bakkar⁹ |
Lucia Pertierra⁸ | Mojmir Vinkler² | Hana Markova^{1,5} | Jan Laczko^{1,5} |
Kateřina Sheardova^{1,10} | Marcela Hortova-Kohoutkova¹ | Jan Frič^{1,11} |
Giancarlo Forte¹ | Petr Kaňovsky¹² | Silvie Belaškova¹ | Jiří Damborsky^{1,4} |
Jakub Hort^{1,5} | Nicholas T. Seyfried^{13,14,15} | Robert Bowser⁹ | Gustavo Sevlever⁸ |
Robert A. Rissman¹⁶ | Richard A. Smith¹⁷ | Marian Hajduch³ | Patrick Pirrotte^{7,18} |
Zdenek Spacil⁴ | Eric B. Dammer^{13,14} | Clara Limbäck-Stokin^{19,20} |
Gorazd B. Stokin^{1,21,22}

¹International Clinical Research Centre, St. Anne's University Hospital, Brno, Czech Republic

²Institute of Mathematics and Statistics, Masaryk University, Brno, Czech Republic

³Institute for Molecular and Translational Medicine, Faculty of Medicine and Dentistry, Palacky University Olomouc, Olomouc, Czech Republic

⁴RECETOX Centre, Faculty of Sciences, Masaryk University, Brno, Czech Republic

⁵Memory Clinic, Department of Neurology, 2nd Faculty of Medicine, Charles University and Motol University Hospital, Prague, Czech Republic

⁶Icometrix, Leuven, Belgium

⁷Collaborative Center for Translational Mass Spectrometry, Translational Genomics Research Institute, Phoenix, Arizona, USA

⁸FLENI, Buenos Aires, Argentina

⁹Department of Neurobiology, Barrow Neurological Institute, Phoenix, Arizona, USA

¹⁰1st Department of Neurology, St. Anne's University Hospital and Faculty of Medicine, Masaryk University, Brno, Czech Republic

¹¹Institute of Hematology and Blood Transfusion, Prague, Czech Republic

¹²Department of Neurology, Faculty of Medicine and Dentistry, Palacky University Olomouc and Research and Science Department, University Hospital Olomouc, Olomouc, Czech Republic

¹³Center for Neurodegenerative Disease, Emory University School of Medicine, Atlanta, Georgia, USA

¹⁴Goizueta Alzheimer's Disease Research Center, Emory University, Atlanta, Georgia, USA

¹⁵Departments of Biochemistry and Neurology, Emory University School of Medicine, Atlanta, Georgia, USA

¹⁶Department of Neurosciences, University of California San Diego, La Jolla, California, USA

¹⁷Center for Neurologic Study, La Jolla, California, USA

¹⁸Mass Spectrometry & Proteomics Core Facility, City of Hope Comprehensive Cancer Center, Duarte, California, USA

¹⁹Department of Cellular Pathology, Imperial College Healthcare NHS Trust, London, UK

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²⁰Imperial College London, Faculty of Medicine, London, UK

²¹Division of Neurology, University Medical Centre, Ljubljana, Slovenia

²²Translational Aging and Neuroscience Program, Mayo Clinic, Rochester, Minnesota, USA

Correspondence

Gorazd B. Stokin, International Clinical Research Centre, St. Anne's University Hospital, Brno, Czech Republic
Email: gbstokin@alumni.ucsd.edu

Present address

Clara Limbäck-Stokin, Neuropathology and Ocular Pathology Department, Oxford University Hospitals NHS Foundation Trust, Oxford, UK

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Abstract

The choroid plexus (ChP) produces and is bathed in the cerebrospinal fluid (CSF), which in aging and Alzheimer's disease (AD) shows extensive proteomic alterations including evidence of inflammation. Considering inflammation hampers functions of the involved tissues, the CSF abnormalities reported in these conditions are suggestive of ChP injury. Indeed, several studies document ChP damage in aging and AD, which nevertheless remains to be systematically characterized. We here report that the changes elicited in the CSF by AD are consistent with a perturbed aging process and accompanied by aberrant accumulation of inflammatory signals and metabolically active proteins in the ChP. Magnetic resonance imaging (MRI) imaging shows that these molecular aberrancies correspond to significant remodeling of ChP in AD, which correlates with aging and cognitive decline. Collectively, our preliminary *post-mortem* and *in vivo* findings reveal a repertoire of ChP pathologies indicative of its dysfunction and involvement in the pathogenesis of AD.

KEYWORDS

aging, Alzheimer's disease, cerebrospinal fluid, choroid plexus, pathology

Highlights

- Cerebrospinal fluid changes associated with aging are perturbed in Alzheimer's disease
- Paradoxically, in Alzheimer's disease, the choroid plexus exhibits increased cytokine levels without evidence of inflammatory activation or infiltrates
- In Alzheimer's disease, increased choroid plexus volumes correlate with age and cognitive performance

1 | NARRATIVE

1.1 | Contextual background

Alzheimer's disease (AD) manifests clinically with behavioral changes and progressive decline in cognitive faculties¹ and is by far the most common cause of dementia in the elderly.² Pathologically, AD brains exhibit synaptic and neuronal loss³ accompanied by reactive glia.^{4–7} Senile plaques and neurofibrillary changes are regarded as the disease-characterizing neuropathological hallmarks.⁸ However, these hallmarks are observed also in people who are cognitively healthy^{9,10} or suffer from different neurodegenerative disorders,¹¹ while lesions typical of other disorders such as α -synuclein inclusions frequently populate AD brains.¹² Despite this clinical and neuropathological spectrum of the Alzheimer's syndrome,¹³ accumulating evidence suggests that the most proximal neurobiological event underlying clinical fea-

tures involves impaired brain connectivity due to inadequate synaptic performance.^{14–17} Accordingly, there is significant interest in understanding the neurobiology of synaptic failure in AD¹⁸ and in particular its intimate relationship with neuroinflammation.^{17,19} Since the choroid plexus (ChP) plays a pivotal role in brain homeostasis including its immune status,^{20–22} we here preliminarily investigate structural health of the ChP and provide early evidence of complex ChP pathology indicative of its dysfunction in patients with AD.

The discovery that genetic defects segregating with kindreds afflicted by familial AD (FAD)²³ perturb formation of amyloid- β peptides ($A\beta$), the major constituents of senile plaques,²⁴ paved the way to the amyloid hypothesis. This posits that aberrant $A\beta$ underlie the pathogenesis of AD.²⁵ Although clinical trials have failed to provide definitive support for the amyloid hypothesis,²⁶ the recognition that FAD mutations impair numerous cellular processes ranging from endosomal activity to axonal transport^{27,28} indicates that, in the

foreseeable future, the mechanisms by which FAD mutations cause AD will ultimately be deciphered. The vast majority of AD cases, however, are sporadic in nature and mechanisms at play go beyond those underlying the pathogenesis of FAD. The major risk factor of sporadic AD is aging,²⁹ which promotes a pro-inflammatory state³⁰ also in the aging brain.^{31,32} Coincidentally, key molecules, risk factors, and physiological processes implicated in the pathogenesis of AD including A β assemblies,³³ tau aggregates within and beyond neurofibrillary changes,^{34–37} apolipoprotein E (ApoE),^{38,39} TREM2,^{40–42} and the blood-brain barrier (BBB)^{43,44} all interact with glial lineages, which mediate the pro-inflammatory state and contribute to synaptic failure including synaptophagy in AD brains.^{19,45,46} The finding that systemic inflammatory signals correlate with cognitive decline^{47,48} and regularly communicate and shape epigenetic programs of the glia in the brain⁴⁹ suggests that pro-inflammatory changes in AD brains can be set off also by peripheral causes. Considering ChP coordinates the crosstalk of the immune signals between the periphery and the brain,^{22,50–56} these reports suggest that ChP dysfunction could contribute to the inflammatory state of AD brains.

The ChP is a grape-like structure composed of a tight-junction-bound epithelium and the underlying stroma rich in fenestrated capillaries of peripheral vascular origin. This specific architecture creates a barrier between the peripheral blood and the cerebrospinal fluid (CSF), which circulates between the ventricles and the surface of the brain.^{21,57,58} Through production of the CSF, ChP monitors and adjusts brain homeostasis³⁹ including the activation status of the brain immune system,^{20,53} proliferation of neural stem cells⁵⁹ and cognitive behavior.^{50,60,61} Aging results in profound structural and functional changes of the ChP in animals^{22,50} and humans^{62–64} alike with ChP for unknown reasons becoming significantly increased in volume.⁶⁵ Changes in the ChP have been extensively described also in animal models of AD,^{52,61,66–68} while strikingly few studies to date examine ChP in patients with AD. These studies show altered structural integrity,⁶⁴ describe transcriptomic and proteomic profiles indicative of inflammatory changes and impaired metabolism,^{62,67,69} and report preliminary correlations between ChP volumes, cognition, and biomarkers of AD.^{70,71} Studies of the CSF also reveal extensive abnormalities including inflammatory changes in the CSF in aging^{72–74} and AD^{74–81} and further support the role of ChP in AD. The basis for these abnormalities is largely unknown. To address these gaps in our knowledge and to critically test for the involvement of the ChP in AD, we performed a series of pilot experiments to tease apart the effects of aging *versus* AD in the CSF and then systematically examine the structure of ChP in patients with AD.

1.2 | Study design and main results

We here ask what changes in the CSF environment surround ChP, systematically test for structural, inflammatory and metabolic changes of the *post-mortem* ChP and last investigate ChP *in vivo* in patients with AD. To answer these questions comprehensively, we use a diverse inventory of data sets and samples from different sources and cohorts.

RESEARCH IN CONTEXT

1. Systematic Review: The authors reviewed literature in PubMed using the terms “cerebrospinal fluid”, “choroid plexus”, “aging” and “Alzheimer’s disease”. In contrast to the cerebrospinal fluid (CSF), where changes during aging and in Alzheimer’s disease (AD) are extensively reported, strikingly little is known about the role of the choroid plexus (ChP). Although severed structural integrity, transcriptomic and proteomic profiles and preliminary correlations between cognition, AD biomarkers, and ChP volumes were reported earlier, a systematic analysis of molecular changes in the ChP associated with AD is currently lacking.
2. Interpretation: Our study reveals a relationship between aging and AD in that changes in the CSF associated with aging are perturbed in the instance of AD. These CSF changes are associated with significant remodeling of the ChP in patients with AD, which in turn correlates with cognitive performance and reflects inflammatory signals and aberrant protein accumulations. Considering the ChP plays a critical role in brain homeostasis, this and previous studies provide mounting evidence of its dysfunction and participation in the pathogenesis of AD.
3. Future Directions: To clarify the role of the ChP in AD it is imperative to research: (1) the cause(s) of the observed ChP pathology in aged and AD brains, (2) the clinically relevant defects in brain homeostasis produced by dysfunctional ChP, and (3) the potential for a therapeutic strategy that normalizes ChP function in the instance of aging and AD.

To investigate changes in the CSF environment surrounding ChP in AD *vis-à-vis* natural aging, we analyzed a large publicly available data set of CSF proteomes consisting of healthy individuals and patients with AD^{82,83} from scratch to take into account the effects of aging. To establish how specific are the CSF changes to AD, we used mass spectrometry to measure CSF proteins in patients with mild cognitive impairment (MCI) or dementia due to AD, an infectious neurological disease, acute Lyme polyradiculoneuritis (ALymeP), and a faster progressing neurodegenerative disorder, amyotrophic lateral sclerosis (ALS).

To test for structural, inflammatory and metabolic changes of the ChP *post mortem* in AD, we harvested ChP from the lateral cerebral ventricles including ChP from the temporal horns of healthy individuals and patients afflicted by AD and other neurological conditions. The formalin fixed paraffin embedded (FFPE) sections of ChP were stained with hematoxylin and eosin (H&E) to examine gross structural changes. To probe for neurodegenerative changes, the ChP were stained with markers for amyloid deposits, neurofibrillary changes and α -synuclein.

To test for inflammation, the FFPE sections were stained with markers for macrophages and T lymphocytes. RNA isolated from snap frozen ChP was evaluated by real-time polymerase chain reaction (RT-PCR) for the expression levels of different markers of immune system activation and lysates from snap frozen ChP were prepared to measure cytokine levels by ELISArray and western blots. To test for metabolic changes in the ChP, the same lysates used to measure cytokine levels were subject to mass spectrometry to quantify a number of CSF resident proteins selected using a random number table in addition to quantifying cholesterol and different lipids.

To investigate ChP in vivo, we first used a 1.5T magnetic resonance imaging (MRI) scanner to establish the intensities, structural changes and volumes of ChP in healthy individuals and patients with AD. Volumes of ChP were measured from three-dimensional (3D) reconstructions as well as manually from a randomly selected subset of originally acquired MRI brain images. A 3.0T MRI scanner was then used to measure ChP volumes at a higher resolution in an independent cohort of healthy individuals and AD patients to further validate our original findings. The ChP volumes were lastly correlated with age and the cognitive performance using the Mini-Mental State Examination (MMSE).

The abnormalities observed in the CSF of patients with AD consisted largely of aberrancies in the same biological processes that changed most significantly with aging in healthy individuals. These findings were most pronounced in the 66- to 75-year-old age group and suggest that ChP in AD is perpetually bathed in CSF exhibiting protein abnormalities that evolve with aging. The H&E stained FFPE sections of ChP showed age-related structural changes, which were indistinguishable between healthy individuals and patients afflicted by AD and other conditions at a light microscopy level. In AD, ChP exhibited significantly increased levels of several cytokines, aberrant accumulation of transthyretin (TTR) and Niemann-Pick intracellular cholesterol transporter 2 (NPC2) and reduced levels of GM1 ganglioside. Therefore, the ChP hosts unexpected inflammatory signals and aberrant accumulation of molecules involved in metabolic pathways in AD. Imaging showed significantly increased intensities and remodeling of ChP in AD patients. ChP volumes measured automatically, using 3D reconstructions, manually or with a 3.0T MRI scanner on a separate cohort were significantly increased in AD patients compared with healthy individuals. The ChP volumes correlated with age and inversely with cognitive performance in AD patients in vivo and thus provided clinical relevance to our *post-mortem* ChP findings in AD.

1.3 | Discussion and study conclusions

The preliminary experiments described here document an unexpected confluence of inflammatory signals and aberrant accumulation of metabolically active ChP resident proteins that are physiologically secreted into the CSF in patients with AD. Additionally, proteomic alterations including evidence of inflammation develop and perturb physiological process of aging in the CSF.⁸⁴ These findings are clinically relevant considering patients with AD show structural changes

of the ChP, which correlate with aging and cognition.⁸⁴ Although these findings require further validation, they reveal a repertoire of ChP pathologies that lay the foundation for testing the role of ChP dysfunction mechanistically in the pathogenesis of AD.

In contrast to previous studies of the CSF in aging^{72,85} and AD,^{76,77,79,82,86} our analyses reveal that the same biological processes change in both of these conditions but to a different extent. This suggests that CSF changes in AD reflect a perturbed process of physiological aging. Changes in the inflammatory pathways identified in AD are more pronounced in other disorders we examined, possibly due to faster clinical progression of both AylmeP and ALS. Despite similarities, deeper analyses disclose subtle differences in inflammatory pathways between AD, AylmeP disease, and ALS,⁸⁷⁻⁸⁹ which indicate an additional layer of inflammatory changes beyond aging. The finding that the most significant CSF changes in sporadic AD occur at the same time when the risk of AD starts increasing⁹⁰ raises the question of whether these changes relate causally to the pathogenesis, reflect brain pathology, or indicate a protective response.^{77,91,92} The observations that CSF in the young exhibits restorative properties⁹³ and that a dampened immune response evolutionarily favors longevity,⁹⁴ support the notion that CSF changes in AD reflect pathological rather than physiological events.

The described abnormalities of the ChP validate and extend previously reported findings in animal models^{67,68} and patients with AD,^{64,69,95} which together with this study provide compelling evidence of ChP dysfunction in AD. Considering ChP provides the brain with nutrients and removes its waste, any dysfunction is expected to significantly perturb brain homeostasis.^{21,58,96} This is further corroborated by the impaired production and altered composition of the CSF in aging⁹⁷ and in AD.^{85,98,99,100} Intriguingly, many of the cytokines that we found to increase in the ChP were previously reported having increased concentrations in the CSF in AD.^{77,101} A lack of inflammatory infiltrates in the ChP suggests that these cytokines originate from the CSF, but could also be locally produced or reflect unresolved inflammation.¹⁰² In all cases, functional ChP is essential for timely sensing and transmission of information about peripheral inflammatory insults to the brain.^{52-56,61,103-105} It is improbable that dysfunctional ChP, bathed in a CSF marked by inflammatory changes,⁸⁴ senses and transmits such information to the brain in AD correctly. Therefore, the ChP in AD likely misconstrues the peripheral inflammatory status and contributes to the pro-inflammatory state of the brain beyond normal aging,^{31,32,106-108} which promotes synaptic failure, impairs brain connectivity, reduces cognitive reserve, and ultimately culminates in cognitive decline.

We identify structural changes of the ChP in vivo in patients with AD, which correlate with aging and inversely with cognitive performance. Our findings are consistent with previously reported associations between ChP volumes, biomarkers of AD and cognition.^{70,71} They extend previously noted association between aging⁶⁵ and ChP by showing that increased ChP volumes with aging undergo further increase and significant remodeling of their shapes in AD. Considering the paucity of in vivo studies of the ChP, it is essential to discern the diagnostic value of ChP changes in AD defined with biomarkers,¹⁰⁹

*post-mortem*¹¹⁰ or syndromically,¹³ and relative to other dementias. Given the lack of efficient treatments for AD, our work suggests that strategies targeting ChP may be a novel and promising therapeutic option in the instance of AD and other neurodegenerative disorders.^{111,112}

1.4 | Limitations, unanswered questions, and future directions

Although the use of *post-mortem* ChP tissue is critical to document human pathology, it comes with several shortcomings. Such tissues cannot provide mechanistic insights and are inevitably subject to a degree of damage caused by delayed collection. To circumvent the latter, we performed MRI imaging studies in patients with AD. Considering this was a pilot study, the results await further validation using larger sample sizes, different cohorts, and optimally longitudinal study design.

Despite the repertoire of ChP pathologies described in AD, the mechanisms underlying ChP damage in the pathogenesis of AD remain unclear. Considering ChP is bathed in the CSF, which is significantly altered in AD, a direct role of the CSF compromising ChP provides the simplest explanation. However, the fact that aging is the major risk factor for AD and simultaneously affects ChP and CSF suggests that inherent changes of the ChP beyond normal aging per se could also be at play. It remains in fact unknown whether upregulation of the host-defense programs¹¹³ occurs properly during aging with enhanced AD risk. Although current studies suggest that increased ChP volumes observed by us and others^{70,71} in AD are not the result of compensatory increase in the CSF production,^{69,98,99} but rather of perturbed local blood flow and permeability,^{114,115} significant further work is needed to provide a mechanistic account of the observed structural changes, the effects of systemic arterial hypertension,¹¹⁶ and correlations with aging and cognitive performance.

Regardless of the mechanisms underlying ChP pathology in AD, the inflammatory, metabolic, and other changes of the ChP and the CSF suggest their contribution in perturbing brain homeostasis including the correct sensing and transmission of immune signals from the periphery to the brain^{22,50} and call for studies of the role of the ChP in the pathogenesis of AD. Regrettably, current lack of bona fide cell and animal models of sporadic AD makes it extremely difficult to study mechanisms underlying ChP pathology and its repercussions. Testing how AD-like composition of the CSF alters ChP, investigating the effects of risk factors of AD on the structure and function of ChP and recapitulating aging and mimicking AD in ChP organoids¹¹⁷ will likely open new vistas in our understanding of the ChP. A more nuanced understanding of transcytosis-mediated relays across the ChP in AD is also needed. Studying ChP longitudinally in patients afflicted by a range of neurodegenerative disorders, correlating its changes with behavior, cognition, and other neurological deficits, evaluating rigorously the effects of cardiovascular and other risk factors, and teasing apart the effects of compromised ChP from the disrupted BBB in AD^{118,119} should be informative.

2 | CONSOLIDATED RESULTS AND STUDY DESIGN

To distinguish alterations of the CSF that can compromise ChP in AD from the effects of aging, we first re-examined a publicly available CSF proteomic data set representative of a wide age range of cognitively intact healthy individuals and patients with AD (Figure 1A). Half of the most significantly changed biological processes identified in the whole cohort consisted of pathways involving inflammation (Figure 1B). When the cohort was split to compare AD patients with healthy individuals, only innate immunity, neutrophil degranulation, complement cascade, and amyloid fiber formation related biological processes ranked higher in significance in AD (Figure 1C). The most prominent changes in biological processes coincided with the age period with the highest incidence of AD and consisted of biological processes involved in the inflammatory response, the immune system, interleukin signaling, cytokine-mediated signaling, innate immune system and immune cell activation (Figure 1E and F). Further mass spectrometry analyses revealed that the same biological processes were altered also in inflammatory and other neurodegenerative disorders of the brain (Figure 2A-C). However, in these disorders, different proteins underlay further exacerbated inflammatory pathways compared with AD (Figure 2D).

To test whether CSF abnormalities in AD are accompanied by compromised ChP, the ChP were harvested from lateral cerebral ventricles including their temporal horns and examined for their gross structure. Light microscopy of H&E stained FFPE sections of ChP showed similar degree of epithelial atrophy, stromal fibrosis, blood vessel thickening, and calcifications in healthy individuals and patients with AD (Figure 3A and B, Figure S1A-D). Furthermore, examination of the neuropathological hallmarks of neurodegeneration found no amyloid deposits, neurofibrillary changes, nor α -synuclein inclusions in the ChP of healthy individuals and patients afflicted by early or late AD or AD with Parkinson's disease (PD) (Figure S1E-T). To examine inflammatory changes, the ChP were evaluated for inflammatory infiltrates, activation of the immune cells and the cytokine levels. H&E stained ChP showed similar number of inflammatory cells in healthy individuals and patients afflicted by AD (Figure 3C). This finding was further confirmed by staining FFPE ChP sections with an anti-CD68 antibody, a bona fide marker of macrophages (Figure 3D). Since lack of inflammatory infiltrates does not inform about the activation of the immune cells in the ChP, we next examined the expression levels of different markers of the immune system activation by RT-PCR. There were no differences in the expression levels of different markers of the activation receptors, cytokines, immune cell activation, as well as of the T-cell activation between healthy individuals and patients with AD (Figure 3E). We last measured cytokines in lysates prepared from snap frozen ChP and found significantly increased levels in several pro- and anti-inflammatory cytokines in AD patients compared with healthy individuals (Figure 3F, Figure S2).

To investigate ChP in AD further, we measured the concentrations of a subset of randomly selected CSF proteins as a proxy measure

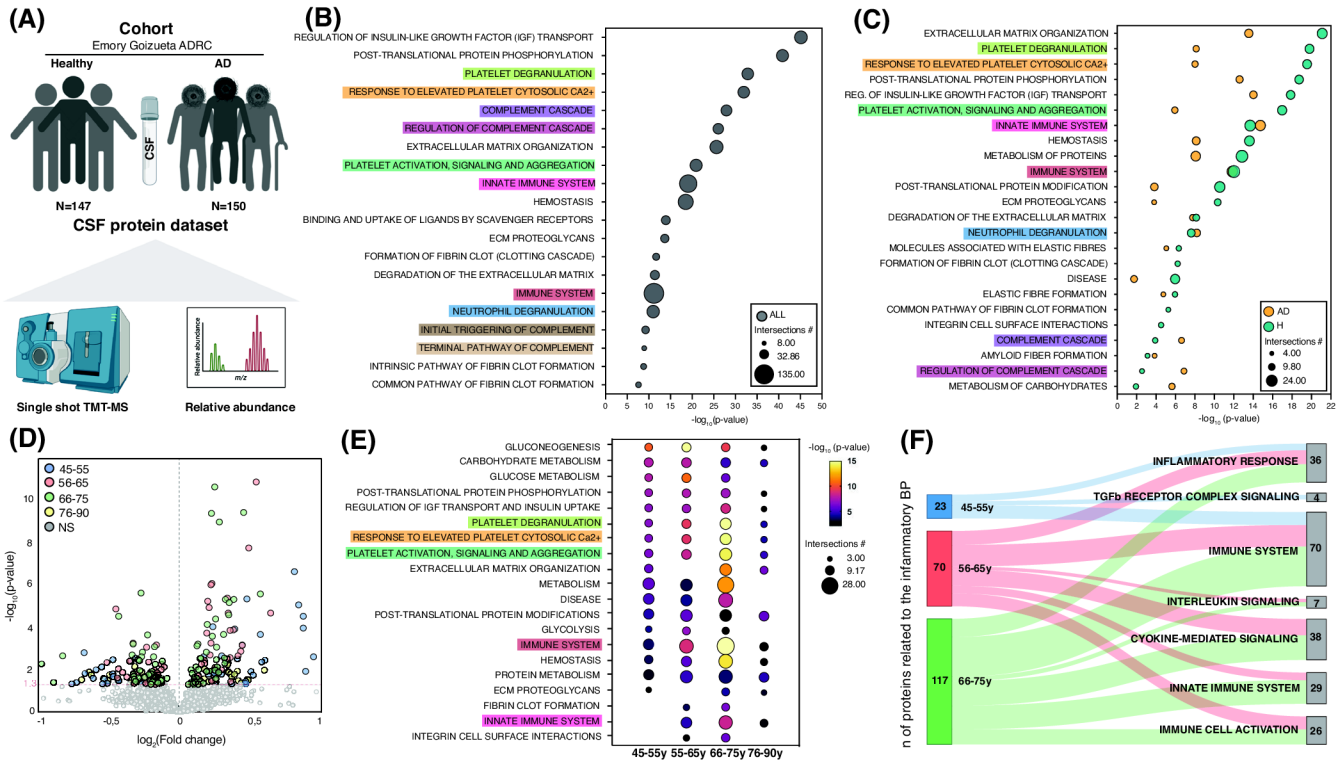


FIGURE 1 CSF in aging and AD. (A) Characteristics of the CSF protein database from the Emory Goizueta ADRC cohort. (B) The most significant Ingenuity pathways based on the CSF proteins changed by aging ($p \leq 0.05$). (C) The most significant Ingenuity pathways based on the CSF proteins changed by aging in healthy individuals versus AD ($p \leq 0.05$). (D) Volcano plot showing significantly changed CSF proteins in AD compared with healthy individuals in 45–55 ($n = 29$), 56–65 ($n = 97$), 66–75 ($n = 133$) and 76–90 ($n = 38$) year-old age groups ($p \leq 0.05$, NS: not significant). (E) The most significant Ingenuity pathways based on the CSF proteins changed by aging in AD compared with healthy individuals in different age groups ($p \leq 0.05$). (F) Sankey diagram showing significantly changed inflammatory pathways in AD compared with healthy individuals across different age groups ($p \leq 0.05$)

of its function. The proteotypic sequence of half of the randomly selected CSF proteins was successfully validated using isotopically labeled standards and their concentrations in the ChP measured using mass spectrometry (Figure 4A). In contrast to other proteins, NPC2 and TTR showed significantly increased levels in the ChP in patients with AD compared with healthy individuals. To investigate these protein changes further, we stained FFPE ChP sections with antibodies against NPC2 and TTR. We observed significantly increased mean intensities of both NPC2 and TTR in the ChP in AD patients compared with healthy individuals, predominantly located in the epithelium (Figure 4B and C). While both proteins have been previously associated with AD,^{120,121} genetic dosage of NPC2 has also been reported to regulate cholesterol and lipid metabolism.¹²² Mass spectrometry measurements showed significantly decreased levels of anti-inflammatory GM1 ganglioside, but not of cholesterol nor of other lipids in the ChP of patients with AD compared with healthy subjects (Figure 4D, Figures S3 and S4A-F).

To test whether changes identified in the ChP *post mortem* have a clinical or *in vivo* correlate in AD, we studied ChP using 1.5T MRI (Figure 5A). The T1 weighted sequences of ChP showed significantly increased intensities in AD patients compared with healthy individuals, which suggests that ChP undergoes changes in its composition

in the course of AD (Figure 5B). Examination of the shapes of 3D reconstructed representations of the ChP is consistent with significant remodeling in AD considering the overall hypertrophy of the ChP accompanied by its atrophy along the anterior margins extending distally toward hippocampi (Figure 5C). Automated volumetric measurements of hippocampi, cerebellar cortices and ChP found significantly reduced volumes of the hippocampi, no changes in cerebellar cortices and significantly increased volumes of ChP in patients with AD (Figures 5D and 5E). To confirm this finding, volumes of 3D reconstructed representations were also measured and showed significantly reduced hippocampal and increased ChP volumes in AD patients (Figure 5F). ChP volumes are in this case smaller, because 3D representations are based on continuous triangulated surfaces and thus more accurate compared with the automated measurements, which are vulnerable to include also artefactual voxels. Manual measurements of ChP in a randomly selected subset of MRI brain images further confirmed this finding (Figure 5G). We last imaged ChP at a higher resolution using a 3T MRI and again reproduced previously observed increased ChP volumes in AD patients compared with healthy individuals in a separate cohort (Figure 5H). Notably, increased ChP volumes correlated with age and inversely with cognitive performance in patients with AD (Figure 5I and 5J).

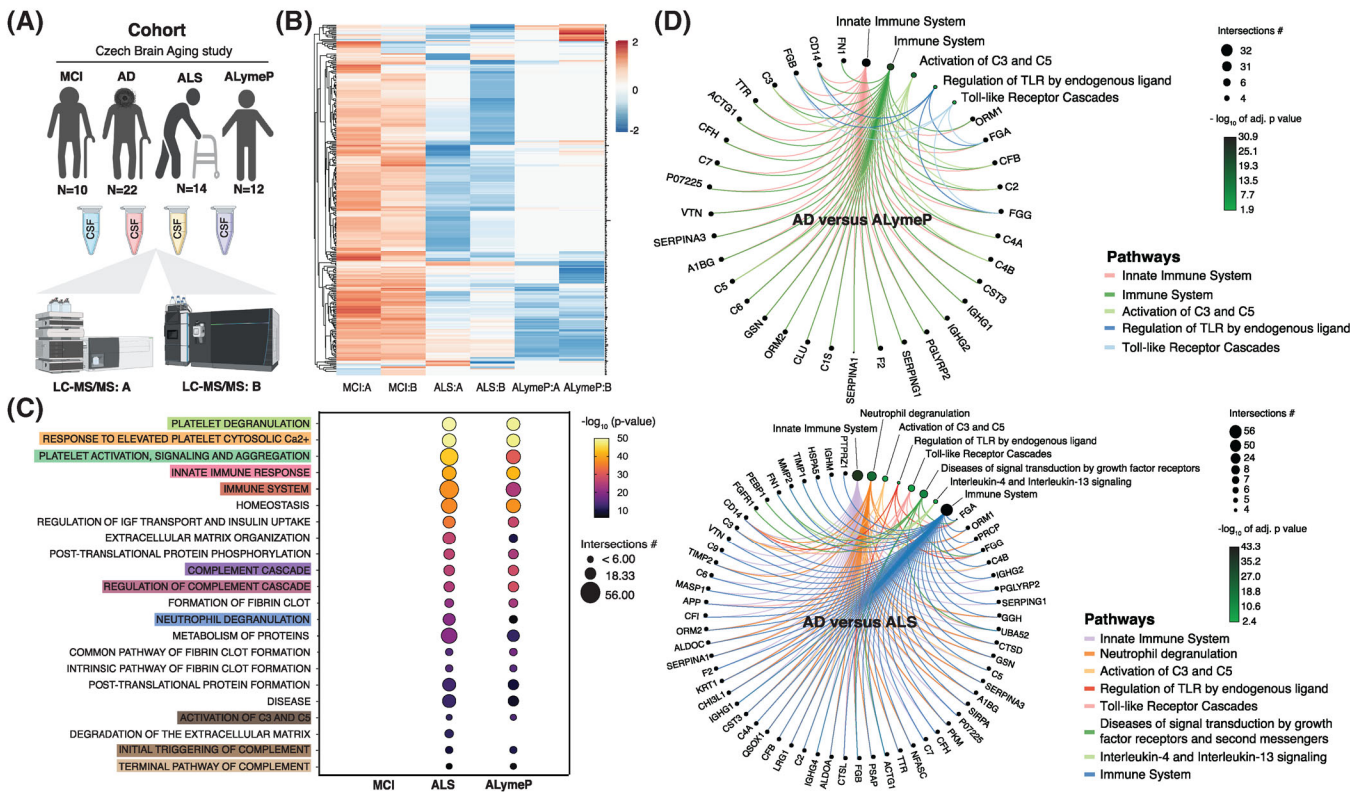


FIGURE 2 CSF in AD and other conditions (A) Characteristics of the Czech Brain Aging Study cohort. (B) Heatmap representing hierarchically clustered CSF proteins identified by two independent mass spectrometers and protocols (A and B) in MCI, ALyMeP, and ALS compared with AD ($p \leq 0.05$ FDR adjusted t -test). (C) The most common significantly changed Reactome pathways in MCI, ALyMeP, and ALS compared with AD ($p \leq 0.05$). (D) Circular netplots showing significantly changed inflammatory CSF proteins and pathways in AD compared with ALyMeP and ALS ($p \leq 0.05$)

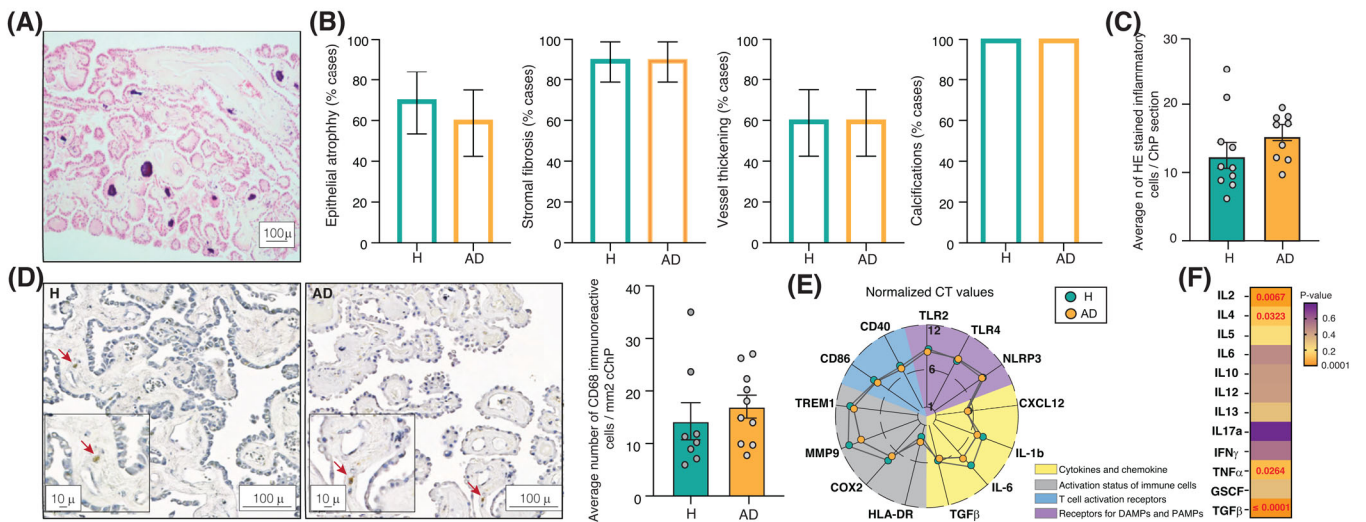


FIGURE 3 Structure and the inflammatory status of the ChP in AD. (A) Representative H&E stained FFPE section of the ChP. (B) Percentage of healthy individuals and AD patients showing epithelial atrophy, stromal fibrosis, vessel thickening and calcifications in the ChP. One technical replicate \pm SEM; two-sample t -test ($p \leq 0.05$). (C) Average number of inflammatory cells in H&E stained ChP from healthy individuals and AD patients. Mean number of cells per six high-power microscopy fields per section \pm SEM; two-sample t -test ($p \leq 0.05$). (D) Anti-CD68 antibody labelled cells in the ChP of healthy individuals and AD patients. Mean number of cells per 10 high-power microscopy fields per section \pm S.E.M.; two-sample t -test ($p \leq 0.05$). (E) Expression levels of markers of the activation of the ChP immune cells in healthy individuals and AD. The lines crossing central, middle and outermost circle represent distances corresponding to normalized CT values of 1, 6, and 12, respectively. (F) ELISA Array measured ChP cytokine levels in healthy individuals and AD. Mean of 3 technical replicates per sample \pm SEM; two-sample t -test ($p \leq 0.05$)

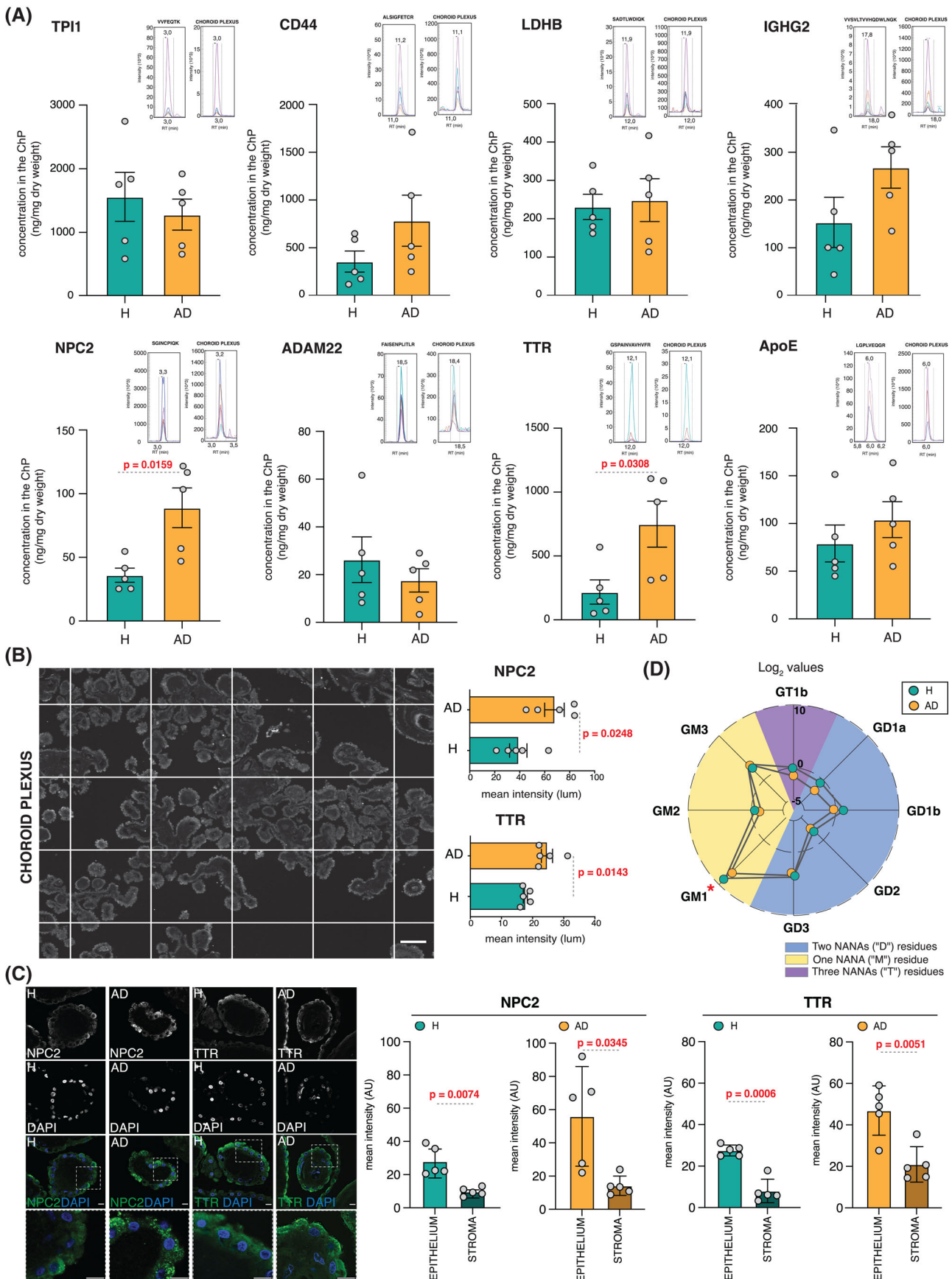


FIGURE 4 Aberrant accumulation of CSF resident proteins in the ChP in AD. (A) Levels of CSF resident proteins in the ChP of healthy individuals and AD patients. Skyline iRT retention time prediction algorithm showing identical chromatogram retention times between

3 | DETAILED METHODS AND RESULTS

3.1 | DETAILED METHODS

3.1.1 | Study participants

Clinical data and samples were obtained from the Emory Goizueta ADRC CSF proteomics of 300 individuals,^{82,83} UCSD Shiley-Marcos ADRC, the Czech Brain Aging Study,¹²³ the Argentina ADNI,¹²⁴ the Barrow Neurological Institute¹²⁵ and the Imperial College Parkinson's UK Brain Bank¹²⁶ (Table S1). Collection of all samples was approved by the Institutional Review Boards of each participating institution with written consent obtained from all the individuals.

3.1.2 | Tissue collection

CSF samples were collected and stored following best practice protocols set forth by the National Institutes of Health Alzheimer's Disease Research Centers (NIH ADRCs) and others^{123,125–128}.

3.1.3 | Cognitive testing

All study individuals underwent comprehensive diagnostics consisting of clinical evaluation including laboratory tests and brain imaging according to NIH and other best practice guidelines.^{123,125} ALymeP was diagnosed when individuals demonstrated clinical characteristics, increased CSF leukocyte count and positive serology for *Borrelia burgdorferi*.

3.1.4 | Pathology

Senile plaques were probed using the primary antibody recognizing amyloid- β (4G8, Covance), neurofibrillary changes using an antibody against phosphorylated tau (AT8, Thermo Fisher Scientific), and Lewy pathology using the α -synuclein clone 42 antibody (#610787, BD Transduction Labs).

3.1.5 | ELISArray

ChP were homogenized using a Bullet Blender (Next Advance). Samples were assessed for their protein content using Pierce BCA protein

assay (Thermo Fisher Scientific). Cytokines were measured using Multi-Analyte ELISArray (QIAGEN). Cytokine levels were determined based on the absorbance values of negative (buffer) and positive (standards for all cytokines examined) controls. All measurements were normalized to the total protein content.

3.1.6 | Western blots

Protein lysates were separated on 4%–20% sodium dodecyl sulfate (SDS)-polyacrylamide gels and transferred to polyvinylidene fluoride (PVDF) membranes (Bio-Rad Laboratories). After blocking, membranes were incubated with antibodies against interleukin 2 (IL-2; D7A5, Cell Signaling Technology), IL-6 (#6672, Abcam), IL-10 (#34843, Abcam), tumor necrosis factor α (TNF α ; #9739, Abcam), and transforming growth factor beta (TGF β 1; #3711, Cell Signaling Technology). Cytokine levels were normalized to β -actin.

3.1.7 | Immunohistochemistry

The PPFE ChP sections were blocked and incubated overnight with primary antibodies against NPC2 (1:50, sc-166449, Santa Cruz), TTR (1:500, A0002, DAKO), CD3 (1:500, NCL-LCD3-565, Novocastra), or CD68 (1:100, ab955, Abcam).

3.1.8 | Gene expression analysis

RNA was prepared using the miRNeasy kit (Qiagen, Germany) and transcribed using the High-Capacity cDNA Reverse Transcription Kit (Thermo Fisher Scientific). Quantitative RT-PCR (qPCR) was carried out with Taqman probes (Taqman Gene Expression Assay, Thermo Fisher Scientific) using TaqMan Gene Expression Master Mix (Thermo Fisher Scientific). qPCR analysis was performed using a LightCycler II (Roche). The Ct values (Δ Ct) of genes of interest were normalized to house-keeping genes.

3.1.9 | Microscopy and image processing

ChP images were acquired at 10 \times using AxioScan.Z1 scanner microscope (Zeiss). Acquired images were analyzed using the Image Pro Premier 3D software (Media Cybernetics). Distribution of NPC2 and TTR was examined using a 63 \times oil-immersion objective with a confocal laser scanning microscope (Zeiss).

proteotypic peptides used to quantify ChP proteins (right) and stable isotopically labeled synthetic peptides (left). Mean of six technical replicates per sample \pm S.E.M.; two-sample *t*-test ($p \leq 0.05$). (B) Mean intensities of NPC2 and TTR in the scanned ChP of healthy individuals and AD patients (magnification bar 100 μ). Mean of 90 technical replicates per sample \pm S.E.M.; two-sample *t*-test ($p \leq 0.05$). (C) Mean NPC2 and TTR intensities in the confocal images of the ChP epithelium and stroma in healthy individuals and AD (magnification bar 10 μ). Mean of 3–12 technical replicates \pm SEM; repeated measures ($p \leq 0.05$). (D) Levels of ChP gangliosides in healthy individuals and AD. Mean of six technical replicates per sample \pm SEM; two-sample *t*-test ($p \leq 0.05$)

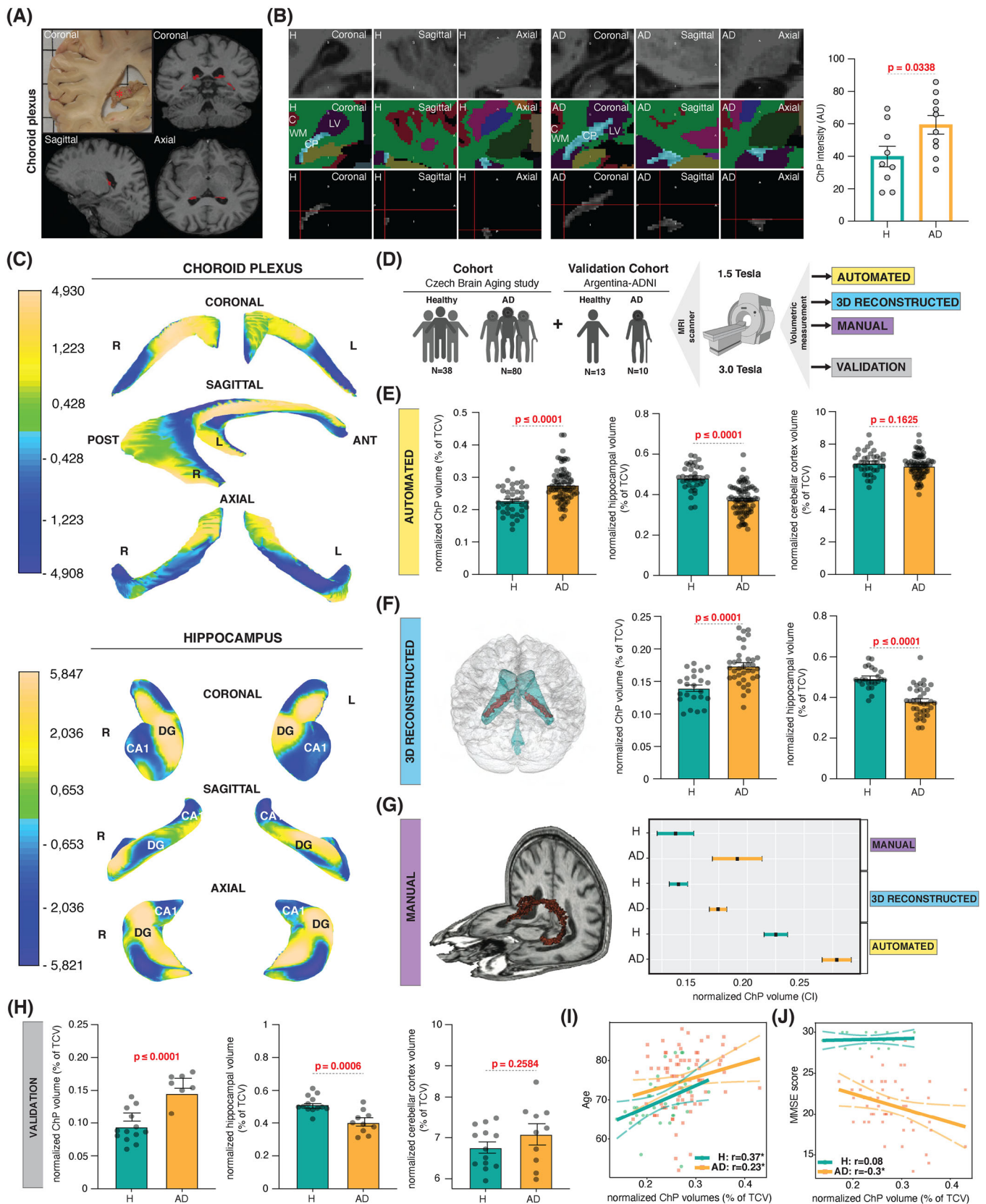


FIGURE 5 Increased intensity, remodeling, and volume of ChP in AD. (A) A coronal view of *post-mortem* ChP in the right lateral ventricle (red asterisk) and T1-weighted sequence derived positions of ChP in coronal, sagittal and axial views (in red). (B) Coronal, sagittal, and axial views of ChP representative of healthy individuals and AD patients visualized using T1-weighted sequence (top), FreeSurfer brain structure map (middle), and following ChP segmentation (bottom). Normalized ChP intensities in healthy individuals and AD. Individual measurements \pm SEM; two-sample

3.1.10 | Quantitative histology

Six high-magnification ChP images per section stained with either H&E or the anti-CD-68 antibody were scored for the number of inflammatory cells. To measure total mean NPC2 and TTR intensities, a grid of 500 × 500 pixel squares was placed over each image. A total of 30 grid squares were randomly selected to sample intensities across the entire ChP. Three rectangular regions of interest per each grid square were examined for a total of 90 regions of interest per image. Mean intensities were calculated using Image Pro Premier 3D software (Media Cybernetics). To examine the distribution of NPC2 and TTR within the ChP, the sections were imaged with a 63× oil-immersion objective using a confocal laser scanning microscope (Zeiss). 1–4 ChP grape-like structures in three independent fields of view were sampled per each slide.

3.1.11 | Mass spectrometry proteomics of the CSF

CSF samples were analyzed using two independent mass spectrometry protocols. According to the first protocol (A), three technical replicates of 500 ng of tryptic peptides per sample were analyzed by LC-MS/MS using an UltiMate 3000 RSLCnano system (Thermo Fisher Scientific) coupled via an EASY-spray ion source (Thermo Fisher Scientific) to an Orbitrap Elite mass spectrometer (Thermo Fisher Scientific). According to the second protocol, one technical replicate of 1 μg of tryptic peptides per sample was analyzed by LC-MS/MS using an UltiMate 3000 RSLCnano system coupled to an Orbitrap Fusion Lumos tribrid mass spectrometer (both Thermo Fisher Scientific).

3.1.12 | CSF proteomic data processing

The raw data were analyzed using MaxQuant software (version 1.5.6.5). The MaxQuant output file (ProteinGroups.txt) was uploaded into Perseus software (version 1.6.1.1). Previously described 300 patient Emory cohort CSF tandem mass tag (TMT)-based quantitative discovery mode proteomics data set^{82,83} was re-analyzed for age-stratified differences. In contrast to previous work, regression of age was not performed in the current analysis.

3.1.13 | Processing of ChP tissue for mass spectrometry assays and global lipid analysis

ChP samples were homogenized using BeadBlaster 24 (Benchmark Scientific Inc.). Isopropanol was added to each sample, which was vortexed, sonicated, and centrifuged at RT. The supernatant was then removed for the ganglioside and other assays (lipid extract). Remaining pellets were dried (Savant SDP121 P, SpeedVac, Thermo Fisher Scientific), reconstituted using glass beads in AmBic buffer with SDC, homogenized, vortexed and sonicated (protein extract). Synthetic isotopically labeled standard peptides containing an enzymatically cleavable tag at C-terminus were added to control for the variance in trypsin digestion.

3.1.14 | Validation of mass spectrometry targets

Gangliosides were quantified using selected reaction monitoring (SRM) assays in lipid extracts, which were internally standardized with isotopically labeled GM1 and GM3, respectively. Proteins were also analyzed using SRM assays for proteotypic peptides as protein surrogates. Tentative identifications of proteotypic peptides detected in ChP and concordant with the Skyline iRT Retention Time Prediction algorithm were validated using their isotopically-labeled analogs.

3.1.15 | Mass spectrometry quantitative protein assays

Dried solid-phase extracts of ChP digests were reconstituted in acetonitrile with formic acid. Samples were analyzed using ESI-UHPLC-SRM mass spectrometry on a triple quadrupole mass analyzer (1290 Infinity II and 6495B, Agilent) in positive ion detection mode.

3.1.16 | Mass spectrometry ganglioside assays

Gangliosides in lipid extracts of ChP were examined using both ESI-UHPLC-SRM positive and negative ion detection modes. Gangliosides were quantified with the stable isotope-labeled GM3 using appropriate response factors.

t-test ($p \leq 0.05$). (C) Mean changes in the shape of the 3D representations of the ChP and hippocampi in AD compared with healthy individuals. Single measurements ± SEM; two-sample *t*-test ($p \leq 0.05$). (D) Experimental design of the ChP volumetry study. (E) Automated measurements of normalized mean ChP, hippocampal and cerebellar cortical volumes in healthy individuals and AD. Individual measurements ± SEM; two-sample *t*-test ($p \leq 0.05$). (F) 3D reconstruction of the ChP (red) within the ventricular system (blue) in the brain. Normalized mean 3D representation-derived ChP and hippocampal volumes in healthy individuals and AD. Single measurements ± SEM; two-sample *t*-test ($p \leq 0.05$). (G) 3D representation of manually traced ChP (in red). 95% confidence intervals of normalized automated, 3D reconstructed and manually traced ChP volumes in healthy individuals and AD. (H) Normalized mean ChP, hippocampal and cerebellar cortical volumes in healthy individuals and AD. Individual measurements ± SEM; two-sample *t*-test ($p \leq 0.05$). (I) Correlation between normalized mean ChP volumes and age in healthy individuals and AD. Individual measurements ± SEM. (J) Correlation between normalized mean ChP volumes and MMSE scores in healthy individuals and AD. Individual measurements ± SEM.

3.1.17 | Mass spectrometry lipid analysis

Orbitrap Fusion (Thermo Fisher Scientific) and UHPLC (Nexera X2, LC-30AD, Shimadzu) with identical C18 column (Waters Corporation) thermostated at 40°C and gradient elution program as specified for ganglioside assays in the positive ion mode were used to measure lipids.

3.1.18 | Cholesterol spectrophotometric assay

Cholesterol was extracted from 1 mg of dry ChP tissue. The sample ultra-filtrate was transferred to a separate vial. Cholesterol was quantified using a colorimetric assay (MAK043, Sigma-Aldrich, USA).

3.1.19 | Brain MRI imaging

T1-weighted 3D high-resolution MRI images were acquired using magnetization-prepared rapid gradient echo (MP-RAGE) and fast spoiled gradient echo (FSPGR) pulse sequences using 1.5T (Siemens Avanto) in the Czech Brain Aging Study and 3.0T scanner (GE Signa HDxt) in the Argentina ADNI.

3.1.20 | MRI intensity measurements

Based on FreeSurfer wrapped in R, all segmented regions of interest and their masks were exported, including ChP intensities. Since raw intensities of ChP preclude their direct use in statistical analyses, we normalized raw intensity of ChP to CSF. The CSF strip algorithm was developed based on the white strip algorithm.¹²⁹

3.1.21 | MRI volumetry measurements

Volumetric measurements were performed using FreeSurfer automated algorithm version v5.3 (<http://surfer.nmr.mgh.harvard.edu/>)¹³⁰⁻¹³² MRI images were further evaluated using 3D reconstructions. Automatic segmentation of FreeSurfer was used to obtain segmentation masks. To examine 3D differences in shape, ChP were represented with 4246 mesh points.

3.1.22 | Manual MRI tracing

Manual segmentation was performed on a slice per slice basis using Slicer (www.slicer.org) with a paintbrush of 0.5 mm. Once the region of interest was fully delineated, potential over- and under-segmented areas were checked in coronal orientation and segmentations corrected. 3D volume rendering was performed to test for appropriate spatial appearance of ChP in line with the expected anatomy.

3.1.23 | Statistical analysis

All statistical analyses were performed using commercially available softwares (R, Prism, SPSS, and SAS). CSF proteomic and brain imaging analyses were performed using R (v.4.1.0). RColorBrewer, igrph, ggraph, and ggplot2 packages were used for differential CSF protein expression analyses. Brain imaging analyses were performed using Whitestripe, FreeSurfer, NeuroBase, fslr, fsbrain, misc3d, rgl, and ggplot2.

3.2 | DETAILED RESULTS

3.2.1 | The effects of aging, AD, and other neurological disorders on the CSF

Since ChP is surrounded by the CSF, pathological CSF alterations can directly compromise its structure and function. To distinguish these alterations in AD from the effects of aging, the major risk factor of AD,²⁹ we investigated the relative abundance of CSF proteins in a well-established CSF proteomic data set of healthy individuals and patients with AD (Figure 1A)^{82,83} Half of the most significantly changed biological processes overall consisted of inflammation-related pathways (Figure 1B). In contrast to healthy individuals, the majority of the most significantly changed biological processes by aging in AD showed reduced ranking in significance except for select pathways involved in inflammation and amyloid fiber formation (Figure 1C). Only pathways related to glucose, hyaluronan metabolism, binding and uptake of ligands by scavenger receptors were changed uniquely in AD patients. Our findings suggest that largely the same biological processes that change naturally with aging become perturbed in AD.

To investigate the relationship between aging, inflammation, and AD further, we divided the CSF proteomic cohort into four age groups (Figure 1D). The most pronounced changes between AD patients and healthy individuals occurred in the 66- to 75-year-old age group, which also showed the highest ranking in significance of the inflammatory pathways compared with all the other age groups (Figure 1E). Furthermore, the proteins and pathways involved in the inflammatory changes in the CSF of AD patients differed greatly between age groups (Figure 1F). These findings reveal an age-dependent evolution of proteins and pathways changing in the CSF of patients with AD compared with healthy individuals.

Studies of the CSF proteomic cohort provide compelling evidence that aging plays a role in the CSF changes in AD, but are not informative regarding their specificity to AD. To interrogate their specificity, CSF of patients with dementia due to AD were compared with CSF from patients with MCI, ALyMeP, and ALS (Figure 2A). CSF samples were run on two separate mass spectrometers and only those CSF proteins that were identified by both approaches were used to build the CSF proteomes of individual disorders (Figure 2B). While there were expectedly no major differences in the CSF proteomes between MCI and AD, predominantly inflammatory pathways had higher

ranking in significance in patients with ALymeP and ALS compared with AD (Figure 2C). Patients with ALymeP and ALS also differed from AD in the proteins involved in the inflammatory pathways of CSF (Figure 2D). These findings suggest that biological processes undergoing changes in the CSF in AD are affected also by other neurological disorders with inflammatory and faster progressing neurodegenerative diseases showing more pronounced CSF changes.

3.2.2 | Structural, inflammatory, and metabolic changes of the ChP in AD

Inflammation invariably alters the structure and function of involved tissues.⁸⁴ Considering the intimate contact between ChP and CSF, the inflammatory CSF abnormalities suggest possible damage of the ChP structure in AD. To examine the structure of the ChP, FFPE sections harvested from lateral cerebral ventricles including samples from the temporal horns were stained with H&E (Figure 3A). Light microscopy revealed comparable degree of epithelial atrophy, stromal fibrosis, blood vessel thickening, and calcifications in the ChP of healthy individuals and AD patients (Figure 3B, Figure S1A-D). Staining for amyloid, neurofibrillary changes, and α -synuclein also demonstrated lack of hallmark lesions in the ChP of patients with early or late AD or AD with PD (Figure S1E-T). ChP, therefore, exhibit no overt structural changes nor pathology in AD.

Staining with H&E (Figure 3C) or with anti-CD68 and anti-CD3 antibodies (Figure 3D), which label macrophages and T lymphocytes, respectively, showed no evidence of inflammatory infiltrates in AD patients compared with healthy individuals. We next investigated the activation status of the ChP immune system by screening RNA from snap frozen ChP for the expression levels of the activation receptors, TLR4, TLR9, and NLRP3; cytokines, IL1 β , IL6, and TGF β ; immune cell activation, TREM1, HLA-DR, COX2, and MMP9; and T-cell activation, CD40 and CD86 using RT-PCR. The expression levels of markers of the activation of the immune system were comparable between healthy individuals and AD patients (Figure 3E). We last measured cytokines in lysates from snap frozen ChP by an ELISArray. In contrast to healthy individuals, AD patients showed significantly increased levels of IL-2, IL-4, TNF α , and TGF β in the ChP (Figure 3F). We separated lysates also by SDS-PAGE and confirmed significantly increased cytokine levels in the ChP of AD patients (Figure S2). ChP in AD, therefore, lack infiltration or activation of inflammatory cells, but display unexpected inflammatory signals.

To assess the function of the ChP in AD, we quantified CSF proteins in the ChP as a proxy measure of metabolic damage. Proteotypic sequences of 8 out of 16 randomly selected CSF resident proteins passed the validation and their ChP levels quantified using mass spectrometry (Figure 4A). Levels of NPC2 and TTR were significantly increased in the ChP of AD patients compared with healthy individuals. Scans of ChP stained with antibodies against NPC2 and TTR further corroborated this finding (Figure 4B), while confocal imaging showed that aberrant accumulation of NPC2 and TTR immunoreactivity in the ChP is confined predominantly to the epithelium (Figure 4C).

Metabolically active proteins that are physiologically secreted from the ChP into the CSF, therefore, aberrantly accumulate in AD. Considering NPC2 deficiency perturbs lipid metabolism¹²² and increases ganglioside levels,¹²¹ we asked whether increased NPC2 levels show decreased ganglioside levels or otherwise perturbed lipid metabolism in the ChP in AD. Measurements of cholesterol, gangliosides, phosphatidylcholines, phosphatidylethanolamines, sphingomyelins, phosphatidylinositols, phosphatidylserines found reduced levels of GM1 ganglioside, but not of cholesterol nor of any other lipid specie in AD patients compared with healthy individuals (Figure 4D, Figure S3 and S4A-F). Considering GM1 harbors anti-inflammatory properties,¹³³ reduced GM1 levels further indicate perturbed inflammatory milieu of the ChP in AD.

3.2.3 | Imaging changes of ChP in AD patients

While our study provides evidence of compromised ChP in AD in *post-mortem* tissue samples, its clinical relevance remains to be established. To test this, we investigated structural changes of ChP in living healthy individuals and AD patients using MRI (Figure 5A). We first measured intensities of T1-weighted MRI images of ChP acquired using a 1.5T MRI scanner. Intensities of ChP were significantly increased in patients with AD compared to healthy individuals (Figure 5B). Considering this observation suggests remodeling of ChP in AD, we next examined changes in the shape of 3D reconstructed representations of the ChP and hippocampi. ChP exhibited significant overall hypertrophy with atrophic changes along the anterior margins extending distally toward the hippocampi in AD patients compared with healthy individuals (Figure 5C). Hippocampi showed previously described atrophic changes across their entire structure.¹³⁴ These experiments demonstrate that ChP undergoes significant structural changes in AD.

To test for changes in the size of the ChP, we next performed volumetric measurements of the ChP, hippocampi, which have been extensively reported to decrease in AD¹³⁵ and cerebellar cortices since previous reports found no major changes in AD¹³⁶ (Figure 5D). An automated volumetric measurement showed that ChP volumes significantly increased, while hippocampal volumes expectedly decreased and cerebellar cortices showed no changes in AD patients compared with healthy individuals (Figure 5E). To test this observation further, we measured volumes of 3D representations of ChP and hippocampi reconstructed from a randomly selected subset of MRI images. While 3D representations of hippocampal volumes demonstrated expected decrease, ChP volumes were significantly increased in AD patients (Figure 5F). To avoid segmentation-derived measurement errors,¹³⁷ we last measured ChP volumes manually. Manual measurements showed no overlap between the 95% confidence intervals of the ChP volumes in AD patients compared with healthy individuals further reinforcing the veracity of our findings (Figure 5G). We last acquired MRI brain images from a separate cohort using a 3.0T MRI scanner. In contrast to the hippocampal volumes, the ChP volumes were significantly increased in AD patients (Figure 5H). Increased ChP volumes

demonstrated stronger association with age in healthy individuals compared with AD patients (Figure 5I). AD patients, but not healthy individuals, also showed significant inverse correlation between ChP volumes and the MMSE scores (Figure 5J). Therefore, the larger the ChP volume, the poorer the cognitive performance in AD patients. Taken together, these experiments demonstrate the clinical relevance of our findings in regard to changes that we have noted *post mortem* in the ChP in AD.

AUTHOR CONTRIBUTIONS

Gorazd B. Stokin designed and planned the study, Maria Čarna and Gorazd B. Stokin analysed all of the data and wrote the manuscript, Isaac G. Onyango performed ChP cytokine experiments and analysis, Clara Limbäck-Stokin and Robert A. Rissman provided ChP samples, Clara Limbäck-Stokin performed all the ChP neuropathological experiments and analysis, Nadine Bakkar, Robert Bowser, Jakub Hort, and Kateřina Sheardova contributed CSF samples, Victoria David-Dirgo, Krystine Garcia-Mansfield, Dušan Holub, Ritin Sharma, and Patrick Pirrotte performed all the CSF mass spectrometry experiments, Jiří Damborsky, Durga Jha, Marketa Nezvedova, and Zdenek Spacil performed all the ChP mass spectrometry and other lipid experiments, Valentina Lacovich and Kateřina Texlova performed ChP immunohistochemistry and analysis, Hernan Chaves, Jakub Hort, Zuzana Nedelska, Lucia Pertierra Hana Markova, Gustavo Sevlever and Martin Vyhnalek were involved in the acquisition and analysis of MRI images, Stanislav Katina and Martin Vyhnalek performed MRI image and shape programming and analysis, including intensities and volumes, Ruben Houbrechts and Thijs Vande Vyvere performed manual ChP measurements, Maria Čarna, Eric B. Dammer, and Dušan Holub performed all the bioinformatics assessments and analysis, Silvie Belaškova, Jiří Damborsky, and Jan S. Novotny performed biostatistical analysis, Robert Bowser, Silvie Belaškova, Jiří Damborsky, Giancarlo Forte, Jan Frič, Jakub Hort, Patrick Pirrotte, Clara Limbäck-Stokin, Nicholas T. Seyfried, Gustavo Sevlever, Zdeněk Spačil, and Richard A. Smith provided continuous input during experiments and data analysis.

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CONFLICTS OF INTEREST

The authors report no competing interests. Author disclosures are available in the [supporting information](#).

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SUPPORTING INFORMATION

Additional supporting information can be found online in the Supporting Information section at the end of this article.

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