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Non-cell autonomous choroid plexus-derived sAPPα regulates adult hippocampus proliferation and plasticity

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Abstract
While Aβ peptides derived from amyloid precursor protein (APP) play a key role in Alzheimer disease pathogenesis, the extracellular sAPPα soluble ectodomain has been shown to stimulate adult neurogenesis and synaptic plasticity. Elevated expression of App in the choroid plexus has been recently reported, suggesting an important role for APP in cerebrospinal fluid. We conditionally knocked down App expression specifically in the adult mouse choroid plexus either by genetic deletion or shRNA expression, which led to reduced proliferation in both subventricular zone and hippocampal dentate gyrus neurogenic niches. Conversely, proliferation in both niches was increased either by viral expression of App in choroid plexus or by infusion of sAPPα but not sAPPβ in cerebrospinal fluid. To test the hypothesis that favoring the production of Aβ specifically in choroid plexus could negatively affect niche functions, we used AAV5 mediated expression of human mutated APP specifically in the choroid plexus of adult wild type mice. These mice showed reduced niche proliferation and, after one year, exhibited behavioral defects in reversal learning. Consistent with impaired memory, electrophysiological analysis revealed impaired synaptic plasticity as evidenced by deficits in hippocampal LTP. Our findings highlight the unique role played by the choroid plexus in regulating brain function, and suggest that targeting APP in choroid plexus may provide a means to improve hippocampus function and alleviate disease-related burdens.
### Introduction

Alzheimer disease (AD) is a neurodegenerative disease that affects the aging human population. Genetic risk factors include AD associated mutations in the amyloid precursor protein (APP) and presenilin 1 (PSEN1 and PSEN2) genes and others (e.g. APOE4 allele) contributing to the onset and development of the disease [16]. Pathologically, the brain of AD patients is characterized by neurofibrillar tangles, consisting of hyperphosphorylated forms of the microtubule-associated protein TAU, and by amyloid plaque deposits, enriched in extracellular and intracellular Aβ40 and Aβ42 (thereafter Aβ) amyloid peptides. Aβ peptides are generated from APP by sequential proteolytic cleavage by β- and γ-secretases, while cleavage by α-secretase releases the secreted APPα ectodomain (sAPPα) from the cell surface. Due to the position of the 3 cleavage sites, the secretion of sAPPα precludes the generation of Aβ peptides. In contrast, cleavage at the β site releases the sAPPβ ectodomain, in parallel with Aβ peptide production.

An important and still under-investigated issue is the physiological function of sAPPα. Indeed, the pathological traits accompanying amyloid plaque formation could potentially be explained by a parallel decrease of sAPPα levels resulting in lost functions that cannot be compensated by sAPPβ [33]. Key functions of sAPPα include its ability to enhance spine density, synaptic plasticity and memory, as well as neurogenesis [24, 28]. Expression of sAPPα in hippocampus has been shown to rescue cognition and synaptic transmission and to mitigate synaptic and cognitive deficits in a pathological context [12, 43, 50]. sAPPα dependent signals are important for neuroprotection through sAPPα binding to transmembrane APP [23], and in synaptic plasticity through binding to partners such as the recently identified GABAAR1α ligand and the α7-nACh receptor [26, 32, 33]. Furthermore, APP plays a role in neurogenesis, either as full-length transmembrane APP [47] or through its sAPPα ectodomain [4]. In the latter study, it was shown that sAPPα acts through binding to the EGF receptor, but the cellular source of the sAPPα ectodomain was not identified.

In addition to neurofibrillary tangles and amyloid plaques, a deficit in adult neurogenesis has been associated with AD [5, 25, 34, 45, 49]. Although adult neurogenesis in humans has been recently challenged [30, 44], it has been demonstrated conclusively in rodents, with two primary neurogenic sites, the subventricular zone (SVZ) and the subgranular zone (SGZ) of the hippocampal dentate gyrus (DG). Neuroblasts produced in the rodent SVZ migrate away into the olfactory bulb where they differentiate into inhibitory neurons. In the hippocampus, neuroblasts integrate locally in DG and differentiate into glutamatergic neurons. Humans constitute an important exception since SVZ-derived neuroblasts may migrate into the striatum [10]. In the transgenic APP/PS1 mouse model of AD, a decrease in adult SGZ neurogenesis was observed in addition to amyloid plaques throughout the brain parenchyma including the hippocampus [51]. Moreover, sAPPα stimulates the proliferation of SVZ and SGZ neural progenitors [1, 4, 8], while viral expression of sAPPα in the hippocampus of the APP/PS1 mouse rescues synaptic failure [12]. Although circumstantial, these results evoke the possibility that an impairment in adult neurogenesis may contribute to healthy and pathological ageing. In fact, even if the recent controversy on adult neurogenesis in the humans cannot be ignored, a recent study convincingly showed that adult DG neurogenesis is maintained at a high level until old age but drops sharply in AD patients, supporting the AD “neurogenesis” hypothesis of cognitive impairment [25].

The cerebrospinal fluid (CSF) has been recognized recently as an important contributor to the neurogenic niche [3, 19, 38]. Trophic factors, transcription factors and guidance molecules are secreted by the choroid plexus (ChPl) into the CSF. We previously showed that sAPPα infused in the CSF recognizes stem cells expressing the EGF-receptor within the SVZ and enhances their proliferation, while no effect on hippocampal neurogenesis was observed [4]. In line with these observations, we aimed to evaluate the importance of the ChPl in regulating adult
neurogenesis and to explore the possibility that the ChPl might constitute an important actor and translational target in healthy and pathological aging. We report that sAPPα produced specifically from the ChPl positively affects proliferation in both neurogenic niches. Conversely, we used serotype 5 adeno-associated virus (AAV5) vectors specifically targeting the ChPl to express human APP bearing the Swedish-Indiana (SwInd) mutations, thus favoring Aβ production, which resulted in reduced proliferation in both niches, caused defects in reversal learning and impaired synaptic plasticity mechanisms in the hippocampus.

Materials and Methods
Animals and ethics
C57Bl/6J mice were purchased from Janvier (France) and Appflx/flx mice were described previously [22]. All colonies were maintained under a 12:12 light/dark cycle with free access to food and water. In environmental enrichment experiments, Appflx/flx mice were placed in cages with running wheels (2 mice per cage) 7 days before surgery and kept for 15 days in these cages before analysis. All animal procedures were carried out in accordance with the guidelines of the European Economic Community (2010/63/UE) and the French National Committee (2013/118). For surgical procedures, animals were anesthetized with xylazine (Rompun 2%, 5 mg/kg) and ketamine (Imalgene 1000, 80 mg/kg) by intraperitoneal injection. This project (no. 00702.01) obtained approval from Ethics committee no. 59 of the French Ministry for Research and Higher Education.

Protein and virus stereotoxic surgery
Vectored Cre recombinase protein, Cre-Tat, was produced as previously described [41]. Adeno-associated virus (AAV) were of serotype 5 and purchased from either SignaGen [AAV5-CMV-eGFP-U6-shRNA(mAβ42) and AAV5-CMV-eGFP-U6-shRNA(scrambled mAβ42)] or Vector Biolabs [AAV5-CMV-eGFP-2A-mAPP[NM_007471.3], AAV5-CMV-mCherry-2A-hAPP(SwInd), and AAV5-CMV-eGFP]. The hAPP(SwInd) sequence is a form of human APP [NM_201414.2] bearing both Swedish (K670N/M671L) and Indiana (V717F) related mutations. Cre-Tat protein (~30 µg), vehicle (protein vehicle is 1.8% NaCl, 15% DMSO; virus vehicle is 0.9% NaCl), or high-titer AAVs (~10^{13} GC/ml) were injected (2 µl per mouse) into the right lateral ventricle (coordinates from bregma: x, -0.58 mm; y, +1.28 mm; z, -2 mm) with a 10 µl Hamilton syringe (0.2 µl/min). For protein infusion (3.5 µg per mouse), sAPPα (Sigma-Aldrich), sAPPβ (Sigma-Aldrich), or vehicle were infused for either 7 or 15 days with 100 µl osmotic mini pumps (Alzet) implanted at the same stereotoxic coordinates as above.

Reversal learning
Spatial memory was assessed by the Morris water maze test [6] in which mice use visual cues to locate an escape platform (9 cm in diameter) in an open circular swimming arena (150 cm in diameter, 40 cm deep) filled with opaque water (Acusol, 20 ± 1 °C). The escape platform was hidden 1 cm below the water surface. Room temperature was kept constant at 24°C and both arena placement and surrounding visual cues were kept fixed during all experiments. Data was acquired by the SMART recording system and tracking software (Panlab). Data processing was automated with NAT (Navigation Analysis Tool), an in-house tool rooted in MATLAB [18]. Mice underwent a two-phase training protocol (Fig. 4e). The first phase consisted of 5 training days, 1 day of rest followed by probe test 1 (initial learning). For the second phase, the platform was moved to a different quadrant, and again training lasted 5 days followed by 1 day of rest before probe test 2 (reversal learning). Training sessions consisted of 4 daily swimming trials (1 h interval between trials) starting randomly from different positions, with each quadrant sampled once per day. For each trial, mice were released at a starting point facing the inner wall.
and given a maximum of 90 s to locate and climb onto the escape platform. Mice that failed to locate the platform within 90 s were guided to it. In either case, mice were allowed to stay on the platform for 30 s. To assess spatial memory, probe tests were performed 24 h after the last training session by tracking mice for 60 s with the platform absent. Measured parameters during the training phases were the average time taken to reach the platform (i.e., mean escape latency) and the average speed. Measured parameters during probe tests were the mean distance traveled by the mouse before arriving at the platform location, % time spent in the target quadrant, and the number of platform location crossings.

**Electrophysiology**

Acute transverse hippocampal slices (400 μm) were prepared as previously described [29]. Briefly, mice were culled by cervical dislocation and decapitation. Brains were rapidly removed and placed in chilled (1-4 °C) artificial cerebrospinal fluid (aCSF) composed of (in mM) 119 NaCl, 2.5 KCl, 0.5 CaCl₂, 1.3 MgSO₄, 1 NaH₂PO₄, 26.2 NaHCO₃ and 11 glucose. After sectioning, hippocampal slices were maintained at room temperature in a storage chamber containing aCSF saturated with 95% O₂ and 5% CO₂ for at least 1 h before the experiments. Slices were then transferred to a submerged recording chamber mounted on a Scientifica SliceScope Pro 6000 microscope equipped for infrared-differential interference (IR-DIC) microscopy and were perfused with aCSF (2 ml/min) at RT. All experiments were performed in CA1 stratum radiatum region of the hippocampus. Field excitatory postsynaptic potentials (fEPSPs) were recorded with glass pipettes (2–5 MΩ) filled with 1 M NaCl. Postsynaptic responses were evoked by stimulating Schaffer collaterals (0.033 Hz) in CA1 stratum radiatum with aCSF-filled glass pipettes. Input/output relationships of evoked excitatory postsynaptic potentials (EPSPs) were assayed by incrementing stimulation strength (5 to 50 μA, 100 μs). The test-shock used in subsequent experiments was chosen to elicit 50% of the maximal slope. Paired-pulse experiments consisted of 2 identical stimuli with increasing inter-pulse intervals (50 to 250 ms). Paired-pulse ratios were generated by plotting the maximum slope of the second fEPSP as a percentage of the first. Long-term potentiation (LTP) was induced by high-frequency stimulation (HFS: 2 trains of 100 pulses at 100 Hz, 30 s inter-train interval).

**PCR and Western blots**

Mice were anesthetized for intracardiac perfusion with PBS. Subsequently, choroid plexus, hippocampus, cortex, retina and/or SVZ samples were extracted in ice-cold PBS, frozen on dry ice, and stored at -20 °C. Genomic DNA, total RNA and proteins were recovered by using the Allprep DNA/RNA/Protein Mini Kit (Qiagen 80004). The efficiency of Cre-induced recombination in Applox/lox mice was verified by PCR with primers F, C and D previously described [22]. For quantitative RT-PCR, cDNA was synthesized from 13 ng total RNA with QuantiTect Reverse Transcription Kit (Qiagen 205313). Quantitative PCR reactions were carried out in triplicate with SYBR Green I Master Mix (Roche S-7563) on a LightCycler 480 system (Roche). Expression was calculated by using the 2^ΔΔCt method with Gapdh as a reference. For Western blot analysis, proteins samples were separated on NuPAGE 4-12% Bis-Tris pre-cast gels (Invitrogen NP0321) at 200 V for 1 h and transferred onto a methanol-activated PVDF membrane at 400 mV for 1 h. Primary antibody anti-APP (22C11 mouse 1:500, Millipore MAB348) was incubated overnight at 4 °C and anti-mouse-HRP-coupled secondary antibody (1:3000, Life Technologies) was incubated for 1.5 h at RT. Signal was detected by SuperSignal West Femto Substrate (Thermo Scientific 34095) with a LAS-4000 gel imager (Fujifilm) and quantified by densitometry with ImageJ.
Histology

Mice were anesthetized for intracardiac perfusion with PBS, and brains were subsequently removed and fixed in 4% paraformaldehyde for 3 days. Coronal sections (60 μm) were obtained by Vibratome (Microm). For immunostaining, floating sections were steamed 10 min in citrate buffer (10 mM sodium citrate pH 6.0, 0.05% Tween), washed in PBS and treated with blocking solution (PBS pH 7, 1% triton-X100, 5% normal goat serum) for 1 h. Primary antibody anti-Ki67 (SP6 rabbit 1:500, Abcam ab16667) was incubated overnight at 4 °C in blocking solution and biotinylated anti-rabbit IgG secondary antibody (1:3000, Vector Laboratories BA-1000) at RT for 1 h. Staining was visualized by using the Vectastain ABC HRP kit (Vector Laboratories PK-6100) with the DAB Peroxidase (HRP) Substrate Kit (Vector Laboratories SK-4100). Ki67-positive cells were counted manually in the DG on 5 sections per mouse (sections 180 μm apart, n = 5 to 8 mice per group). Cells in SVZ were counted by Stereoinvestigator (Stereology Software, MBF Bioscience) on 8 sections per mouse (sections 180 μm apart, n = 5 to 8 mice per group).

Statistical Analysis

Morris water maze training data (escape latency and speed) were analyzed with two-way repeated measures ANOVA with Statview 5.0.1 (SAS). Probe test data were analyzed for normality (D’Agostino & Pearson omnibus normality test) and unpaired t test was used for group comparisons with Statview 5.0.1 (SAS). Electrophysiological data were analyzed by ANOVA with Statistica 6.1 and Statview 5.0.1. Histological and biochemical data were analyzed by unpaired t test or ANOVA (as described in Figure legends) with Prism v6 (GraphPad). All data are given as mean ± standard error of the mean (SEM).

Results

**App is highly expressed in the adult choroid plexus**

Given that ventricular infusion of recombinant sAPPα has been shown to affect SVZ proliferation [4, 8], and that the ChPl has been shown to express APP [21], we hypothesized that the ChPl is a potential endogenous source for CSF-borne sAPPα. Indeed, App is one of the most highly expressed ChPl genes [2]. We confirmed high App expression levels in the ChPl by qPCR. In comparison to the hippocampus (that has been shown previously to strongly express APP) and the SVZ, we found higher and elevated levels in ChPl in lateral and 4th ventricles of 4-month-old adult mice (Fig. 1a). Interestingly, Transthyretin and Apoe, two genes involved in App functions [7, 40] are also very highly expressed in the ChPl [2, 39].

**App knock-down in the choroid plexus decreases adult proliferation**

In order to knock down App expression locally and specifically in ChPl, we performed Cre-Tat intracerebroventricular (icv) injections in 10-month-old App^flox/flox^ mice (Fig. 1b). Cre-mediated deletion of the App locus and App expression were evaluated 15 days later in various structures including retina, hippocampus, cortex and ChPl (Fig. 1b-f). The specificity of Cre-Tat targeting selectively the ChPl was confirmed by the absence of recombination in other structures (Fig. 1c), as previously demonstrated for another floxed mouse model [41]. Consequently, only in ChPl do we observe a decrease in App mRNA (by 37 ± 7%, Fig. 1d) and APP protein (by 52 ± 3%, Fig. 1e-f).

To evaluate the impact of App knock-down on cell proliferation, ~3-month-old App^flox/flox^ mice were injected with vehicle or Cre-Tat and subsequently implanted with 15-day osmotic minipumps for CSF infusion of either vehicle or sAPPα (Fig. 1g). Compared to vehicle injected/infused controls, animals injected with Cre-Tat and infused with vehicle showed a significant reduction in the number of proliferating cells in the SVZ (Fig. 1h). This decrease
was rescued by infusion of sAPPα (Fig. 1h), suggesting that knock-down of sAPPα leads to impaired neurogenesis. In contrast, this experimental paradigm did not alter DG proliferation (Fig. 1i-j), even under enriched environment conditions with free access to running wheels known to stimulate hippocampal proliferation [46].

In order to reduce App levels more robustly in the ChPl, a viral vector expressing shRNA against the mouse βA42 sequence was injected into the ventricles of ~3-month-old wild type mice (Fig. 2a). As previously reported [48], icv injection of AAV5 results in specific ChPl targeting, and indeed we did not detect expression of co-expressed eGFP marker protein in the parenchyma (Fig. 2b). Eight weeks post-injection, App (mRNA) and APP protein decreased nearly 5-fold in ChPl with no change in hippocampus and cortex levels (Fig. 2c-e). This decrease led to a significant reduction in the number of proliferating cells in both SVZ and DG (Fig. 2f-i).

**sAPPα gain-of-function in either cerebrospinal fluid or choroid plexus increases adult proliferation**

To further investigate the impact of sAPP on adult neurogenesis, either sAPPα or sAPPβ was infused for 7 days in the lateral ventricles of ~3-month-old wild type mice (Fig. 3a). APP can be cleaved either along the non-amyloidogenic pathway to give rise to sAPPα or along the amyloidogenic pathway to produce its sAPPβ counterpart, and the icv injection of the two sAPP forms has been previously shown to have opposite effects on neurogenesis [8]. In our paradigm, infusion of sAPPα but not of sAPPβ increased the number of proliferating cells both in the SVZ and DG of adult mice (Fig. 3b-e). To confirm that CSF and choroid plexus sAPPα gain-of-function are correlated, icv injections of AAV5 expressing mouse App were performed for ChPl-specific over-expression (Fig. 3f). After 8 weeks, the relative App expression was increased by approximately 2-fold (Fig. 3g), resulting in a significant increase in the number of proliferating cells in both the SVZ and DG (Fig. 3h-i).

**Choroid plexus expression of APP(SwInd) impairs behavior**

Because altering APP expression selectively in the ChPl impacted neurogenic niches, we hypothesized that favoring the production of Aβ specifically in ChPl could negatively affect niche functions. To explore this possibility, we overexpressed a mutated form of human APP (Swedish K670N/M671L and Indiana V17F mutations) specifically in the ChPl of wild type mice at 3 months of age (Fig. 4a). The consequences of this gain-of-function were evaluated 3 and 12 months after injection (when mice were 6 and 15 months old, respectively). We found strong hAPP(SwInd) mRNA expression in ChPl after 3 months which increased more than 3-fold by 12 months (Fig. 4b). Strikingly, proliferation was significantly decreased at 3 months post-injection in both SVZ and DG of hAPP(SwInd) expressing mice. However, no difference was seen at 12 months post-injection (Fig. 4c-d), although this is likely due to the extremely low levels of proliferation also observed in control mice that may preclude further reduction. Indeed, proliferation already dramatically decreases from the time of injection at 3 months (‘No injection’ in Fig. 4c-d) to 6 months (3 months post-injection) in both niches, and further decreases in the DG at 15 months (12 months post-injection) as previously reported [9]. This decrease in proliferation was not due to amyloid plaque accumulation, as none was observed in the cortex or hippocampus of mice expressing the human APP(SwInd) in the ChPl (data not shown).

We also evaluated spatial memory by using the Morris water maze reversal learning paradigm at either 3 months or 12 months post-injection (Fig. 4e). Mice from either group had not been subjected to previous spatial memory tests. During the learning phases, all groups showed significant improvement in latency to find the hidden platform after 5 days of training, but no differences were observed between groups (control and hAPP(SwInd)) in either latency or
swimming speed at both ages (Supp. Fig. 1). During probe tests, platforms were removed and performance was evaluated by the time spent in the target platform quadrant, the number of platform area crossings, and the mean total distance taken to reach the platform area (Fig. 4f-h). At either 3 months or 12 months post-injection, hAPP(SwInd) mice showed no difference in performance compared to controls during probe test 1 (initial learning). Performance after probe test 2 (reversal learning) was unchanged in 6-month old mice (3 months post-injection) but showed significant differences for all parameters at 15 months (12 months post-injection). The time spent in the trained target quadrant was decreased, as was the number of platform area crossings, while the mean distance to find the platform area was increased, indicating that reversal learning is impaired after long-term ChPl expression of hAPP(SwInd).

**Choroid plexus expression of App(SwInd) impairs synaptic plasticity**

Compromised reversal learning may be rooted in plasticity-dependent deficits in hippocampal-dependent spatial memory. Interestingly, mouse models of AD show a marked decrease in hippocampal CA1 synaptic plasticity in the form of long-term potentiation (LTP) at 4 months of age [20]. We thus assessed synaptic function at Schaffer collateral CA3 to CA1 synapses in the hippocampus of 15-month old mice expressing hAPP(SwInd) in the ChPl after receiving AAV5 icv injections at 3 months of age (Fig. 5a). We found no change in basal synaptic transmission in CA1 as shown by comparable input/output curves (F>1) (Fig. 5b). However, paired-pulse facilitation, a form of short-term plasticity reflecting presynaptic function, is increased in mice expressing mutated APP thus indicating a decrease in pre-synaptic release probability (Fig. 5c). Consistently, induction of LTP by high-frequency stimulation is significantly impaired in AAV5-hAPP(SwInd) injected mice (Fig. 5d), revealing impaired plasticity which corroborates observed deficits in spatial memory (Fig. 4f-h).

**Discussion**

The present study takes its origin in the growing interest for the ChPl, a structure increasingly recognized for its physiological importance beyond its classical “kidney of the brain” functions as it secretes CSF containing a plethora of trophic compounds [13]. Among these compounds are growth factors, guidance cues and morphogens, some of which gain access to the SVZ and participate in the regulation of neuroblast migration [11, 31, 39]. sAPPα infused into the CSF has been shown to increase proliferation and progenitor cell numbers in the SVZ [4, 8], with binding sites on type C and type A progenitor cells [4]. Given the very high expression of APP by the ChPl, we hypothesized that sAPPα could be secreted by the ChPl into the CSF and participate in adult neurogenesis. By conditionally knocking down *App* expression specifically in the ChPl through either genetic deletion or shRNA expression, we establish that ChPl APP has neurogenic activity not only in the SVZ but surprisingly also in the DG. Furthermore, this reduced proliferation can be reversed by the infusion of sAPPα into the CSF. Thus, sAPPα secreted by the ChPl can not only gain access to cells within the “superficial” SVZ, which contact lateral ventricles, but also to cells within the “deeper” DG within the parenchyma. Such transport has been reported for transcription factors secreted into the CSF by the ChPl [41]. Together, these results strongly suggest that sAPPα secreted by the ChPl functions as a neurogenic factor, further confirming ChPl as a major actor in regulating adult neurogenesis. The full range of functions of the APP family have yet to be fully identified, but several studies strongly suggest important developmental and physiological roles [28]. Some functions involve *cis* or *trans* dimer interactions between transmembrane APP molecules, and others involve heterodimers between transmembrane APP and secreted sAPPα [23]. These interactions modulate important processes including cell adhesion, synapse formation, synapse stabilization and cell survival [28]. Full-length transmembrane APP or sAPPα also interact with a large
number of other trans and cis interactors, thus acting as ligands or receptors in a variety of signaling pathways [28]. Some of these properties are attributed to the extracellular C-terminal 16 amino acids of sAPPα, as compared to sAPPβ. For example, acute in vitro or in vivo application of sAPPα but not sAPPβ rescues hippocampal LTP in the adult brain of conditional double knockout mice lacking APP and the related protein APLP2 [17, 33]. This C-terminal sequence facilitates synaptic plasticity in the hippocampus through binding to functional nicotinic α7-nAChRs [26, 33]. In keeping with previous results [8], we also find that in vivo icv infusion of sAPPα, but not sAPPβ, positively impacts neurogenic niche cell proliferation in the adult mouse brain. While the reason for this functional divergence between sAPPα and sAPPβ for proliferation remains unknown, a shift in APP processing towards the amyloidogenic pathway would clearly impair both synaptic plasticity and neurogenesis.

Mouse models developed to analyze the role of specific human APP mutations typically employ mutated genes often expressed throughout the brain and even the body. Our study is unique in that it explores whether expressing mutated APP only in the ChPl of adult mice (3 months old) could affect neurogenic niche proliferation and trigger learning defects. We found that ChPl expression of the human APP(SwInd) mutant, which favors sAPPβ and Aβ peptide formation, decreases proliferation at 3 months after infection (6 months old) with no change at 12 months after infection (15 months old), probably due to the fact that neurogenesis is naturally low at 15 months and cannot be further reduced. However, a reversal learning test showed a significant decrease in the cognitive performance of these mice at 15 months. Accordingly, electrophysiological analysis at 15 months revealed decreased pre-synaptic release probability and impaired synaptic plasticity. No amyloid plaques were observed in these mice, leading us to speculate that the functional outcome of increased sAPPβ and Aβ accumulation is rooted in either synaptic deficits, reduced neurogenesis, and/or neuronal aging [42]. Given that sAPPβ icv infusion does not affect proliferation in the neurogenic niches, it is more likely that the observed deficits are due to increased Aβ (soluble or oligomers). Finally, while we observe no change in proliferation at 15 months, we cannot discount a negative long-term effect of ChPl APP(SwInd) on the shaping of hippocampal synaptic circuits due to impaired proliferation during aging.

Our findings strengthen the unique role played by the ChPl in regulating adult neurogenesis, and it is important to consider the morphological and transcriptomic alterations of the ChPl during normal aging and in late-onset AD that can impact brain homeostasis [35-37, 39]. These alterations can lead to changes in blood-CSF barrier properties, in ChPl function regulation, as well as in CSF composition and turnover. Furthermore, adult neurogenesis failure clearly plays a role in the development of AD in familial and possibly sporadic AD patients [14, 15, 25, 27, 34, 45] and could be in part responsible for the decrease in neurogenesis observed in AD patients [25]. From a translational viewpoint, the fact that expressing a mutated APP gene exclusively in the ChPl can alter the cognitive abilities of mice raises the possibility that modifying the expression of APP or targeting APP mutations specifically in the ChPl, a structure accessible from the venous compartment [41], may represent a novel means to alleviate the burden associated with AD.

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**Figure Legends**

**Figure 1.** Choroid plexus App loss-of-function in App<sup>flx/flx</sup> mice decreases adult neural progenitor proliferation.

- **a** Quantitative PCR analysis of App expression in ChPl from ventricles (LV and 4V), SVZ, and Hc. Values are normalized to GAPDH and expression in Hc (Veh n = 4, Cre-Tat n=7).
- **b** Schematic of App ChPl knock-down model involving single icv injection of Veh or Cre-Tat in adult App<sup>flx/flx</sup> mice.
- **c** App specific recombination in LV choroid plexus after Cre-Tat icv injection. PCR of genomic DNA extracted from indicated structures. Grey arrow indicates App wild type locus, black arrow indicates deleted App locus upon Cre recombination.
- **d** Quantitative PCR analysis of App expression normalized to GAPDH after Cre-Tat or Veh icv injection (Veh n = 10, Cre-Tat n=10).
- **e-f** Western blot analysis (e) of various brain structures after Cre-Tat or Veh icv injection for quantification (f) of APP protein levels normalized to actin (Veh n = 10, Cre-Tat n=10).
- **g** Schematic of sAPPα rescue paradigm for evaluating SVZ cell proliferation. sAPPα was infused by 15-day osmotic mini-pump implanted immediately after Cre-Tat icv injection.
- **h** Analysis of SVZ cell proliferation by quantification of Ki67 positive cells after Veh or Cre-Tat icv injection followed by infusion of Veh or sAPPα (Veh/Veh n=7, Cre-Tat/Veh n= 6, Cre-Tat/sAPPα n=7).
- **i** Schematic of App ChPl knock-down with environmental enrichment (wheeling run) for evaluating DG cell proliferation.
- **j** Analysis of DG cell proliferation by quantification of Ki67 positive cells after App recombination in mice in normal conditions or after physical exercise (Veh n=7, Cre-Tat n=8).

*p*<0.05; **p**<0.01; *t* test in **d**, **f**, and **j**; one-way ANOVA in **h**; all values, mean ±SEM. LC, lateral ventricle; 4V, 4<sup>th</sup> ventricle; ChPl, choroid plexus; Hc, hippocampus; Cx, cortex; Ret, retina; SVZ, subventricular zone; DG, dentate gyrus; icv, intracerebroventricular; Veh, vehicle.

**Figure 2.** Knock-down of choroid plexus App expression reduces cell proliferation in neurogenic niches.

- **a** Schematic of App ChPl knock-down model involving a single icv injection of AAV5 expressing shRNA targeting the mouse Aβ42 sequence in wild type mice. Control virus expresses a scrambled Aβ42 shRNA devoid of targets in mouse.
- **b** An icv injection of AAV5 leads to specific choroid plexus expression. Note the absence of eGFP in brain parenchyma. Scale bar, 80 µm.
- **c** Quantitative PCR analysis of App expression in the ChPl, Hc, and Cx after ChPl App knock-down (shRNA-mAβ42 n=4, shRNA-scrambled n=4).
- **d-e** Western blot analysis (d) of various brain structures after ChPl App knock-down for quantification (e) of APP protein levels normalized to actin (shRNA-mAβ42 n=4, shRNA-scrambled n=4).
- **f-i** Analysis of cell proliferation in SVZ (f,g) and DG (h,i) by quantification of Ki67 positive cells after ChPl App knock-down (shRNA-mAβ42 n=7, shRNA-scrambled n=5). Arrows (f) highlight regional differences in SVZ cells. Scale bars, 100 µm.

**t** test; **p**<0.01; ***p**<0.001.
Figure 3. sAPPα gain-of-function in the cerebral ventricles increases cell proliferation in neurogenic niches.

a Schematic of sAPP icv infusion by 7-day osmotic mini-pump in adult wild type mice.
b-c Analysis of cell proliferation in SVZ (b) and DG (c) by quantification of Ki67 positive cells after icv infusion of sAPPα or Veh (Veh n=9, sAPPα n=9).
d-e Analysis of cell proliferation in SVZ (d) and DG (e) by quantification of Ki67 positive cells after icv infusion of sAPPβ or Veh (Veh n=5, sAPPβ n=5).
f Schematic of App ChPl gain-of-function model involving a single icv injection of AAV5 expressing mouse App or eGFP (control) in wild type mice.
g Quantitative PCR analysis of App expression in ChPl after AAV5 icv injection (AAV5-mAPP n=4, AAV5-eGFP n=3).
h-i Analysis of cell proliferation in SVZ (h) and DG (i) by quantification of Ki67 positive cells after icv injection (AAV5-mAPP n=6, AAV5-eGFP n=5).
*p<0.05; **p<0.01; ***p<0.001; t test; all values, mean ± SEM. ChPl, choroid plexus; Hc, hippocampus; SVZ, subventricular zone; DG, dentate gyrus; icv, intracerebroventricular; Veh, vehicle.

Figure 4. Choroid plexus hAPP(SwInd) expression decreases proliferation in neurogenic niches and impairs reversal learning.

a Schematic of mutant APP ChPl model involving a single icv injection of AAV5 expressing human mutated APP or eGFP (control) in wild type mice.
b Quantitative PCR analysis of ChPl viral expression of hAPP(SwInd) in WT mice at 3- and 12-months post-injection (n=6 per group).
c-d Quantification of Ki67 positive cells in SVZ (c) and DG (d) after ChPl viral expression (AAV5-eGFP n=6, AAV5-hAPP(SwInd) n=12).
e Reversal learning paradigm.
f-h Behavioral responses after probe test 1 (Initial) and probe test 2 (Reversal) quantified by time (f), activity (g) and distance (h) (6 months: AAV5-eGFP n=9, AAV5-hAPP(SwInd) n=11; 15 months: AAV5-eGFP n=11, AAV5-hAPP(SwInd) n=10).
*p<0.05; **p<0.01; ***p<0.001; t test; all values: mean ± SEM. SVZ, subventricular zone; DG, dentate gyrus; NE, not expressed; icv, intracerebroventricular.

Figure 5. Choroid plexus hAPP(SwInd) expression impairs hippocampal LTP.
a Schematic of mutant APP ChPl model involving a single icv injection of AAV5 expressing human mutated APP(SwInd) or eGFP (control) in wild type mice.
b Comparaison of stimulus-response (input/output) relationship in CA1 region between AAV5-hAPP(SwInd) (N=5, n=13, black squares) and AAV5-eGFP (N=7, n=10, red dots) injected mice.
c Comparison of paired pulse facilitation between AAV5-hAPP(SwInd) (N=6, n=13) and AAV5-eGFP (N=8, n=13) injected mice.
d LTP was induced by high frequency stimulation (HFS) at Schaffer collateral-CA1 synapses after 20 min baseline recording of slices from mice injected with AAV5-hAPP(SwInd) (N=4, n=9) or AAV5-eGFP (N=5, n=7). Representative traces showing responses before (dashed line) and 60 min after tetanus delivery (bold line).
*p<0.05; ANOVA; all values: mean ± SEM; N, number of animals; n, number of slices; icv, intracerebroventricular. Calibration bars: 10 ms, 0.2 mV.
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Figure 1

(a) ChPl Hc Cx Ret. 0.0 0.5 1.0 1.5 App expression (cf. GAPDH)

(b) icv inj Veh or Cre-Tat Ac-C Cre-Tat

(c) ChPl Hc Cx Ret. 0.0 0.5 1.0 1.5 2.0 APP mRNA

(d) ChPl Hc Cx Ret. 0.0 0.5 1.0 1.5 2.0 APP mRNA (cf. GAPDH)

(e) ChPl Hc Cx Ret. 0.0 0.5 1.0 1.5 2.0 APP protein (cf. actin)

(f) ChPl Hc Cx Ret. 0.0 0.5 1.0 1.5 2.0 APP protein (cf. actin)

(g) icv inj Veh or Cre-Tat Adult + icv sAPPα infusion

(h) Cre-Tat - + + sAPPα - - + SVZ: Ki67+ cells 0 10000 20000 30000 40000

(i) icv inj Veh or Cre-Tat Adult + enrichment

(j) DG: Ki67+ cells 0 20 40 60 80 100

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Figure 2

Virus:
- AAV5-eGFP-U6-shRNA(scrambled)
- AAV5-eGFP-U6-shRNA(mAβ42)

Viruses:
- Scrambled shRNA(mAβ42)
- mAβ42
- Scr

mAβ42 expression (d.GAPDH)

Ki67+ cells

APP protein (cf actin)

Adult wild type mice

icv inj AAV5

LV

Scrambled

DG
Figure 3

(a) icv protein infusion

(b) 7 d icv inj AAV5

(c) Ki67+ cells

(d) Ki67+ cells

(e) Ki67+ cells

(f) icv inj AAV5

(g) APP695 expression (ΔΔCt, GAPDH)

(h) Ki67+ cells

(i) Ki67+ cells

Infusion:
- Veh
- sAPPα
- sAPPβ

Viruses:
- AAV5-CMV-eGFP
- AAV5-CMV-eGFP-2A-mAPP

Adult wild type mice

Adult wild type mice

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Figure 4

Viruses:
- AAV5-CMV-eGFP
- AAV5-CMV-mCherry-2A-hAPP(SwInd)

b) Human APP expression (cf. GAPDH)

c) DG: Ki67+ cells

** Months post-injection

Init. learning

Test 1

Day 6

Day 13

Test 2

Rev. learning

Days 8 to 12

Probe

f) Time in target quadrant (%)

Init. learning

Test 1

Day 6

Day 13

Test 2

Rev. learning

Days 8 to 12

Probe

h) Mean distance (cm)

Init. learning

Test 1

Day 6

Day 13

Test 2

Rev. learning

Days 8 to 12

Probe
Figure 5

(a) icv inj AAV5  
Electrophysiological test  
Adult wild type mice  

Viruses:  
- AAV5-CMV-eGFP  
- AAV5-CMV-mCherry-2A-hAPP(SwInd)

(b)  
Stimulation intensity (µA)  

(c)  
IEPSP  
ISI (ms)  

(d)  
IEPSP (% change)  
Time (min)  

Control hAPP(SwInd)
**Fig. S1** Morris water maze escape latency and speed measured during acquisition phases. 

**a-d** At 3 months post viral injection, recordings performed during the 5 day initial learning phase (**a,c**) or reversal learning phase (**b,d**) to determine displacement speed (**a,b**) and mean escape latency (**c,d**).

**e-h** At 12 months post viral injection, recordings performed during the 5 day initial learning phase (**e,g**) or reversal learning phase (**f,h**) to determine displacement speed (**e,f**) and mean escape latency (**g,h**). Two-way repeated measures ANOVA; all values ±SEM.