Dipeptidyl peptidase 4 contributes to Alzheimer’s disease–like defects in a mouse model and is increased in sporadic Alzheimer’s disease brains

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Abstract

The amyloid cascade hypothesis, which proposes a prominent role for full-length amyloid β peptides in Alzheimer’s disease, is currently being questioned. In addition to full-length amyloid β peptide, several N-terminally truncated fragments of amyloid β peptide could well contribute to Alzheimer’s disease setting and/or progression. Among them, pyroGlu3–amyloid β peptide appears to be one of the main components of early anatomical lesions in Alzheimer’s disease–affected brains. Little is known about the proteolytic activities that could account for the N-terminal truncations of full-length amyloid β, but they appear as the rate-limiting enzymes yielding the Glu3–amyloid β peptide sequence that undergoes subsequent cyclization by glutaminyl cyclase, thereby yielding pyroGlu3–amyloid β. Here, we investigated the contribution of dipeptidyl peptidase 4 in Glu3–amyloid β peptide formation and the functional influence of its genetic depletion or pharmacological blockade on spine maturation as well as on pyroGlu3–amyloid β peptide and amyloid β 42–positive plaques and amyloid β 42 load in the triple transgenic Alzheimer’s disease mouse model. Furthermore, we examined whether reduction of dipeptidyl peptidase 4 could rescue learning and memory deficits displayed by these mice. Our data establish that dipeptidyl peptidase 4 reduction alleviates anatomical, biochemical, and behavioral Alzheimer’s disease–related defects. Furthermore, we demonstrate that dipeptidyl peptidase 4 activity is increased early in sporadic Alzheimer’s disease brains. Thus, our data demonstrate that dipeptidyl peptidase 4 participates in pyroGlu3–amyloid β peptide formation and that targeting this peptidase could be considered as an alternative strategy to interfere with Alzheimer’s disease progression.

Alzheimer’s disease (AD) is the main age-related neurodegenerative disease, the etiology of which remains a matter of questions. One of the main hypotheses referred to as the “amyloid cascade hypothesis” (1) postulates a key role of amyloid β (Aβ) peptide accumulation (2) in various brain zones, including those controlling memory, a psychometric paradigm altered in AD and that worsens as the disease progresses. However, the consistent failure of Aβ-centric clinical trials aimed at reducing Aβ load by immunotherapy or pharmacological targeting of its generating enzyme γ-secretase (3, 4) casts some doubts about the validity of this hypothesis (5). It remains that, undoubtedly, genetic evidence brought a crystal-clear demonstration that amyloid precursor protein (APP)-linked fragments could participate in the pathology. Several APP-linked catabolites distinct from genuine full-length Aβ (flAβ) were shown to account for some of early lesions of AD-affected brains and could well contribute to the pathology (6–11).

Aβ itself is the product of a physiological APP processing (2, 12, 13) and displays physiological/protective functions (14–16) but could be converted into shorter toxic fragments contributing to the pathology (17). Particularly interesting is the pyroGlu3–Aβ (pE3-Aβ) fragment that derives from an N-terminal dipeptidyl truncation of Aβ yielding a Glu3-Aβ (E3-Aβ) peptide that is subsequently cyclized by glutaminyl cyclase (QC) (18, 19). Numerous anatomical and biochemical evidences suggested that QC is involved in the conversion of E3-Aβ into pE3-Aβ, but the most straightforward demonstration of its key role was brought by studies showing that its pharmacological blockade or genetic depletion alleviates AD-like cognitive deficits in an AD mouse model (20).

Dipeptidyl aminopeptidase 4 (DPP4) is an exopeptidase that was formerly described, by fluorogenic dipeptides substrates screening, as an exopeptidase preferentially cleaving X-Pro sequence (21) and hence, was first referred to as a post–proline dipeptidyl aminopeptidase (22). However, it appeared that most of its main natural substrates harbor an N-terminal dipeptide X-Ala. Strikingly, this structural motif corresponds to the N-terminal sequence (Asp-Ala) of the Aβ peptide. Therefore, DPP4 appears as a good enzyme candidate responsible for the rate-limiting removal of this dipeptide. Indeed, few studies showed that DPP4 could, in concert with QC, yield pE3-40/42Aβ, in vitro (23) and that DPP4 inhibitors could prove useful as an AD treatment (24, 25).

Here, we show by MS that human recombinant DPP4 releases the N-terminal dipeptide of synthetic Aβ40 and in agreement, that a DPP4 specific inhibitor potentiates the recovery of flAβ generated by APP-expressing human cells. We show that the pharmacological blockade of DPP4 by its specific
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inhibitor sitagliptin rescues dendrite morphological alterations in organotypic slices derived from mice infected with lentiviruses harboring the APP bearing the Swedish mutation (APPSwe) sequence. Furthermore, both sitagliptin and shRNA directed toward DPP4 reduce the number of Aβ42-positive plaques and Aβ40/42 loads in triple transgenic AD (3xTg-AD) mouse brain. We also establish that DPP4 genetic reduction and/or sitagliptin alleviate cognitive defects assessed by the Morris water maze (MWM) and Barnes maze tests in 3xTg-AD mice. Finally, we document a Braak stage-dependent and transient augmentation of DPP4 activity in a cohort of sporadic AD brains. Altogether, our data bring novel insights supporting the possibility that DPP4 could contribute to AD pathology.

Results

Pharmacological blockade of DPP4 potentiates Aβ full-length recovery in cells

By using antibodies that interact only with the free N-terminal aspartyl residue of Aβ (FCA18, (26)), we previously established that aminopeptidase A (APA) inhibitors potentiate the recovery of flAβ generated by human cells overexpressing APPswe (14). Here, we show as a control that, as expected, flAβ recovery was highly enhanced when neprilysin, one of the main Aβ-degrading enzymes (27, 28), was blocked with its highly specific and potent inhibitor phosphoramidon (Fig. 1). Furthermore, we assessed the potential effect of the aminopeptidase M (PL250) (29), APA (PL302) (30), and DPP4 (P32/98) inhibitors (31). PL302 and P32/98 but not PL250 significantly potentiated the recovery of flAβ (Fig. 1). Interestingly, we observed an additive effect of PL302 and P32/98 (Fig. 1). These data indicate that APA and DPP4 but not aminopeptidase M contribute to the N-terminal truncation of flAβ and that these catalytic events are likely independent.

DPP4 removes the N-terminal dipeptide of Aβ40-An MS approach

We analyzed the capability of human recombinant DPP4 (rDPP4) to generate Aβ3-40 from synthetic Aβ40. By MS analysis, we found that DPP4 converts Aβ1-40 synthetic peptide into Aβ3-40, thus releasing the N-terminal dipeptide of Aβ40 (Fig. 2, A and B). Importantly, the Aβ3-40/Aβ1-40 ratio

Figure 1. Effects of pharmacological modulation of endogenous DPP4 in HEK APPwt cells on full-length Aβ expression. Full-length Aβ immunoreactivity in HEK APPwt cells treated without (control) or with phosphoramidon (PA), PL302, PL250, and P32/98 (aminopeptidase A, aminopeptidase M, and DPP4 inhibitors, respectively) was quantitated by densitometry. Values are expressed as the percentage of control untreated cells (taken as 100) and are the means ± SEM of 6 to 17 determinations obtained from seven independent experiments. *p < 0.05 and ****p < 0.0001 (Mann–Whitney test). Aβ, amyloid β; APPwt, WT APP; DPP4, dipeptidyl aminopeptidase 4; flAβ, full-length Aβ; HEK, human embryonic kidney.
Figure 2. DPP4 releases the N-terminal dipeptide of Aβ40. HPLC chromatograms of Aβ1-40 (A) and Aβ3-40 (B) peptides and their spectral deconvolution revealing m/z values (4327.1515 and 4141.0908, respectively) recovered after a 6-h incubation of human recombinant DPP4 with synthetic Aβ40. C, Aβ3-40/Aβ1-40 ratio is represented over time in presence or absence of rDPP4, with or without p32/98 inhibitor. D, quantification of Aβ3-40/Aβ1-40 ratio observed after 8 h. Values are the means ± SEM of 5 to 6 determinations obtained from three independent experiments. (**p < 0.01; one-way ANOVA with Dunn’s multiple comparisons post-test). Aβ, amyloid β; AA, average area; DPP4, dipeptidyl aminopeptidase 4; ns, not statistically significant; RT, retention time; SN, signal noise.
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increased with time and was prevented by the DPP4-specific inhibitor p32/98 (31) (Fig. 2, C and D).

DPP4 inhibition rescues alterations of dendritic spine morphology

We have evaluated the putative impact of DPP4 on spine morphology of hippocampal organotypic slices infected with lentiviruses expressing green, WT APP, or APPswe-green constructs (see expressions in Fig. 3B, upper panels). The spine morphology (Fig. 3B, lower panels) was classified within three categories: mature (mushroom/thin, Fig. 3C, red bars), stubby (Fig. 3C, yellow bars), and immature spines (filopodia, Fig. 3C, green bars). First, of note, the expression of the various constructs did not affect neuronal viability as assessed by NeuN expression (Fig. S1). APPswe-expressing slices show a lower number of mature spines than WT APP (APPwt) and green, and a concomitant increase of dendritic filopodia (compare APPwt NT and APPswe NT conditions in Fig. 3C). Of importance, the morphological alterations observed in APPswe-expressing slices were fully rescued by pretreatment with the highly potent and selective DPP4i inhibitor sitagliptin (32), (Fig. 3, B and C, compare APPswe NT versus Sita). This suggested the possibility that DPP4-mediated sitagliptin-sensitive production of pE3-Aβ could account for the alteration of dendritic spines in these organotypic slices. To support this view, we checked whether organotypic slices harbor functional DPP4 and if they produce pE3-Aβ. Our data show that indeed, organotypic slices express pE3-Aβ-like in situ immunoreactivity (Fig. S2A) that is present in insoluble fractions prepared from these slices (Fig. S2B) and display sitagliptin-sensitive DPP4-hydrolyzing activity (Fig. S2C). It is of importance to note that our recent study indicated that synthetic pE3-Aβ could indeed alter dendritic morphology and mimic that triggered by APPswe expression (33). Overall, this set of data indicates that DPP4 contributes to the morphological alterations affecting spines in an ex vivo AD model.

Genetic reduction and pharmacological blockade of DPP4 reduce pE3-42Aβ and Aβ plaque loads

We attempted to reduce DPP4 expression and activity by shRNA approach. We designed several shRNA probes targeting DPP4 (shDPP4) as well as an shRNA corresponding to a control scramble sequence (shScr). We first examined their ability to lower DPP4 expression and activity in N2a cells. Among them, one sequence (see Experimental procedures) was selected because its transfection reduced the number of DPP4-positive N2a cells by about 30% when compared with shScr (Fig. S3, A and B). Then, we produced lentiviruses harboring the shScr and shDPP4 sequences coupled with GFP. This allowed to confirm that stereotaxical delivery of shRNAs adequately yielded GFP expression in the subiculum of the hippocampus of mouse brains (Fig. S3C). shDPP4 infection triggers an about 25% reduction of DPP4 activity in WT mouse brain (Fig. S3, D and E).

We have examined the biochemical and anatomical impacts of DPP4 gene reduction in 3xTg-AD mice infected or not at the age of 3 months with the above-selected shDPP4- or shScr-expressing lentiviruses. At the age of 12 months, DPP4 down-regulation drastically reduces the number of Aβ42-positive plaques (Fig. 4, A and B) without affecting their mean perimeter (Fig. 4C) and areas (Fig. 4D). Interestingly, we also observed a decreased expression of pE3-Aβ in 3xTg-AD mice infected with shDPP4 (Fig. S4). Accordingly, as expected at this age, both Aβ40 and Aβ42 recoveries were much higher in the insoluble fraction (which mostly contains Aβ in an aggregated state) prepared from shScr-3xTg-AD than in shScr-WT infected mice (Fig. 4, E and F). However, both Aβ species were drastically lowered in these insoluble fractions upon shDPP4 infection (Fig. 4, E and F).

We attempted to examine the influence of chronic treatment with sitagliptin. At first, we aimed at assessing whether intraperitoneal administration of sitagliptin could significantly inhibit cerebral DPP4. Indeed, we show that an 8-day treatment of 3-month-old 3xTg-AD mice triggers a reduction of about 30% of cerebral DPP4 activity (Fig. S5). Of note, the extent of sitagliptin-mediated inhibition of central DPP4 was similar to the one achieved by shDPP4 (Fig. S3, D and E). Although partial, this inhibition could however reflect full blockade of DPP4. Thus, we previously established that Gly-Pro–7-amido-4-methylcoumarin undergoes DPP4-unrelated cleavages in crude tissues (34) that could account for shDPP4 and sitagliptin-resistant hydrolyzing activity. This was performed at an earlier age, when Aβ42-positive plaques are clearly less numerous than at the age of 12 months. Thus, the quantification of the number of plaques was more difficult and suffers from higher randomness. Nevertheless, although it did not reach statistical significance, we observed a consistent reduction of Aβ42-positives plaques (Fig. 5, A and B), mean area (Fig. 5C) and perimeter (Fig. 5D). In agreement with a genuine reduction of plaques number, Aβ42 was recovered in a lower amount in the insoluble fraction prepared from sitagliptin-treated 3xTg-AD mice (Fig. 5F), while Aβ40 lowering did not reach statistical significance (Fig. 5E). Overall, our combined genetic and pharmacological approaches indicate that DPP4 down-regulation lowers Aβ-positive plaque formation and Aβ aggregation in 3xTg-AD mice.

Genetic and pharmacological reduction of DPP4 partly alleviates learning and memory defects in 3xTg-AD mice

To determine whether DPP4 activity could modulate memory and learning in vivo, we used the 3xTg-AD mouse model that gathers several hallmarks of AD pathology, that are, senile plaques, neurofibrillary tangles, memory deficit, and synaptic plasticity defects (35). We used two well-known spatial learning tests, the MWM and Barnes maze, that are considered as sensitive means to detect memory deficits in these mice (36), to examine the influence of genetic and pharmacological reduction of DPP4 activity. As expected, 3xTg-AD mice were affected in their learning ability as reflected by a higher latency to reach the platform than WT mice in the MWM (Fig. 6A). Interestingly, shDPP4 partly rescues these alterations (Fig. 6A). This is further supported by the randomness in the trajectories of shScr-treated 3xTg-AD mice (Fig. 6B, left panel) that was corrected in shDPP4-treated mice.
Figure 3. DPP4 pharmacological blockade influences synaptic morphology. Organotypic slices (see procedure in panel A) were infected with green (B, left panel), APPwt (B, middle panel), or APPswe (B, right panel) lentivirus coexpressing green protein (B) treated or not for 24 h with sitagliptin (100 μM). C. graph represents the percentage of spines corresponding to mushroom (red bars), stubby (yellow bars), and immature (filopodia, green bars) categories and are the means ± SEM of 10 to 22 determinations obtained from two independent experiments. **p < 0.01, ***p < 0.005, ****p < 0.0001, (the Kruskal–Wallis test with Dunn’s multiple comparisons post-test). Scale bars in B upper panels correspond to 300 μm. APPswe, APP bearing the Swedish mutation; APPwt, WT APP; DPP4, dipeptidyl aminopeptidase 4; ns, not statistically significant.
Figure 4. Aβ42-positive plaques and Aβ load are decreased in 3xTg-AD mouse brain after shRNA-mediated reduction of endogenous DPP4. A, immunohistochemical analysis using the anti-Aβ42 antibody in 12-month-old 3xTg-AD mice infected with lentiviruses bearing either shScr or shDPP4. Scale bars in panel A correspond to 500 μm (4×) or 100 μm (10×). B, graph represents the number of Aβ42-positive plaques per square millimeters. C and D, graphs represent the mean perimeter (C) and area (D) of Aβ42-positive plaques. B–D, data are the means ± SEM of 5 (shScr) or 7 (shDPP4)-injected mice (Mann–Whitney test). *p < 0.05. Aβ40 (E) and Aβ42 (F) measured by ELISA in insoluble fractions prepared from 12-month-old 3xTgWT and 3xTg-AD transgenic mice infected with shScr or shDPP4 lentiviruses as in panel A. Values are the means ± SEM of 6 (WTshScr and WTshDPP4) or 14 to 15 (shScr- and shDPP4-injected 3xTg-AD) determinations obtained from two independent experiments and are expressed in pg per mg of tissue. ****p < 0.0001 (Mann–Whitney test). 3xTg-AD, triple transgenic AD; Aβ, amyloid β; AD, Alzheimer’s disease; DPP4, dipeptidyl aminopeptidase 4; shDPP4, shRNA probes targeting DPP4; shScr, shRNA corresponding to a control scramble sequence.
Figure 5. Influence of sitagliptin on Aβ42-positive plaques. A, immunohistochemical analysis using anti-Aβ42 antibody in 10- to 11-month-old 3xTg-AD mice chronically treated or not with sitagliptin. Scale bars in panel A correspond to 500 μm (4×) or 100 μm (10×). B, graph represents the number of Aβ42-positive plaques per square millimeters. Panels C and D represent the mean area (C) and perimeter (D) of Aβ42-positive plaques. B–D, data are the means ± SEM of 6 (control, physiological serum) or 8 (sitagliptin)-treated mice (Mann-Whitney test). Note that although there is a clear apparent reduction of the number of Aβ42-positive plaques, statistical analysis did not reach significance (Mann-Whitney test). Aβ40 (E) and Aβ42 (F) measured by ELISA in insoluble fractions prepared from 10- to 11-month-old WT and 3xTg-AD mice chronically treated with sitagliptin as described in Experimental procedures. Values are means of 4 (WT serum and WT sitagliptin) or 13 to 15 (3xTg-AD serum and 3xTg-AD sitagliptin) determinations obtained from two independent experiments and are expressed in pg of Aβ40/42 per mg of proteins. *p < 0.05; Mann-Whitney test. 3xTg-AD, triple transgenic AD; Aβ, amyloid β; AD, Alzheimer’s disease; ns, not statistically significant.
in probe tests performed at 48 h where treated mice swim in
the adequate north west zone virtually similar to WT mice
(Fig. 6B, right panel). Of note, chronic treatment with sita-
gliptin indicated that pharmacological blockade of DPP4 also
partly rescued the latency to reach the target zone in the
Barnes test, a paradigm that was drastically altered in shScr-
treated 3xTg-AD mice (Fig. S6A). Accordingly, sitagliptin
partly improved the distribution of the number of entries in
every hole in sitagliptin-treated 3xTg-AD mice (Fig. S6B). Of
importance, neither sitagliptin nor shDPP4 (data not shown)
affects locomotor activity or anxiety parameters as measured
by the rotarod and open field test (Fig. S7).

**Figure 6. shDPP4 treatment partly restores learning and memory deficits in 3xTg-AD mouse model.** A, graphical representation of the latency to reach the platform in Morris water maze for 3xTgWT (light green and red curves) and 3xTg-AD (dark green and red curves) mice infected with lentiviruses expressing shScr (light and dark red curves) or shDPP4 (light and dark green curves). B, swimming trajectories of indicated treated mice. Panel C shows the number of entries in each quadrant of the Morris water maze. (WT shScr: n = 11; WT shDPP4: n = 12; 3xTg-AD shScr: n = 9; 3xTg-AD shDPP4: n = 11). *p < 0.05; **p < 0.01; ***p < 0.005 (InVivoStat test). 3xTg-AD, triple transgenic AD; AD, Alzheimer’s disease; DPP4, dipeptidyl aminopeptidase 4; NE, north east; NW, north west; SE, south east; shDPP4, shRNA probes targeting DPP4; SW, south west.

DPP4 activity is increased in brains of patients with AD

It has already been described that pE3-42Aβ is present in
abundance in brains of patients with sporadic and familial AD
(37). Thus, according to our data, we assumed that there
should be an enhanced activity of DPP4 in AD-affected brains.
Here, we compared DPP4 activity in hippocampi of controls
and patients suffering from sporadic AD at different Braak
stages (Table S1). We observed a higher inhibitor-sensitive
DPP4 activity in AD brains than in control patients. This in-
crease appears to be AD Braak stage-dependent, culminated at
stages I-III and then declined to return to control values at
later stages (Fig. 7).
Discussion

The main conclusion of our study is that the reduction of DPP4 could be beneficial in the context of AD. It is supported by five lines of independent *in vitro*, *ex vivo*, and *in vivo* evidence. First, recombinant DPP4 releases the N-terminal dipeptide of synthetic Aβ. Second, inhibition of DPP4 in human cells potentiates the recovery of secreted FlAβ. Third, in APPswe-expressing neurons that display drastic alterations of dendritic spines morphology, pharmacological blockade of DPP4 fully restores a normal phenotype. Fourth, both genetic and pharmacological approaches drastically reduce Aβ42-positive plaques and Aβ40/42 load in insoluble fractions prepared from 3xTg-AD brains. Fifth, shDPP4 and sitagliptin partly rescue learning and memory deficits. Overall, this consistent network of evidence strongly suggests that DPP4 could contribute to the generation of E3-Aβ in AD. It should be noted that DPP4 was documented as the rate-limiting enzyme yielding E3-Aβ amenable to cyclization by QC. These concerted enzymatic events have been previously documented *in vitro* (23).

DPP4 inhibitors have been proposed as means to interfere with cognitive disorders (25). Most of studies deal with type 2 diabetes, a well-recognized risk factor for cognitive dysfunctions and dementia (38). This is mainly due to the propensity of DPP4 to hydrolyze many substrates (39) including GLP-1 (40, 41). GLP-1 is a 30-amino acid peptide which harbors an His-Ala N-terminal dipeptide that fulfills the requirement of DPP4 specificity. Indeed, GLP-1 is one of the preferred substrates of DPP4 (42). Interestingly, although GLP-1 is mainly produced in the intestinal tract, it is also present in the brain and more particularly in the hippocampus and frontal cortex (43). Of note, GLP-1 expression is decreased in patients with AD as well as in AD mouse brain (41). Furthermore, it has been shown that GLP-1 overexpression triggers protective effects and that its DPP4-mediated proteolysis drastically shortens its lifetime (40, 41). Accordingly, DPP4 inhibition favors GLP-1 signaling and improves AD-like cognitive deficits in several AD mouse models (39–41) including the 3xTg-AD mice (41). Our data indicate that, besides GLP-1, DPP4 could also participate to AD pathology by targeting Aβ.

In AD mouse models, several anatomical and functional paradigms including apathy-like phenotype, long term potentiation perturbation, and endolysosomal dysfunction (9, 44, 45) have been shown to occur early and independently of Aβ (7, 46) and, for some of them, are drastically potentiated by γ-secretase inhibitors (7). Recent evidence indicates that the β-secretase-derived fragment of APP (namely C99) could account for these dysfunctions (11). However, at late stage of AD-like pathology, an Aβ-linked contribution to learning and memory defects was observed. The firm demonstration of this Aβ-linked counterpart was brought by the comparison of 3xTg-AD and 2xTg-AD mice (46). The latter mice display similar accumulation of C99 but, unlike 3xTg-AD, they never produce detectable amounts of Aβ (due to the absence of PS1 mutation (46)). Of note, although 3xTg-AD and 2xTg-AD mice display similar early perturbations, the former mice exhibit higher learning and memory defects. Our data indicate that these Aβ-linked alterations of behavior could be accounted for by pE3-Aβ peptides. Of importance, we show that both genetic reduction and pharmacological blockade of DPP4 drastically reduce Aβ42-positive plaques and Aβ42 recovery. This is likely due to the reduction of pE3-Aβ. Thus, it has been shown that very low amounts of pE3-42Aβ are sufficient to serve as Aβ42 seed (47), accelerate its aggregation (23), and thereby, the occurrence of plaques.

**Figure 7.** DPP4 enzymatic activity is transiently increased in sporadic AD-affected brains. DPP4 enzymatic activity was measured by fluorimetry as described in Experimental procedures in human brain samples from different Braak stages patients (see Table S1) (controls: n = 10; AD I-III: n = 9; AD IV-V, n = 10; AD VI: n = 14; *p < 0.05, **p < 0.01, ***p < 0.0005, and ****p < 0.0001). (Two-way ANOVA test with Dunnett’s multiple comparisons test). AD, Alzheimer’s disease; DPP4, dipeptidyl aminopeptidase 4.
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It should be emphasized that cognitive alterations are not fully rescued by genetic or pharmacological downregulation of DPP4. This could be explained by the fact that shRNA DPP4 triggers only a partial reduction of the enzyme activity (see Fig. S1) but not by a poor sitagliptin bioavailability after gavage administration (48). Another likely possibility is that besides DPP4, other activities involved in Aβ truncation could occur. One such candidate could be the exopeptidase APA. Thus, this enzyme displays a high affinity for acidic residues such as the N-terminal aspartyl residue of Aβ (22). This hypothesis is supported by our previous study (14) (and present Fig. 2) showing that APA inhibitors enhanced the recovery of cellular flAβ. Furthermore, our recent work showed that the APA inhibitor RB150 and short hairpinRNA directed towards APA both rescue dendritic morphology, lower pE3-Aβ-positive and Aβ-positive plaques and loads, and improve memory defects in 3xTgAD mice (33). Interestingly, APA and DPP4 inhibitor–induced potentiation of flAβ recovery was additive, indicating that the two enzymes independently target flAβ. Thus, the shRNA DPP4-resistant cognitive alterations could be due to residual bioactive DPP4 and APA. It should be noted that we have recently shown that the reduction of APA in vivo partly alleviates memory deficits in 3xTg-AD mice (data not shown).

DPP4 is increased in sporadic AD-affected brain, but this increase occurs relatively early at Braak stages I-III and returned to virtually control values at later stages. These data agree well with previous studies showing that DPP4 is upregulated in AD brain neurons and is associated with plaques (49) but not at the late demented stage in both frontal and parietal cortices of patients with AD (50). The increase of DPP4 indeed coincides with the appearance of plaques (our unpublished data), and hence, one can envision that the pE3-Aβ generated could well initiate the seeding of Aβ, its aggregation and the occurrence of plaques. This fits well with the observation that only very low amounts of pE3-42Aβ could serve as a seed of Aβ42 aggregation and deposition (47) and that low amounts of pE3-42Aβ are sufficient to induce hippocampal neurodegeneration in transgenic mice (51). Thus, DPP4 transient increase could be sufficient to initiate a vicious cycle by which AD anatomical lesions progress.

**Experimental procedures**

**MS**

Synthetic Aβ40 from Bachem (15 ng/μl) was incubated for various time periods with human recombinant DPP4 (2 ng/μl) (R&D System) or for 6 h with or without the DPP4 inhibitor p32/98 (100 μM). Reactions were stopped after addition of formic acid (0.1%) and then samples were subjected to spectrometry analysis. Briefly, Aβ peptides were separated with UPLC system (Thermo Fisher) on a C18 column in an appropriate gradient. MS data were acquired with a Q-Exactive plus mass spectrometer (Thermo Fisher) operating in the full-scan mode. Finally, Aβ fragments were identified using Xcalibur Quan Browser software (version 4.1.31.9).

**Immunoprecipitation procedure**

APPwt-overexpressing human embryonic kidney 293 cells were grown in Dulbecco’s modified Eagle’s medium (DMEM)–fetal calf serum (FCS) (10%)–penicillin/streptomycin medium. Cells were plated in 6-well dishes and allowed to secrete for 8 h in Opti-MEM/FCS (1%) containing phosphoramidon (10 μM) to prevent Aβ degradation by neprilysin (28, 52), with or without aminopeptidase M (pl250), APA (pl302), and DPP4 (p32/98) inhibitors at 5 μM. After media centrifugation, supernatants were completed with one-tenth of RIPA buffer (Tris 50 mM; pH 7.4, containing NaCl (150 mM), EDTA (1 mM), Triton X100 (1%), deoxycholate (0.5%), and SDS (0.1%)) and incubated overnight with a 100-fold dilution of FCA18 antibody (26) and protein A agarose beads (VWR). Beads were washed twice with RIPA buffer 1× and once with Tris (10 mM, pH 7.5) and subjected to Tris/tricine 16.5% polyacrylamide gels. Proteins were transferred onto nitrocellulose and incubated overnight with the 2H3 monoclonal antibody (anti-Aβ1-12 provided by Dr. D. Schenk, San Francisco, CA) at a 1/1000 dilution. After secondary antibody incubation with a goat anti-mouse peroxidase-conjugated antibody (1/2000 dilution), chemiluminescence was recorded using LAS-3000 (Raytest) and quantifications were performed using the Multi Gauge software.

**Organotypic hippocampal slice preparations and culture**

Organotypic hippocampal slices were prepared from 5- to 7-day-old C57Bl6/Rj mice (Janvier Labs). Brains were quickly dissected to retrieve hippocampi, and then, both hemispheres were grown in Dulbecco’s Modified Eagle Medium (97.5% Earles’ Balanced Salt solution (EBSS) and 2.5% EBSS-Hepes). Slices were transferred on sterile hydrophilic membrane Millicell discs (FHLC01300, Millipore) placed in semi-porous cell culture inserts (0.4 μm, Millipore) containing the culture medium (Minimum Essential Medium Eagle + Glutamax-1 (50%), EBSS (18%), EBSS/D-glucose 13% (5%), penicillin-streptomycin (5000 U/ml, 1%), horse serum (25%), and nystatin (10,000 U/ml, 0.06%). Two hours after plating, slices were infected with lentiviruses encoding for green, APPwt, and APPsw proteins, respectively. Slices were kept at 37 °C, with CO2 (5%) for 9 days before imaging analysis. For sitagliptin treatment, slices were incubated for 24 h at a concentration of 100 μM.

**Imaging of dendrites and quantitation of spine morphology**

Slices were left attached on the Millicell membrane and mounted with the mounting solution, cover-slipped, and dried before imaging. Spine morphology was assessed on an average of 14 different dendrite segments (obtained from two independent experiments) taken with Plan-Apochromat 63×/1.40 oil DIC M27 lens, zoom 3.0, and pinhole at 52 μm. The quantification of spine morphology (mature, stubby, and filopodia) was performed manually using “false” color images ZEN software as previously described (53, 54).
Organotypic slice homogenates and immunofluorometry

After sitagliptin treatment (100 µM, 11 days), two organotypic slices (4 confetti discs per slice) were pooled in 200 µl of Tris 10 mM/RIPA buffer (Tris 10 mM, pH 7.4, containing NaCl (150 mM), EDTA (1 mM), Triton X-100 (1%), deoxycholate (0.5%), and SDS (0.1%)). After 30 min in ice dry, confetti discs were removed and supernatants were mechanically homogenized by Potter. DPP4 activity was measured as described in DPP4 activity measurements (see below). For NeuN immunostaining, organotypic hippocampal slices were fixed with 4% of paraformaldehyde and then permeabilized overnight in PBS containing Triton X-100 (0.4%). Hippocampal slices were incubated in the saturation blocking buffer (PBS/Triton 0.1%-bovine serum albumin 5%) for 8 h and then hybridized with the anti-NeuN antibody (d:1/2500, Abcam, rabbit) in PBS/Triton for 48 h. After three PBS washes, secondary antibodies were added overnight with 4′,6-diamidino-2-phenylindole (DAPI) in PBS/Triton 0.1%. Finally, slices were rinsed in double distilled water and then placed on the glass slide with the aqueous mounting medium. For pE3-xÅB immunostaining and before the permeabilization, organotypic hippocampal slices were included in paraffin with automatic tissue processor and then were deparaffinized with three successive xylene baths (1 h per bath), two ethanol 100%, one ethanol 90%, and one ethanol 70% baths for 5 min. This sample preparation allowed unmasking with formic acid for 10 min. The pE3-xÅB antibody was diluted at 1/50 (IBL international) in PBS/Triton 0.1%.

Cytation imaging

Mouse neuroblastoma N2a cells (ATCC, CCL131) were grown in DMEM FCS (10%)-penicillin/streptomycin medium. Cells were plated in 6-well plates at a density of 100,000 cells/well. Five days after infection, cells were fixed for 20 min with paraformaldehyde (4%), washed three times with PBS, permeabilized for 5 min with Triton X-100 (0.1%), and blocked for 1 h with BSA (5%)/Tween (0.05%). The primary antibody against DPP4 (dilution of 1/500, Abcam ab28340) was added overnight, and then cells were rinsed three times with PBS and incubated for 1 h with the secondary anti-goat antibody (Interchim, dilution of 1/500 + DAPI (1/20,000)). Cells were then rinsed again with PBS 1× and fixed with the VectaMount medium (Vector Laboratories) before Cytation imaging analysis with the Biotek Cytation 5 microscope, which automatically acquired six pictures per well with the same acquisition parameters (×20 magnification, numeric aperture 0.45, brightfield, at following excitation and emission wavelengths, respectively: DAPI (377; 447); GFP (469; 525); Texas Red (586; 647)). We used ImageJ software to quantify the fluorescence intensity Texas Red in GFP-positive cells.

DPP4 activity measurements

Hippocampi were homogenized in Tris buffer (10 mM, pH 7.5). DPP4 activity was measured in 50 µg of proteins, with or without the DPP4 inhibitor P32/98 (100 µM) using Gly-Pro-7-amido-4-methylcoumarin as the substrate (100 µM, Santa Cruz Biotech) in the assay buffer (Tris 50 mM, pH 7.5, a final volume of 100 µl). Fluorescence was recorded at 360-nm excitation and 460-nm emission wavelengths as described (50).

Animals

3xTg-AD (35) and 3xTg-WT (non transgenic) mice were generated from breeding pairs provided by Dr Franck LaFerla (Irvine). Animals were housed with a light/dark cycle (12 h:12 h) and were given free access to water and food. All experimental procedures were in accordance with the European Communities Council Directive of 22 September 2010 (2010/63/EU) and approved by the French Ministry of Higher Education and Research (Project number APAFIS#9645-2017012315473838) and by Côte d’Azur University Animal Care and Use Committee.

Viral production and stereotaxic injections

Lentiviral particles were produced as previously described (55). Viral titers were assessed using p24 ELISA (VPK-107; Cell Biolabs). Green, green-APPwt, and green-APPswe lentiviruses (cloned in the lentiviral vector with an IRES-ZsGreen fluorescent tag (pHAGE-CMV-MCS-IRES-ZsGreen)) (56) were under the control of the CMV promoter. shRNA constructs TCATCACCCTGCCCATAGTTCTGCTGAGC (shDPP4) and GCACTACAGAGCCTCACTAGATAGTT (shScr) were inserted in the vector pGFP-C-sh Lentil under the U6 promoter (Cat. No. T R30023, OriGene). Gene-specific mouse shRNA sequences are TL500553 CD26 (Gene ID 13482) Mouse shRNA under the U6 promoter. Two- to three-month-old WT and 3xTg-AD mice were anesthetized by intraperitoneal injection with a mixture of ketamine (100 mg/kg) and xylazine (10 mg/kg). Mice were bilaterally stereotaxically injected into the subiculum region (3 µl) with high titers of viral particles (1–1.5 × 1010 viral particles) at the following coordinates Sub: a/p: ± 3.8, m/l ± 2.5, d/v: −2.0 (The Mouse Brain in stereotaxic coordinates, Second Edition, Elsevier Academic Press).

In vivo pharmacological treatments

A selective and specific inhibitor of DPP4, sitagliptin (provided by Sigma-Aldrich, PHR1857.14) was administered once a day by intraperitoneal injections at a dose of 15 mg/kg (i.e., 0.3 mg/day for a 20-g mouse) for a total of 6 weeks of treatment before animal sacrifices. To evaluate cerebral DPP4 activity, sitagliptin was administrated daily for 8 days and then DPP4 activity was measured as described above.

Barnes maze and MWM

For Barnes maze, spatial learning and memory were studied using a dry land–based rodent procedure (57). Mice performed during 3 to 4 days on a spatial acquisition protocol, allowing animals to retrieve a box placed under a hole of the Barnes maze apparatus, before being assessed for memory with a probe trial in which mice had a total time of 60 s to find the target place. Several parameters were evaluated including the
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latency to reach the target site, number of quadrant crossing, latency, and distance in the quadrant of the target site.

MWM task was assessed in a circular pool filled with an opaque solution and placed in a room surrounded by visual clues. Maze was performed as previously extensively described (46). Data were analyzed with the ANY-maze 6.1 software.

Rotarod performance and open field tests

Locomotor performance of mice was tested on a rotarod apparatus (Bioseb, model LE8200) as previously described (46). Three trials were performed during a period of 5 min for each animal. Mice anxiety and exploratory behavior were recorded by an Open Field test. Briefly, time spent in the center of the box (40 × 40 cm) is recorded during 10 min per animal.

Immunohistochemistry

3xTg-WT or 3xTg-AD mice infected with lentiviruses or injected with pharmacological treatment were anesthetized as described. Mouse brains were embedded in paraffin and then sliced with a microtome apparatus at a thickness of 8 μm. Slices were treated with formic acid and then incubated overnight at 4 °C with a recombinant anti-beta amyloid 1 to 42 antibody that does not cross-react with Aβ37, 38, 40, and 43 (dilution 1/1000, H31L21, Invitrogen) or with anti-pEA antibody (dilution:1/50, rabbit, IBL international). After several washes, sections were incubated for 1 h with an anti-rabbit horseradish peroxidase. Slices were then revealed with the diaminobenzidine–ImmPACT system (Vector Laboratories).

Human brain sample preparation

Human brain samples were obtained from the “NeuroCEB” Brain Bank. All procedures using human brain samples were performed in accordance with the ethical standards of both institutional and national research committees as well as the 1964 Helsinki declaration and amendments. In accordance with the French Bioethical Agreement (ACB 2013-1887), individual consents were signed by the patients or their close relatives in their name. Cases were anonymized, and information regarding age, sex, and neuropathology is provided in Table S1. Tissue lysates were obtained after mechanical homogenization of temporal cortex sections. Powder was resuspended in Tris 10 mM, pH 7.5, and samples were mechanically homogenized by a Potter, sonicated, and analyzed for DPP4 enzymatic activity as described above.

Insoluble fraction preparations and Aβ quantitation by ELISA

Dissected hippocampi from 3xTg-WT and 3xTg-AD mice or organotypic hippocampal slices were homogenized in RIPA buffer containing a mix of protease inhibitors as previously described (58). Proteins were centrifuged after mechanical homogenization by a Potter (100,000g, 1 h, at 4 °C). Pellets containing insoluble material were resuspended in formic acid (70%), centrifuged (100,000g, 1 h, 4 °C), and then, supernatants were brought to pH 7.5 by addition of Tris HCl (1 M, pH 10.8) containing betaine (25 mM). This fraction was referred to as the insoluble fraction. Human Aβ40 and Aβ42 peptides levels were measured in insoluble fractions using sandwich enzyme-linked immunoasorbent assay kits (BioSource, Invitrogen) as described (59).

Statistical analysis

Organotypic slices and MS were analyzed by one-way ANOVA and the Kruskal–Wallis (*p ≤ 0.05; **p < 0.01; ****p < 0.005) test to study the spine morphology. DPP4 activities were compared by Wilcoxon (*p ≤ 0.05) and Mann–Whitney tests (*p ≤ 0.05). For behavioral experiments, all statistical analyses were realized using InVivoStat software. For each statistical analysis, normal distribution of residuals and homogeneity of variance were analyzed by checking normal probability plot and residual versus predicted plot. For Barnes maze and MWM task, escape latency was analyzed by mixed-model ANOVA with repeated measure (genotype x injection x training day) and followed by Benjamini–Hochberg’s correction. Concerning statistics of probe phase MWM and Barnes maze, open field, and rotarod, the influences of both injection and genotype were considered by two-way ANOVAs (genotype and injection) with multiple comparisons adjusted by Benjamini–Hochberg’s method. For open field statistical analyses, data were log-10-transformed to respect homoscedasticity assumptions. For immunohistochemistry, immunoprecipitation experiments, Aβ42-positive plaque number and morphology, and ELISA, we used the Mann–Whitney test. The number, area, and perimeter of plaques were analyzed by automatic-macro program on ImageJ software, and every point corresponds to an average of ten pictures for each mouse. The DPP4 activity in humans was performed using the two-way ANOVA test with Dunnett’s multiple comparisons test.

Data availability

All data are contained within the article and in the accompanying supporting information.

Supporting information—This article contains supporting information.

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