

The mechanism of γ -Secretase dysfunction in familial Alzheimer disease

Lucía Chávez-Gutiérrez^{1,2},
Leen Bammens^{1,2}, Iryna Benilova^{1,2},
Annelies Vandersteen^{3,4},
Manasi Benurwar^{1,2}, Marianne Borgers^{1,2},
Sam Lismont^{1,2}, Lujia Zhou^{1,2}, Simon Van
Cleynenbreugel^{1,2}, Hermann Esselmann⁵,
Jens Wiltfang⁵, Lutgarde Serneels^{1,2},
Eric Karran², Harrie Gijzen⁶,
Joost Schymkowitz^{4,7},
Frederic Rousseau^{4,7}, Kerensa Broersen^{3,8}
and Bart De Strooper^{1,2,*}

¹VIB Center for the Biology of Disease, Leuven, Belgium, ²Center for Human Genetics (CME) and Leuven Institute for Neurodegenerative Diseases (LIND), University of Leuven (KUL), Leuven, Belgium, ³Faculty of Science and Technology, MIRA Institute for Biomedical Technology and Technical Medicine, University of Twente, AE Enschede, The Netherlands, ⁴Switch Laboratory, Department of Cellular and Molecular Medicine, KULeuven, Leuven, Belgium, ⁵Department of Psychiatry and Psychotherapy, LVR-Clinics Essen, University of Duisburg-Essen, Essen, Germany, ⁶Janssen Research & Development, a Division of Janssen Pharmaceutica NV, Beerse, Belgium, ⁷VIB Switch Laboratory, Flanders Institute for Biotechnology (VIB), Leuven, Belgium and ⁸Vrije Universiteit Brussel, Brussel, Belgium

The mechanisms by which mutations in the presenilins (PSEN) or the amyloid precursor protein (APP) genes cause familial Alzheimer disease (FAD) are controversial. FAD mutations increase the release of amyloid β (A β)42 relative to A β 40 by an unknown, possibly gain-of-toxic-function, mechanism. However, many PSEN mutations paradoxically impair γ -secretase and ‘loss-of-function’ mechanisms have also been postulated. Here, we use kinetic studies to demonstrate that FAD mutations affect A β generation via three different mechanisms, resulting in qualitative changes in the A β profiles, which are not limited to A β 42. Loss of ϵ -cleavage function is not generally observed among FAD mutants. On the other hand, γ -secretase inhibitors used in the clinic appear to block the initial ϵ -cleavage step, but unexpectedly affect more selectively Notch than APP processing, while modulators act as activators of the carboxypeptidase-like (γ) activity. Overall, we provide a coherent explanation for the effect of different FAD mutations, demonstrating the importance of qualitative rather than quantitative changes in the A β products, and suggest fundamental improvements for current drug development efforts.

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*Corresponding author. VIB, Center for the Biology of Disease, Center for Human Genetics, University of Leuven (KUL), Herestraat 49, Leuven, Flanders B-3000, Belgium. Tel.: +32 16 346227; Fax: +32 16 347181; E-mail: Bart.destrooper@cme.vib-kuleuven.be

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Introduction

A central and still unresolved debate with important therapeutic implications in the field of Alzheimer disease (AD) research revolves around the question of how mutations in presenilin (PSEN), the catalytic core of the γ -secretases (De Strooper *et al*, 1998), cause disease. The γ -secretases are intramembrane cleaving protein complexes (Hebert *et al*, 2004; Shirovani *et al*, 2004) responsible for the generation of amyloid β (A β) from the amyloid precursor protein (APP). A β peptides of different lengths accumulate in amyloid plaques in the AD brain. More than 150 familial Alzheimer disease (FAD) mutations have been mapped to the genes encoding *PSEN1* or *PSEN2* (<http://www.molgen.ua.ac.be/ADMutations>), pointing to a crucial role of the γ -secretase complexes in the disease. Apart from PSEN, a mature and active γ -secretase complex consists of three additional subunits: Nicastrin (Nct), PSEN enhancer 2 (Pen-2), and either anterior pharynx 1 (APH-1) A or B (for a review, see Tolia and De Strooper, 2009). The γ -secretase complexes proteolyse type 1 transmembrane proteins, among them the APP, the Notch receptors and ligands, the Erb4 receptor and N-Cadherin (Wakabayashi and De Strooper, 2008).

As a rule, FAD PSEN mutations increase the relative amount of A β 42 versus A β 40 in *in vivo* and *in vitro* paradigms (Borchelt *et al*, 1996; Duff *et al*, 1996; Scheuner *et al*, 1996; Murayama *et al*, 1999), which led to propose that PSEN mutations act via a toxic gain-of-function mechanism. However, more refined analyses have made clear that the change in A β ratio does not necessarily reflect an increase in A β 42 production, but can also be the consequence of a decrease in A β 40 levels. Actually, many mutations reduce one or both products of the γ -secretase in steady-state conditions (Song *et al*, 1999; Bentahir *et al*, 2006; Shen and Kelleher, 2007; Shimojo *et al*, 2007; Heilig *et al*, 2010). These observations have led to an opposite hypothesis in which FAD mutations in PSEN cause dementia through a loss of function of γ -secretase, resulting in decreased proteolytic processing of different substrates and compromising intracellular signalling pathways (Shen and Kelleher, 2007; Kelleher and Shen, 2010). In fact, the current model for γ -secretase successive proteolysis (Takami *et al*, 2009) may link a loss of function to misprocessing of APP and abnormal generation of A β (De Strooper, 2007; Wolfe, 2007). However, the fact that less efficient proteolytic processing of APP may lead to alterations in the A β profile and AD is contraindicated in the light of the classical amyloid hypothesis, which stresses the importance of quantitative accumulation of either total

A β or A β 42 (Hardy and Selkoe, 2002). Moreover, a recent report has shown that reduced γ -secretase activity does not increase the production (accumulation) of longer A β peptides (Quintero-Monzon *et al*, 2011).

Importantly, the biophysical and biochemical properties of A β vary strongly with its length. Longer A β 42 has a much stronger tendency to aggregate than the shorter A β 40 (Jarrett and Lansbury, 1993; Jarrett *et al*, 1993). Furthermore, the relative ratio of A β 40 to A β 42 influences strongly the biological effects of the A β mixture *in vitro* and *in vivo*, even when total A β amounts are kept equal (Kuperstein *et al*, 2010). Whereas A β 40 appears to act protectively in various toxicity assays (Wang *et al*, 2006; Kim *et al*, 2007), longer A β peptides promote aggregation and neurotoxicity (McGowan *et al*, 2005). In fact, it has been suggested that the ratio (A β 42/A β 40) is more important than the absolute amounts of A β 42 (Tanzi and Bertram, 2005). Similar to A β 42, A β 43 is potently amyloidogenic and neurotoxic (Saito *et al*, 2011). While it is commonly found in AD brains (Welander *et al*, 2009), its potential relevance in disease was only recently addressed (Saito *et al*, 2011). Thus, qualitative changes in A β (De Strooper, 2007; Wolfe, 2007) are at least as important as the quantitative alterations proposed by the original amyloid hypothesis (Hardy and Selkoe, 2002).

In contrast, the 'simple' loss-of-function hypothesis proposes that A β alterations are only an epiphenomenon of the *PSEN* mutations, and that inefficient cleavage of membrane proteins by γ -secretase complexes is the fundamental upstream cause of the neurodegenerative process (Shen and Kelleher, 2007; Kelleher and Shen, 2010). This hypothesis finds support in (a) experimental results with *Psen* knockout mice (Saura *et al*, 2004), where progressive neurodegeneration occurs without A β deposition, and (b) in three case reports in which missense mutations in *PSEN* genes displayed neurodegenerative clinical phenotypes but no A β accumulation (discussed in Shen and Kelleher, 2007; Kelleher and Shen, 2010). However, this last argument has been considerably weakened by follow-up studies showing that neurodegeneration was likely caused by a second mutation in the progranulin gene in one case (Boeve *et al*, 2006), whereas in a second case abundant amyloid deposition in the frontal lobe appeared at autopsy (for further discussion, see Bergmans and De Strooper, 2010).

On the other hand, recent observations in patients suffering from familial acne inversa in China (Wang *et al*, 2010) and independently in Great Britain (Pink *et al*, 2011) raise doubts about the validity of the 'simple' γ -secretase loss-of-function hypothesis. This condition appears to be associated with the haploinsufficiency of γ -secretase subunit genes (*Nicastrin*, *Pen2*) and most likely involves a deficiency in Notch cell signalling. However, none of the acne-affected individuals had AD symptoms. These observations indicate that reduced γ -secretase activity is not sufficient to cause AD, although further follow-up studies in these families are needed. Alternative mechanisms for the loss-of-function hypothesis have been proposed over the years (for an overview, see De Strooper and Annaert, 2010). For instance, several reports indicate alterations in subcellular trafficking or turnover of selected membrane proteins (Wilson *et al*, 2004; Esselens *et al*, 2004) or defective acidification of phagolysosomal compartments associated with *PSEN* loss of function (Lee *et al*, 2010). In addition, disturbances in cellular Ca²⁺

homeostasis by direct effects on the Ca²⁺ leakage function of *PSEN* (Zhang *et al*, 2010) or indirect effects on Ca²⁺ signalling pathways (reviewed in Bezprozvanny and Mattson, 2008) have been associated to *PSEN* loss of function. However, these hypotheses do not provide an explanation for the mutations in *APP* and also do not take into account that all tested FAD mutations affect the prime function of *PSEN*, which is proteolysis.

From this brief overview it is clear that further in-depth investigation of the effects of clinical mutations on the function and structure of γ -secretase is required, especially given the relevance of such analysis for further drug development.

Addressing this important question implies multidisciplinary approaches, in which deep structural and functional studies dissect the mechanisms of FAD mutations. Solving the 3D-structure of the protease complex would allow studying how FAD mutations affect the structure, and possibly the function. However, this is a huge challenge as important technical and experimental barriers need to be overcome. On the other hand, dissecting γ -secretase activity by kinetic analysis can yield important mechanistic insights into how FAD mutants regulate enzyme function.

In vitro reconstitution of γ -secretase activity has provided initial insights into the enzymatic mechanism. Ihara and co-workers have provided compelling evidence for sequential processing of substrates by γ -secretase (Sato *et al*, 2003; Qitakahara *et al*, 2005; Kakuda *et al*, 2006; Yagishita *et al*, 2008). The most direct evidence was the identification of particular tri- and tetra-peptides generated from the *APP*-CTF stub by the γ -secretase (Takami *et al*, 2009). Their model proposes that *APP* can be sequentially cut along two production lines: A β 49 > A β 46 > A β 43 > A β 40 and A β 48 > A β 45 > A β 42 > A β 38 (Figure 1A). Accordingly, the endoproteolytic activity (first ϵ -cleavage) releases the *APP* intracellular domain (AICD) and A β 48 or A β 49. These long A β s are then shortened by consecutive carboxypeptidase-like γ -cleavages, which progressively decrease A β hydrophobicity and increase the probability of its release into the extracellular environment. In agreement, it has been shown that the endoproteolytic cleavage site determines the product line preference of the γ -secretase *in vivo* (Funamoto *et al*, 2004), and therefore the series of A β products. Also, presenilinase cleavage (the autocatalytic activation of *PSEN*) results in the generation of tripeptides in accordance with this model (Fukumori *et al*, 2010). The ϵ -cleavage in the *APP* substrate is analogous to the Notch S3 cleavage site (Sastre *et al*, 2001; Weidemann *et al*, 2002) and most likely other γ -secretase substrates are processed in similar ways.

In the current study, we used a cell-free assay to analyse how clinical mutations in *PSEN1*, *PSEN2* and *APP* affect the activity of the γ -secretase complex. Dissection of the different activities of the γ -secretase complex allowed us to reach a coherent explanation for the effects of the tested FAD mutations. We coupled kinetic studies of the endopeptidase activity to the analysis of the carboxypeptidase-like cleavage to show that FAD mutations have widely variable effects on the efficiency of the first cleavage, which releases the intracellular signalling domains of substrates. This observation rules out an impairment in the endopeptidase (ϵ) mechanism as necessary for the pathological effect of FAD mutations. In contrast, all FAD *PSEN* and *APP* mutations alter the proces-

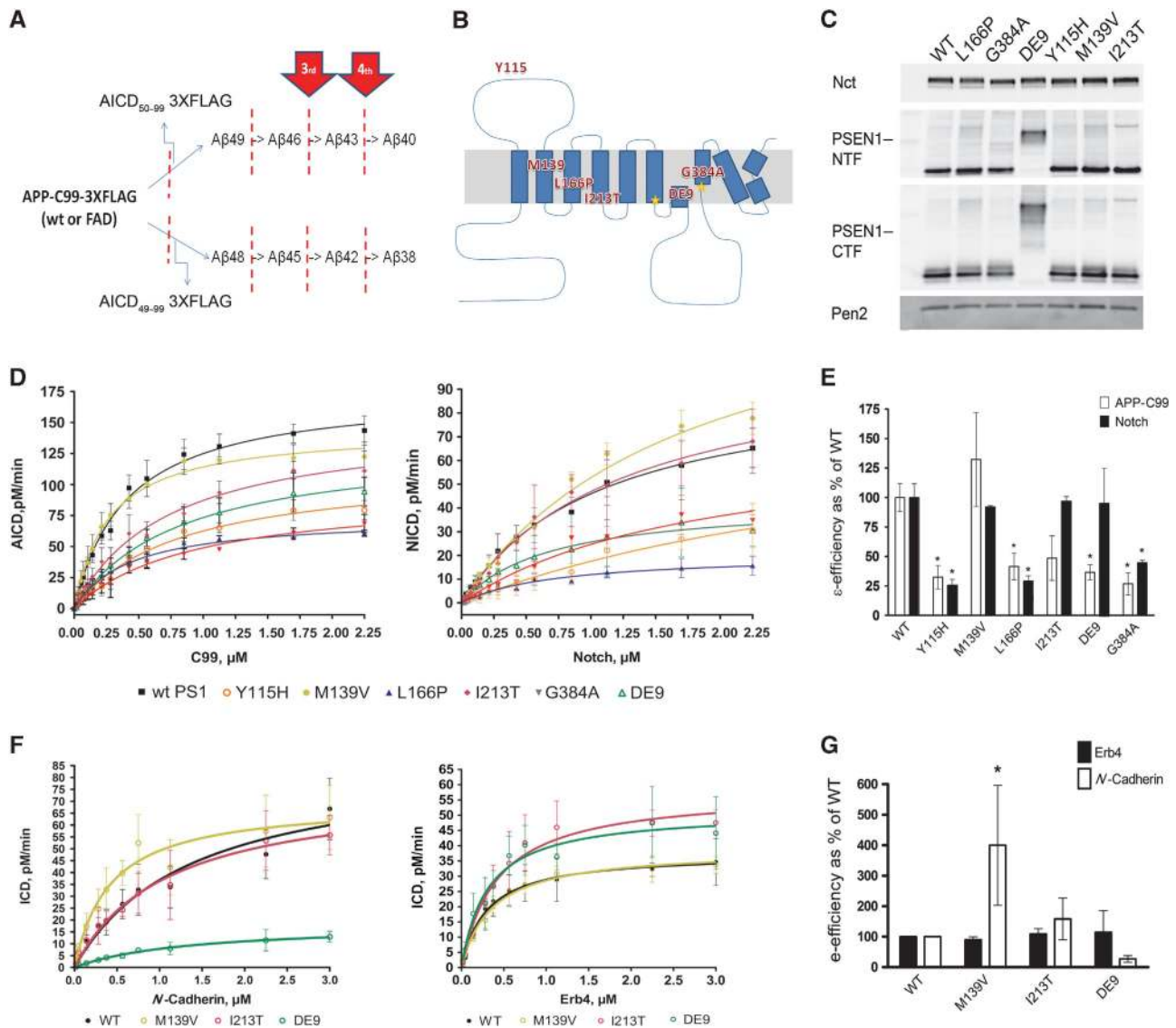


Figure 1 FAD-*PSEN1* mutations do not consistently decrease the enzymatic efficiency of the endopeptidase cleavage. (**A**, **B**) Schematic overviews of APP processing and location of FAD-*PSEN1* mutations used in the current study. (**C**) Expression levels of Nct, PSEN1-NTF, PSEN1-CTF and Pen-2 in *Psen1/2*^{-/-} mEFs transduced with human wt or FAD-*PSEN1* mutants using a replication-defective recombinant retroviral expression system (Clontech) and selected with puromycin (5 μg/ml). Western blotting and densitometric analysis of the CHAPSO-solubilized membrane proteins from the different *PSEN1* cell lines indicate that wt and mutant *PSEN1* rescued γ -secretase complex to similar extents. In order to determine specific activities for the wt or FAD complexes, γ -secretase activities were normalized to PSEN1 CTF fragment levels or full-length PS1 levels for the DE9 mutant. (**D**) Kinetic curves for wt and PS1-FAD mutants using purified APP-C99-3XFLAG or Notch-3XFLAG substrates (mean \pm s.e.) or (**F**) Erb4-3XFLAG and N-Cadherin-3XFLAG substrates (mean \pm s.d.). Detergent-extracted membranes were incubated in 0.25% CHAPSO reaction buffer with varying concentrations of purified substrate for 4 h at 37 °C. *In vitro* generated ICD-3XFLAG were analysed by quantitative western blot analysis (see Materials and methods). (**E**) FAD-*PSEN1* ϵ -enzymatic efficiencies for APP-C99 and Notch substrates (mean \pm s.e.). Enzymatic efficiencies unequivocally demonstrate that loss of function at the ϵ -cleavage is not a constant among *PSEN1* mutations. (**G**) FAD-*PSEN1* mutations that did not affect the generation of NICD did not change significantly the processing of Erb4 (mean \pm s.d.) either. In contrast, N-Cadherin processing was significantly upregulated by the M139V (mean \pm s.d.). (**E**, **G**) Experiments were repeated 3–5 times. Statistical significance of the data was tested with one-way analysis of variance (ANOVA) and Dunnett's post test, taking the corresponding WT efficiency as control group, **P* < 0.05.

sing of APP, regulating the generation of A β by three different mechanisms.

Results

FAD-PS1 mutations do not consistently impair the endopeptidase activity of the γ -secretase

We analysed the effects of FAD mutations PSEN1-Y115H, -M139V, -L166P, -I213T, -G384A and delta-exon9 (DE9) on the kinetic constants of the ϵ -cleavage of APP, Notch, Erb4 and N-

Cadherin substrates. The selected mutations are spread throughout the *PSEN1* primary sequence (Figure 1B). Importantly, blockage by the transition state analogue L-685,458 (TSA, InhX) demonstrated the specificity of the assays (Supplementary Figure 1A). To determine the kinetic constants of wt and FAD γ -secretase complexes, we used CHAPSO-extracted membranes from *Psen1/2*^{-/-}, rescued with wt or FAD-mutant *PSEN1* as source of enzyme (Figure 1C) and purified APP C99-3XFlag, Notch-3XFlag, Erb4-3XFlag or N-Cadherin-3XFlag as substrates.

Table 1 Kinetic parameters for human PSEN1 γ -secretase complexes using APP-C99, Notch, Erb4 or N-Cadherin as substrates

	APP-C99 substrate		Notch substrate		Erb4 substrate		N-Cadherin substrate	
	Km \pm s.e., μ M	V _{max} \pm s.e., pM/min	Km \pm s.e., μ M	V _{max} \pm s.e., pM/min	Km \pm s.d., μ M	V _{max} \pm s.d., pM/min	Km \pm s.d., μ M	V _{max} \pm s.d., pM/min
PS1 wt	0.40 \pm 0.05	175.6 \pm 8.4	1.08 \pm 0.17	95.7 \pm 7.5	0.31 \pm 0.07	37.72 \pm 6.18	1.46 \pm 0.36	88.37 \pm 10.95
Y115H	0.81 \pm 0.18	113.3 \pm 11.3 ^a	3.92 \pm 1.97	86.49 \pm 31.4	—	—	—	—
M139V	0.27 \pm 0.04 ^a	144.5 \pm 6.8 ^a	1.78 \pm 0.21 ^a	146.9 \pm 10.1 ^a	0.40 \pm 0.23	39.76 \pm 6.36	0.42 \pm 0.19 ^a	71.16 \pm 20.14
L166P	0.43 \pm 0.07	74.03 \pm 4.2 ^a	0.97 \pm 0.2	23.76 \pm 2.4 ^a	—	—	—	—
I213T	0.73 \pm 0.18	151.1 \pm 14.7	1.26 \pm 0.26	106.1 \pm 11.2	0.45 \pm 0.13	58.68 \pm 5.83 ^a	1.02 \pm 0.11	74.95 \pm 11.76
DeltaE9	0.82 \pm 0.18 ^a	133.5 \pm 13.5 ^a	0.67 \pm 0.24	42.8 \pm 6.5 ^a	0.33 \pm 0.06	40.84 \pm 8.12	1.70 \pm 0.43	21.97 \pm 8.39 ^a
G384A	0.92 \pm 0.18	93.87 \pm 8.7 ^a	1.85 \pm 0.42	71.04 \pm 9.5	—	—	—	—

Kinetic values are derived from the curves displayed in Figure 1 and were determined by nonlinear curve-fitting using GraphPad Prism 4 software (see Material and methods section).

^aSignificant changes according to the 95% CI ($P < 0.05$). *In vitro* activity assays were performed using CHAPSO-extracted membranes from Psen1/2^{-/-} MEFs stably transduced with human wt or FAD PSEN1 mutants and purified substrates-3XFLAG, $n \geq 3$.

The kinetic data fit the Michaelis–Menten reaction curves (Figure 1D and F), and Km (affinity constant) as well as V_{max} (maximal velocity) were determined (Table I). Since γ -secretase activities are normalized to enzyme levels, V_{max} can be taken as k_{cat} and enzymatic efficiencies calculated as k_{cat}/Km . The results reveal diverse effects of the FAD–PSEN1 mutations on this important kinetic parameter. Y115H, L166P and G384A mutants decrease γ -secretase efficiencies by 75% for both APP and Notch, while I213T and DE9 only affect APP, and M139V does not show any effect on the ϵ -cleavage (Figure 1E). Moreover, FAD–PSEN1 mutations that do not affect Notch endoproteolysis do not impair Erb4 cleavage either, while only the M139V significantly increases the processing of N-Cadherin (Figure 1G). Thus, the tested FAD–PSEN1 mutations have no consistent inhibitory effect on the endoproteolytic cleavage of γ -secretase substrates, indicating that reduced release of intracellular domains and signalling cannot explain their AD-causing effects.

FAD–PSEN mutations impair the fourth γ -secretase cleavage in both product lines

Next, we asked whether FAD–PSEN1 mutations lead to APP misprocessing at the γ -cleavage sites. We use Ihara's model (see Introduction) for further description of our work since it explains very well our observations. Kinetic analysis of the carboxypeptidase-like activity is challenging to perform since controlling substrate concentrations, that is, the intermediary A β products, is experimentally not possible yet. Nevertheless, we measured two γ -products in each production line: A β 43, A β 42, A β 40 and A β 38 (Figure 1A) at saturating APP-substrate concentration. Thus, substrates (A β 43 and A β 42) and products (A β 40 and A β 38) of the fourth γ -secretase cleavage in both pathways are analysed and provide a relative number for the γ -cleavage efficiency. Importantly, as some of the clinical mutants affect the ϵ -cleavage, we normalized the A β product levels (A β 38, A β 40, A β 42 or A β 43) towards total AICD (Figure 2A and B). AICD reflects the total initial A β substrate (A β 49 + A β 48) produced and processed in each reaction. Low A β 40 and A β 38 levels and high, long A β levels (>A β 42) are found in all the FAD-linked mutations tested, including the M139V, which does not affect the ϵ -efficiency. Interestingly, the M139V mutation affects the processing of APP only at the level of A β , indicating that endo- and

carboxypeptidase-like activities of the γ -secretase can be dissociated.

Total 'secreted' A β , defined as the sum of A β 38, A β 40, A β 42 and A β 43, decreases significantly in the Y115H, L166P, DE9 and G384A mutations (Figure 2B), implying the concomitant accumulation of longer A β precursors generated in cycles 2 and 3. Qualitative analysis of the A β profiles in urea-based SDS–PAGE confirmed this observation (Figure 2C and Supplementary Figure 2B). We finally determine product/substrate ratios for the fourth enzymatic turnover (A β 38/A β 42 and A β 40/A β 43) (Figure 2D), which demonstrates that the FAD–PSEN1 mutations investigated here dramatically impair the fourth γ -secretase cleavage in both product lines. Our data imply that FAD–PSEN1 mutations cause AD by qualitative shifts in A β profiles and not by general loss of function of the enzyme complex (Shen and Kelleher, 2007; Wolfe, 2007). The dysfunction at the carboxypeptidase-like activity of the complex not only explains the widely documented increase of the A β 42/A β 40 ratio, but also suggests a pathological relevance of an increase in A β 43, which has been reported recently *in vivo* with the PSEN–R278I (Saito *et al*, 2011).

In order to confirm our *in vitro* data, mouse embryonic fibroblasts (MEFs) derived from Psen1/2^{-/-} mice (Herreman *et al*, 2000) rescued with human WT- or FAD–PSEN1 mutants were transiently transduced with APPsw. Secreted A β levels (sA β) were quantified by enzyme-linked immunosorbent assay (ELISA). Figure 2E shows drastic reductions in the A β 38/A β 42 and A β 40/A β 43 ratios for all FAD–PSEN1 mutations, confirming that the fourth enzymatic turnover of the γ -secretase is actually impaired in cells (native γ -secretase conditions). To investigate whether FAD–PSEN2 mutations also affect the fourth enzymatic turnover of the γ -secretase, we performed kinetic analyses with human WT–PSEN2 or FAD N141I–PSEN2 mutant. The effect of the mutation on the endopeptidase efficiency of the γ -secretase complex does not reach statistical significance (Figure 3A) (mean \pm s.e.: 100 \pm 39.9, $n = 4$ or 46.6 \pm 3.9, $n = 3$ for WT- or FAD N141I–PSEN2, respectively, two-tailed *t*-test, $P = 0.3$). Although we cannot discard that this difference is biologically meaningful, the effect on the fourth catalytic cycle is unequivocal. In particular, the A β 40 product was decreased to undetectable levels (Figure 3B). Similar to the mutations in PSEN1, the N141I–PSEN2 reduces the A β 38/

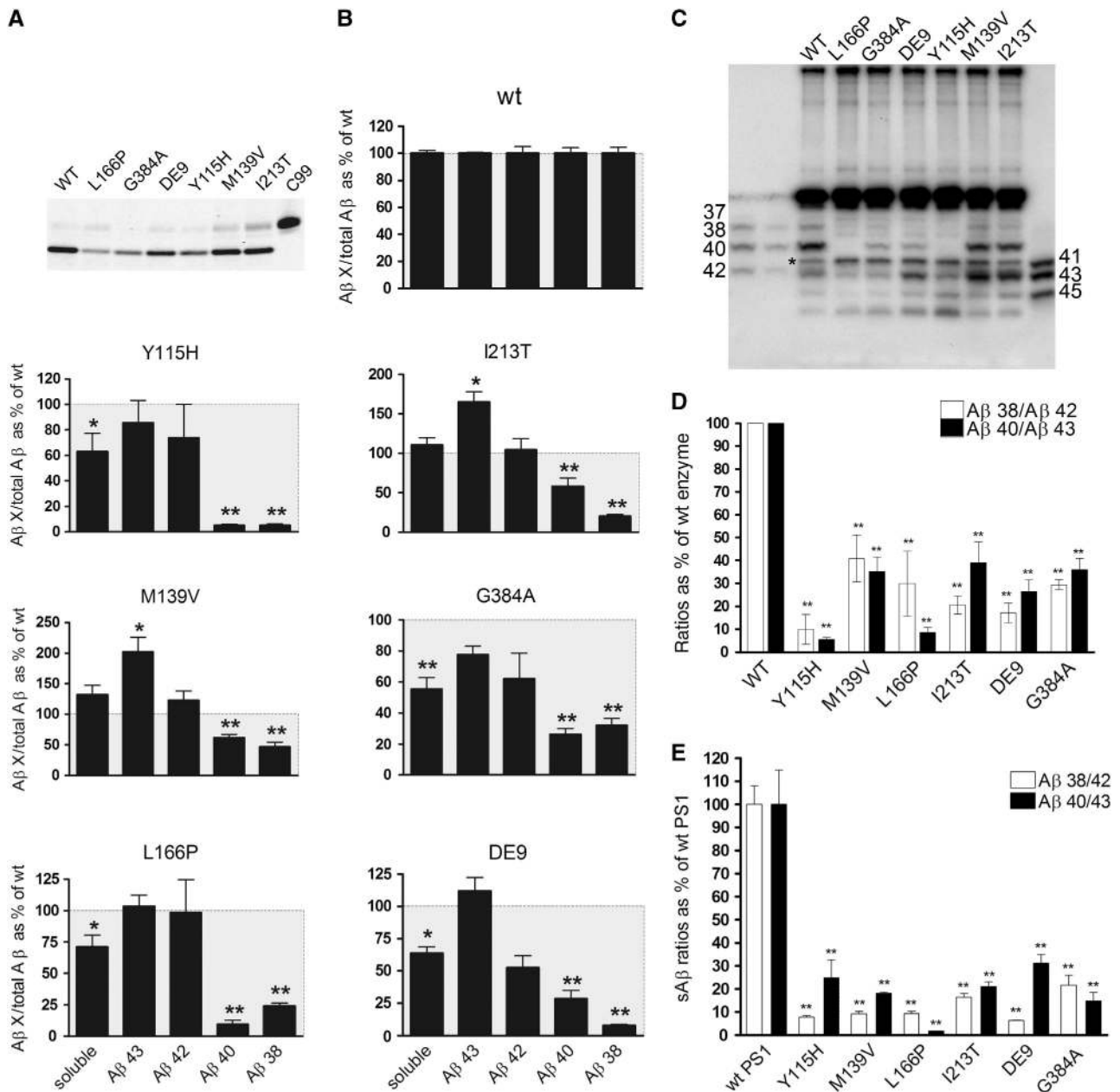


Figure 2 FAD-*PSEN1* mutations impair the fourth enzymatic turnover. AICD levels (moles per min) generated by the wt or FAD mutant complexes (A) were used to normalize Aβ products (moles per min) in order to determine accurately Aβ generation relative to C99 substrate. Aβ profiles (B) thus represent Aβ products corrected for the initial endoprotease activities, plotted as percentage of the wt Aβ products (mean ± s.e.). Soluble Aβ (sum of Aβ38, Aβ40, Aβ42 and Aβ43 peptides) gives information about the efficiency of the γ-cleavages: lower levels (< 100%, grey box) suggest that longer peptides (> Aβ43) accumulate in the reactions. (C) In agreement with the ELISA quantifications, total Aβ analysed in urea-based gels show increments in Aβ42 and Aβ43, and reductions in Aβ40 and Aβ38 in FAD-*PSEN1* mutations, relative to wt. (*) Indicates a non product band that is present in the C99 substrate (see Supplementary Figure 2). (D) Aβ product/substrate ratios determined *in vitro* for the FAD-*PSEN1* mutations show an impairment at the fourth γ-secretase turnover (mean ± s.e.). Experiments in (B) and (D) were repeated 4–6 times. Statistical significance of the data tested with one-way ANOVA and Dunnett’s post test taking the corresponding WT as control group; **P*<0.05, ***P*<0.01. (E) Aβ product/substrate ratios determined *in vivo* confirm impairment at the fourth enzymatic cycle: wt or FAD-*PSEN1* mEF cell lines were transiently transduced with APP^{swe}, extracellular media collected at 24 h after infection and sAβ measured by ELISA (mean ± s.e.). Statistical significance: *n*=4, ANOVA and Dunnett’s post test, ***P*<0.01.

Aβ42 and Aβ40/Aβ43 ratios (Figure 3C), confirming an impairment in the carboxypeptidase-like activity.

Since high Aβ43 and Aβ42 (substrates in this cycle) accumulate *in vitro* or are released *in vivo*, we speculate that FAD-*PSEN* mutations promote a premature release of the Aβ43/Aβ42 peptides.

FAD-APP mutations change the product line preference of the γ-secretase

We then asked whether similar mechanisms could explain the effect of mutations in the APP substrate. The tested mutations are located close to the γ-secretase cleavage site, that is, T43I, V44A, I45T, V46F and V46I (Figure 4A) and all

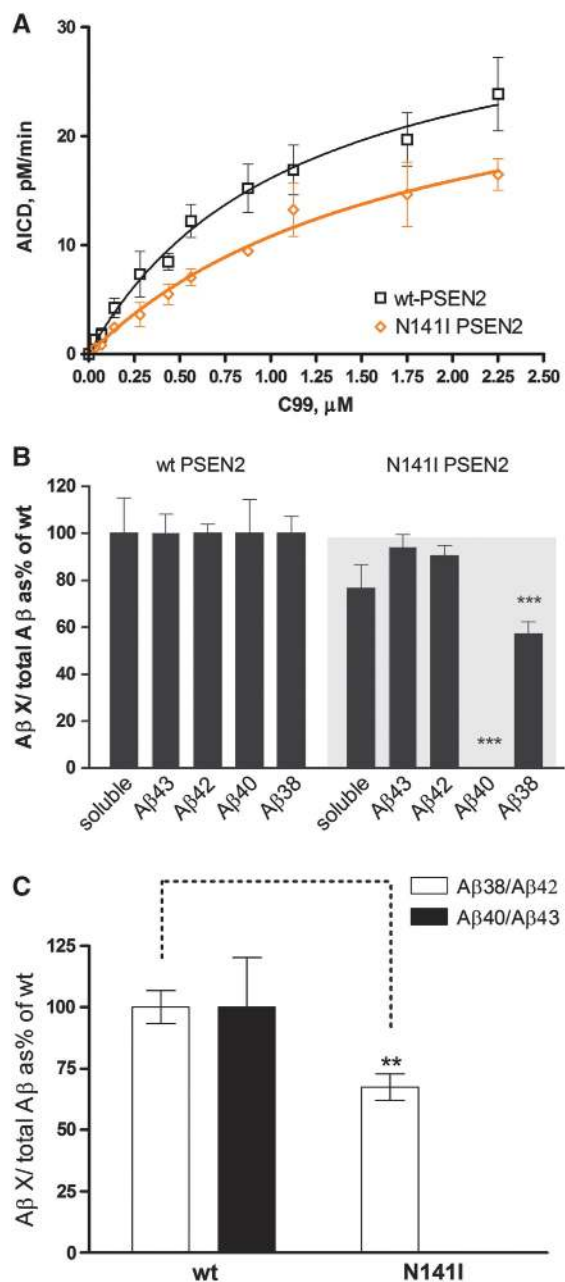


Figure 3 FAD-*PSEN2* N141I impair the fourth enzymatic turnover. (A) Kinetic curves for wt and *PSEN2*-FAD N141I mutant using purified APP-C99-3XFLAG as substrate (mean \pm s.e.). (B) A β profiles represent A β products corrected for the initial endoprotease activities, plotted as % of the wt A β products (mean \pm s.e.). Soluble A β (sum of A β 38, A β 40, A β 42 and A β 43 peptides) suggests accumulation of longer peptides (>A β 43) in the mutant reactions. (C) A β product/substrate ratios determined *in vitro* for the FAD-*PSEN2* mutation show an impairment at the fourth γ -secretase turnover (mean \pm s.e.). In (B) and (C) statistical significance (two-tailed *t*-test) is indicated by ** P <0.005 and *** P <0.001. Note that N141I abolishes A β 40 generation.

produce wild-type A β 38, A β 40, A β 42 and A β 43 peptides, except for the T43I mutation. Kinetic analyses of the ϵ -cleavage show that APP-T43I, V44A and -I45T mutants produce less AICD per mol mutant APP compared to wt substrate, while the other mutations do not affect the ϵ -enzymatic efficiency (Figure 4B and C). In order to analyse

accurately the A β profiles from wt and FAD substrates, A β levels were normalized to the amount of AICD generated in the reaction. FAD A β levels, corrected for the initial amounts of substrates, were then plotted as a percentage of the wt enzyme (Figure 4D). Importantly, and in contrast to FAD-*PSEN*, all the tested *APP* mutations do not affect A β 38/A β 42 or A β 40/A β 43 ratios (Figure 4E). The I45T mutant is the exception, showing increased A β 38/A β 42 ratio, which would be consistent with an impairment in the processing of A β 45 (mutant peptide) to A β 42. However, FAD-*APP* mutations result in high A β 40/A β 38 compared to wt *APP* (Figure 4F). Thus, all investigated mutations change the product line preference by shifting *APP* processing towards the A β 38 production line. The APP-V44A and -I45T substrates in particular show an additional accumulation of longer A β precursors (generated in cycles 2 and 3), as deduced from soluble A β in Figure 4D. The change in the product line can be explained if these *APP* mutations shift the position of the ϵ -cleavage to generate more A β 48, the initial substrate in the A β 38 production line. A neo-epitope antibody against AICD₅₀₋₉₉ (characterized in Supplementary Figure 3) was generated, and allowed us to confirm the product line preference (Figure 5A and B).

Figure 5C shows that the FAD-*APP* mutations consistently shift the position of the ϵ -cleavage towards AICD₄₉₋₉₉, promoting the A β 38 product line, and therefore causing increments in the A β 42 and A β 38 products. Importantly, HEK cells transiently transfected with FAD-mutant C99 substrates increase the A β 42 and A β 38 levels in the extracellular medium while decreasing A β 40, compared to control (wt C99) (Supplementary Figure 4). Figure 4G actually shows that FAD-*APP* mutations change the product line preference (A β 40/A β 38 ratio), but do not alter the fourth enzymatic turnover (A β 38/A β 42 ratio). Similar results were obtained from primary neuronal cultures transiently expressing wt-, -I45T- or V46F-APP (Figure 4H). These results indicate that our observations in the cell-free assay can be extrapolated to the *in vivo* situation. The FAD-*APP* data imply that promoting the A β 38 production line is pathogenic.

It has been shown that small changes in the composition of A β mixes affect critically their aggregation kinetics and toxic effects (Kuperstein *et al*, 2010), for example, a minor increase in the A β 42:A β 40 ratio stabilizes toxic oligomeric species. Since *APP*-FAD mutations increase both A β 42 and A β 38, we asked whether A β 38 could have similar biophysical attributes as A β 40, and therefore could alleviate the potential toxic effects associated to A β 42. However, and in contrast to A β 40, A β 38 has a predicted higher tendency to aggregate (<http://www.tango.crg.es>) (Fernandez-Escamilla *et al*, 2004). To validate this prediction, A β -aggregation assays were performed using thioflavine T fluorescence as readout. We compared the behaviours of A β 40 and A β 38 peptides by analysing the aggregation properties of the A β 42:A β 40 (1:9) and A β 42:A β 38 (1:9) mixes. In contrast to A β 40, A β 38 drives aggregation of A β mixes to higher extents (Supplementary Figure 5A). We then compared the effects of different A β peptides on spontaneous synaptic transmission in the primary mouse hippocampal neurons. Our results show that A β 38, similar to A β 42 and A β 43, but to a lesser extent, elicits acute synaptotoxicity (Supplementary Figure 5B). Although further work is needed to investigate the effects of A β 38

in vivo, these data confirm that individual A β peptides have widely divergent biophysical and biochemical properties.

Effects of inhibitors and modulators on the γ -secretase activity

Our data indicate that various mechanisms affecting the A β spectrum generated by γ -secretase are responsible for the pathogenic effects of the FAD mutations. Therefore, we asked to what extent γ -secretase inhibitors (GSI) and modulators (GSM) that were tested in clinical trial or are under development (De Strooper *et al*, 2010) affected the different parameters discussed above. To evaluate γ -secretase inhibition under equal kinetic conditions, we took advantage of the *in vitro* system and performed activity assays at $1 \times K_m$ substrate concentrations for APP C99 or Notch substrates (0.4 and 1.1 μM , respectively). Under these conditions, the GSI semagacestat (LY-450139), begacestat (Notch sparing GSI) and avagacestat (Notch sparing GSI) efficiently inhibit A β generation in the two production lines (Supplementary Figure 6). However, semagacestat, which failed in phase III trial (<https://www.investor.lilly.com/releasedetail2.cfm?releaseid=592438>) because of cognition and skin problems, is more selective for Notch than for APP (AICD IC_{50} = 257.8 nM and Notch intracellular domain (NICD) IC_{50} = 24.62 nM (95% confidence interval (CI): 190.2–349.5 nM for APP and 15.74–38.51 nM for Notch, $n = 5$), whereas the transition state inhibitor L-685,458 affects both substrates to an equal extent (Figures 6A and B). Surprisingly, and in contrast to previous claims (Martone *et al*, 2009; Gillman *et al*, 2011), the selectivity of both of the ‘Notch sparing’ GSI is not significantly different for APP and Notch substrates (Begacestat: AICD IC_{50} = 61.71 nM and NICD IC_{50} = 138.4 nM (95% CI: 24.77–153.7 nM for AICD and 73.29–261.3 for NICD, $n = 3$) and BMS708163: AICD IC_{50} = 6.82 nM and NICD IC_{50} = 20.03 nM (95% CI: 4.06–11.46 nM for AICD and 7.76–51.7 nM for NICD, $n = 3$)) (Figure 6C and D).

Recently, γ -secretase modulators (GSM) have been evaluated as an alternative to GSI (Weggen *et al*, 2001). GSM lower A β 42 and increase A β 38, but the precise mechanism of action has not been elucidated. One NSAID and two arylimidazole-derived GSM (Oehlrich *et al*, 2010) (E-2012 and a close analogue) did not affect the endopeptidase activity (Figure 6E and Supplementary Figure 7A) but, as expected, reduced A β 42 and increased A β 38. Analysis of the product/substrate ratios for the fourth enzymatic turnover shows that these drugs increase specifically this cycle (Figure 6F and Supplementary Figure 7B) and act, therefore, in the opposite way to the clinical FAD–PSEN mutations. The A β ratios indicated that the GSM evaluated in this study act mainly on the fourth cycle of the A β 38 production line. In fact, our data show them as activators of the γ -secretase (GSA). Taking into account the changes in the FAD–APP A β profiles and the possibility that A β 38 may be part of the pathogenic mechanism, it is crucial to evaluate to what extent the differential effects of the GSA on the A β production lines are problematic.

Discussion

This study settles several issues that have been heavily debated in the literature. Dissection of the different activities of the γ -secretase complex allowed us to characterize the

mechanisms that are regulated in a consistent fashion by FAD mutations in PSEN and APP. Previous reports have employed steady-state analyses to evaluate the effect of these mutations on the γ -secretase. In general, these studies have employed transfected cells to measure, in culture or *in vitro*, changes in secreted product levels or follow the intracellular generation of ICD products by coupling it to reporter systems. Although these approaches are informative, they do not truly reflect the kinetic features of the mutated γ -secretase complexes or APP substrates. For instance, substrate concentration and accessibility is not controlled in such assays.

By analysing the catalytic efficiency of the γ -secretase complex (wt or mutated) in an *in vitro* assay, with both enzyme and substrate in solution, we find here that mutations in PSEN1 and PSEN2 affect γ -secretase at three levels. We see (1) a variable inhibitory effect at the initial endoproteolytic ϵ -cleavage step, which releases the intracellular domains of substrates such as APP, Notch, Erb4 and N-Cadherin. (2) A consistent effect on the consecutive carboxypeptidase-like γ -cleavage with all PSEN mutations causing a ‘premature’ release of (intermediary) substrates/products, explaining why longer A β is generated by these mutants. Interestingly, our data suggest that both A β 42/A β 38 and A β 43/A β 40 ratio increments are pathologically relevant. (3) Additionally, some of the mutations in PSEN and all mutations investigated here in the APP sequence (selected for their close position to the γ -cleavage site in APP) affect the initial position of the ϵ -site, that is, whether γ -secretase cleaves preferentially at position 49–50 or 51–50 in the APP sequence. While these three mechanisms explain for the first time the abundantly documented increase in A β 42/A β 40 ratio associated with all FAD mutations, they also provide a set of entirely novel insights, as we discuss in the following paragraphs.

Our study gives definite numbers on the catalytic efficiency of the γ -secretase complex at the ϵ -site and unequivocally shows that ‘loss of function’—lower catalytic efficiency—is not consistent across the FAD mutations tested. In this regard, we wish to draw attention to the fact that point mutations in PSEN might affect protein stability, and therefore solubilization of ‘less stable’ FAD γ -secretase complexes might result in an accentuated ‘loss of function’. Thus, we cannot exclude that the enzymatic efficiencies of particular FAD complexes (less stable) are underestimated in the conditions used in the current work, which would even strengthen our conclusion that ‘loss of function’ is not necessary for the FAD pathogenic mechanism. Taking all into consideration, our results indeed rule out the possibility that loss of intracellular signalling is necessary and sufficient to cause AD, as postulated by the ‘simple’ loss-of-function hypothesis. Interestingly, the effects at the ϵ -cleavage site are also variable for different substrates tested (K_m values for APP, Notch, Erb4 and N-Cadherin, Table I), suggesting that some of the clinical mutations affect the substrate specificity mechanism. This is in particular clear for the M139V and DE9 mutations. DE9 removes part of the hydrophobic domain VII (HDVII) of PSEN1, which is located in the active site of the γ -secretase (Tolia *et al*, 2008), while the M139 residue is located in the second transmembrane domain of PSEN, which contributes to the formation of the initial substrate-binding site (Watanabe *et al*, 2010). On the contrary, our results show that the L166P mutation affects the catalytic rate of the enzyme but does not change the substrate specificity of the complex,

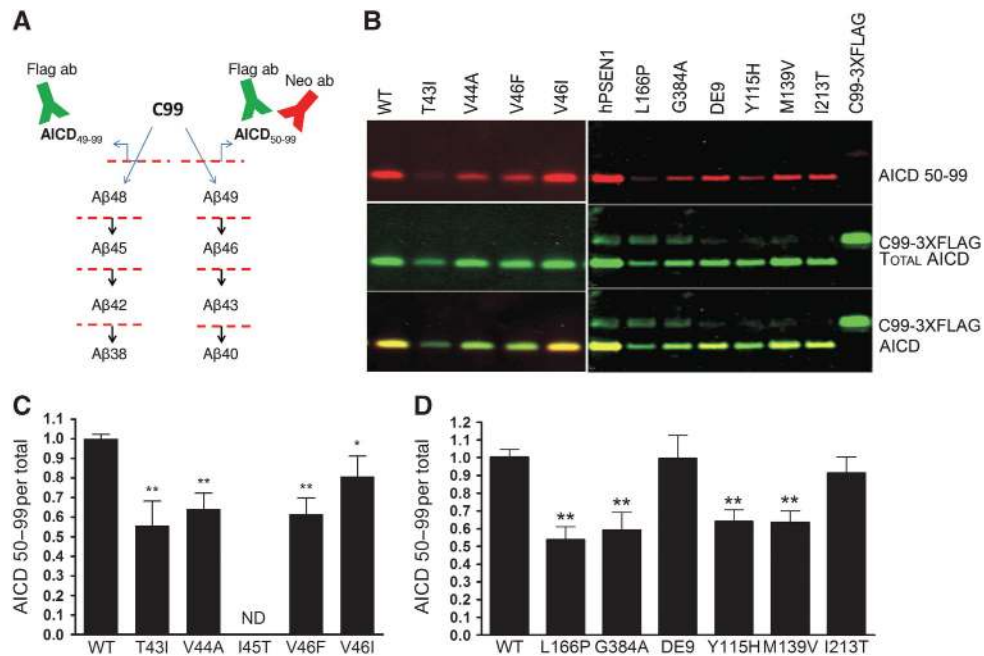


Figure 5 Shift in the ϵ -cleavage position contributes to the FAD-associated phenotype. **(A)** Detection of AICD₅₀₋₉₉ and total AICD using a neo-epitope antibody and the FLAG-M2 antibody, respectively. **(B)** SDS-PAGE/western blot analysis of AICD products from either wt and FAD substrates (left panel) or wt and FAD-*PSEN1* mutants (right panel). Signals for the AICD₅₀₋₉₉ neo-epitope antibody or the FLAG-M2 antibody are shown in red and green, respectively. Overlapping neo-epitope and FLAG antibody signals are displayed in yellow. **(C)** AICD₅₀₋₉₉/total AICD ratios indicate that FAD-APP mutations promote the A β 38 product line by shifting the ϵ -cleavage position. The I45T could not be included in the analysis because of extremely low AICD signals (ND, not determined). **(D)** This pathogenic mechanism is also observed in some FAD-*PSEN1* mutations. Statistical significance of the data ($n=5$) tested with ANOVA and Dunnett's post test, taking AICD generated in WT conditions as control group; * $P<0.05$, ** $P<0.01$.

ratios. Our data therefore give a mechanistic explanation for the decrease in short A β s (<40) reported in the cerebrospinal fluid of carriers of *PSEN1*-A431E10 (Portelius *et al*, 2010) or the alterations in the lengths of A β peptides produced *in vitro* by FAD-*PSEN1*-containing complexes (Murphy *et al*, 2002). Moreover, biophysical observations have shown that FAD-*PSEN1* mutations alter the conformation of the γ -secretase complex (Berezovska *et al*, 2005). Based on our biochemical data we propose that changes in the active site of FAD-*PSEN1* mutations promote the premature release of the A β 43 or the A β 42 peptides from the γ -secretase complexes.

A third mechanism by which FAD-APP mutations act in particular is the shift in the initial ϵ -cleavage site resulting in an increased A β 48 > A β 38. Likely, the product preference results from differential docking modes of the APP substrate into the active site. We confirmed the shift towards A β 48 > A β 38 in living cells expressing wt or mutant APP

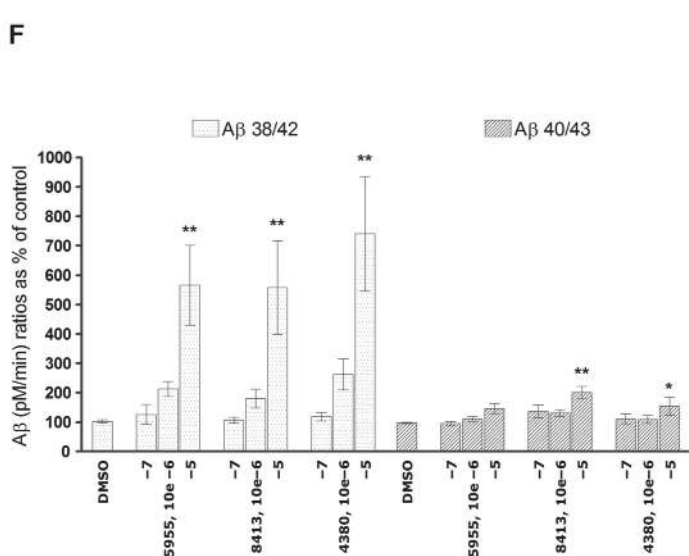
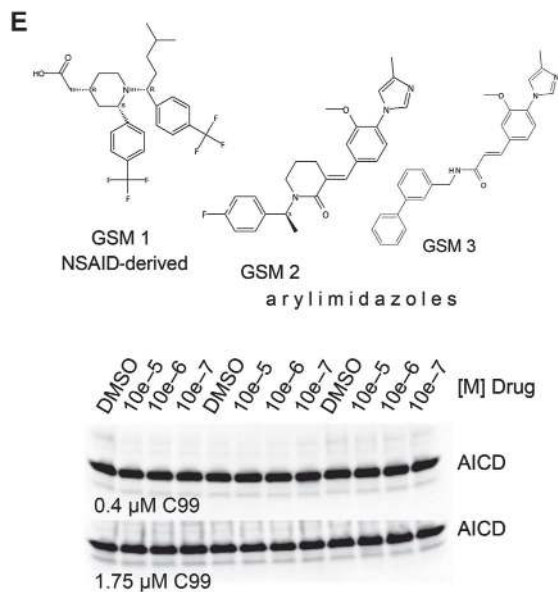
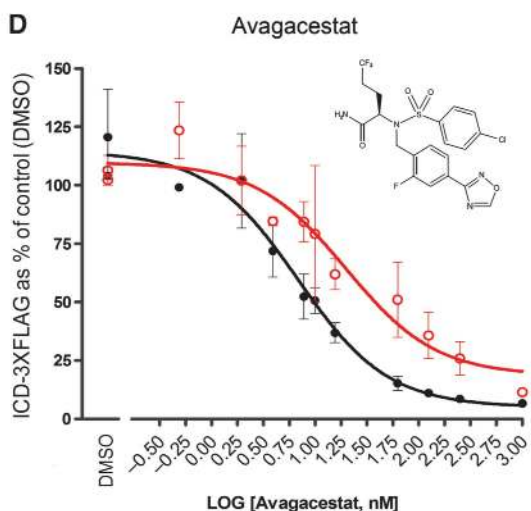
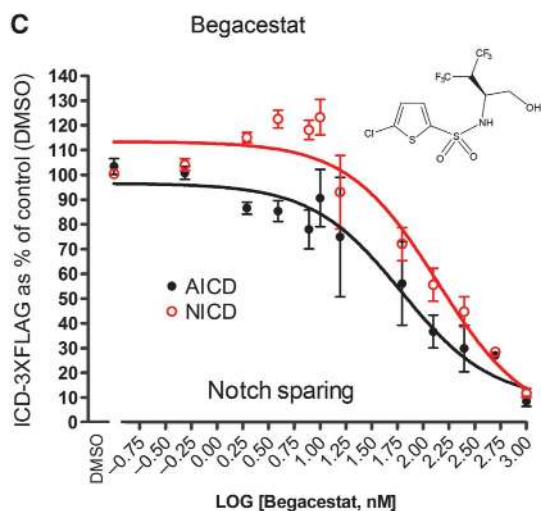
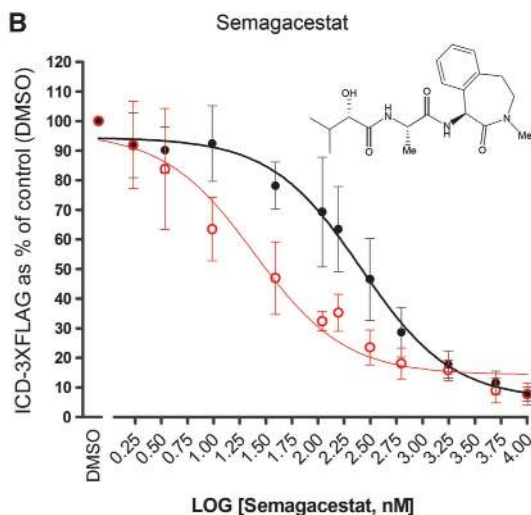
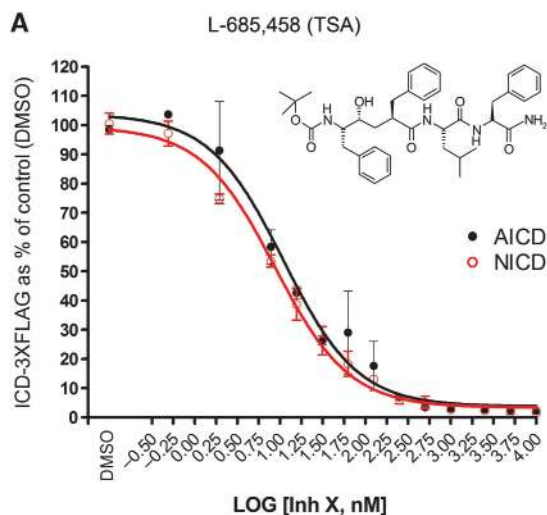
substrates. This result corroborates our claim that the product line preference is an intrinsic property of the γ -secretase complex that remains unaltered in our cell-free assay. Interestingly, this shift in initial docking and production lines is also observed in four of the six *PSEN1* mutants (Figure 5C and D). The fact that some FAD-*PSEN1* mutations combine these two mechanisms (impaired fourth cycle and change in the product line preference) explains the direct and indirect correlations between A β 38 and A β 42 levels reported in the past (Cziri *et al*, 2008; Page *et al*, 2008).

Our study thus demonstrates that FAD mutations cause qualitative changes in the A β profiles by various mechanisms (Bentahir *et al*, 2006; De Strooper, 2007), and that decreased release of intracellular domains (Kelleher and Shen, 2010) is not an essential part of the AD pathogenic mechanism. Nevertheless, as indicated above, the most aggressive *PSEN1* mutations, for example, the L166P, negatively impact the

Figure 4 FAD substrate mutations shift APP processing towards the A β 38 product line. **(A)** Schematic overview of FAD-APP mutations used in this study. **(B)** Kinetic curves for the ϵ -processing of APP. Detergent-extracted membranes from *Psen1/2^{-/-}* mEFs rescued with human wt *PSEN1* were incubated in 0.25% CHAPS buffer, with varying concentrations of purified wt or FAD-APP substrates. AICD-3XFLAG levels were analysed by quantitative western blot analysis (see Materials and methods). **(C)** Enzymatic efficiencies for FAD-APP-C99 substrates (mean \pm s.e.) prove that AICD generation is affected in three out of five FAD-mutant substrates. **(D)** FAD A β product profiles suggest consistent increments in A β 42 and A β 38. Soluble A β levels (sum of A β 38, A β 40, A β 42 and A β 43 peptides) suggest accumulations of longer A β peptides in the γ -processing of the V44A and I45T mutants. The T43I mutation disrupts the epitope for the anti-A β 43-specific antibody, thus neither A β 43 nor soluble A β levels could be determined (ND). **(E)** A β product/substrate ratios reveal that APP mutations do not consistently affect the fourth γ -secretase turnover, but change the product-line preference as indicated by the A β 40/A β 38 ratio **(F)**. **(G)** sA β in the conditioned media of HEK293 cells transiently transfected with wt or FAD-C99 mutants were quantified by ELISA (see Supplementary Figure 4). sA β ratios indicate that APP-FAD substrate mutants change the product-line preference towards the A β 48 > A β 38 (A β 40/A β 38) in living cells, but do not affect the fourth catalytic turnover of the γ -secretase (A β 38/A β 42) (mean \pm s.e., $n=5$). **(H)** Primary cultured neurons were transduced with SFV expressing WT APP or the indicated mutant substrates (mean \pm s.d., $n=3$). **(C-H)** Statistical significance tested with one-way ANOVA and Dunnett's post-test, taking the corresponding WT as control group; * $P<0.05$, ** $P<0.01$.

endopeptidase activity as well, and therefore it is not unlikely that 'partial loss' of γ -secretase function at Notch or other γ -secretase substrates acts as an aggravating factor in FAD.

Moreover, our A β product profiles evidence the generation of longer A β peptides (>A β 43) by the most aggressive FAD complexes. However, whether particular changes in the FAD



A β profiles can be correlated to the age of onset is an interesting but unaddressed question.

In a final series of experiments, we have also assessed to what extent different γ -secretase-directed drugs such as GSI and GSM affect the three mechanisms identified in the current work. When investigating the transition state analogue L-685,458 (InhX), semagacestat, and the Notch-sparing begacestat and avagacestat, we found that the four compounds lowered all γ -cleavages to a similar extent and did not change the A β ratios. However, when we assayed the effects of these GSI on Notch processing, which is considered to be the major liability of GSI, we surprisingly found that semagacestat was more effective in inhibiting Notch than APP. This is particularly significant when considering that a phase III clinical trial with semagacestat was interrupted last year because of severe side effects including worsened cognition and increased incidence of skin cancer. Similarly, the Notch-sparing compounds begacestat and avagacestat did not show significant higher selectivity for APP compared to the Notch substrate in our assay. These data raise serious concerns about the interpretation of inhibitory studies that relied on cellular or *in vivo* data, which in general do not allow direct quantitative comparisons as done with our assay. Importantly, our data do not discard the selective inhibition of APP at the ϵ -cleavage as a plausible strategy for drug development, but basically indicates that the approaches that have been used to reach this aim need to be revisited.

We also tested GSMs and found that all three candidates keep full functionality at the endopeptidase cleavage and regulate the carboxypeptidase-like activity by activating the fourth cycle of the γ -secretase, resulting in an increased processing of the aggregation-prone A β s towards shorter A β peptides. Our data, however, suggest some caution with this strategy as the tested compounds differentially affect the A β 48 > A β 38 versus the A β 49 > A β 40 pathway. The relative increase of A β 38 observed with all compounds needs further scrutiny, as APP clinical mutations also promote this production line and our initial data provided here (Supplementary Figure 5) suggest that A β 38 is less benign than A β 40 with regard to its interaction with A β 42. However, further research is needed to evaluate whether the weak (but significant) effects we see with A β 38 translate into an increased toxicity *in vivo*.

In conclusion, our work provides an important step forward towards the understanding of the mechanisms by which FAD mutations in *PSEN* and *APP* cause AD. Our findings support strongly the hypothesis that although these mutations affect γ -secretase in various ways, they all lead to qualitative shifts in the A β profiles, which provides a common denominator for the pathogenic effect of all FAD mutations.

Materials and methods

Antibodies

Rabbit polyclonal antibodies human PSEN1–NTF (B19.3), PSEN2–CTF (B24), APH-1a (B80.3), PEN-2 (B126.1) and APP C-terminus (B63.3) and monoclonal 9C3 against Nicastrin have been described (Annaert *et al*, 2001; Esselens *et al*, 2004). Rabbit monoclonal neopeptide AICD was obtained from Lilly Company. Other antibodies purchased were as follows: anti-FLAG M2 from Sigma, goat-anti-mouse IRDye800 from Rockland, goat-anti-rabbit Alexa680 from Invitrogen, 82E1 from Demeditec Diagnostics, MAB5232 and MAB1563 against PSEN1–CTF and human PSEN1–NTF from Chemicon, biotinylated anti-mouse IgG from Vector laboratories, streptavidin-HRP from GE Healthcare and 1E8 from Nanotools, Teningen, Germany. ELISA capturing antibodies purchased were as follows: JRF AB038 for A β 1–38 from Janssen; JRF/cAb40/28 for A β 1–40 from Janssen; JRF/cAb42/26 for A β 1–42 from Janssen; and A β 1–43 from Signet Labs Inc.. Detection antibody was obtained from Janssen; huAB25-HRPO (Zhou *et al*, 2011).

GSI and GSM

L-685,458 (Inhibitor X) was purchased from Calbiochem. Begacestat and the GSMs were synthesized according to the procedures reported in either primary publications or patents. GSM 1 was synthesized according to WO2006043064, GSM 2 (E-2012) according to WO2006112552, and WO2006046575, and GSM 3 according to WO2005115990. LY-450139 (semagacestat) and BMS-708163 (avagacestat) were obtained from Haoyuan Chemexpress Co. Limited, Shanghai.

Cell culture

Psen1/Psen2-deficient ($-/-$) MEF (*Psen1/2* $^{-/-}$ mEF) (Herreman *et al*, 2000) were cultured in Dulbecco's modified Eagle's medium/F-12 containing 10% fetal bovine serum. *Psen1/2* $^{-/-}$ mEF rescued with wt (human) *PSEN1* or L166P, G384A and DE9 as well as wt (human) *PSEN2* or N141I were reported before Bentahir *et al* (2006). The Y115H, M139V and I213T FAD–*PSEN1* cell lines were generated accordingly. mEF–*PSEN1* cell lines were transduced with a recombinant adenovirus Ad5/CMV-APP bearing human APP-swe, as previously described (Chavez-Gutiérrez *et al*, 2008). Neuronal cultures derived from E14 embryos and Semliki Forest virus transfection procedures have been described previously (Annaert *et al*, 1999). Semliki Forest viruses (SFV) were produced as described (Annaert *et al*, 1999). Briefly, brains from E14 embryos were trypsinized and plated on 6-cm dishes (Nunc) precoated with poly-L-lysine (Sigma–Aldrich). Cultures were maintained in neurobasal medium (Gibco) supplemented with B27 (GibcoBRL) and 5 mM cytosine arabinoside to prevent glial cell proliferation. After 3 days, neurons were transduced with SFV expressing wt or FAD mutant APP. After 1 and 3 h, post-infection media were refreshed. After 24 h, sA β were analysed by ELISA.

Expression and purification of substrates-3xFLAG

Substrate purification was performed as previously described (Chavez-Gutiérrez *et al*, 2008). Notch-, Erb4- and N-Cadherin-based substrates were designed to be similar in size to the APP substrate (C99–3XFLAG). Purity was assessed by SDS–PAGE and Coomassie staining (GelCode reagent, Pierce).

In vitro activity assays using solubilized γ -secretase

In vitro activity assays were done as previously described (Chavez-Gutiérrez *et al*, 2008), with minor modifications. MEF's microsomal fractions were prepared in 50 mM citric acid, pH 6.7, 0.25 M

Figure 6 Analysis of GSI and GSM. Dose-response inhibitory assays for (A) the transition state analogue (TSA) L-685,458 (InhX), (B) semagacestat, and the Notch-sparing compounds (C) begacestat and (D) avagacestat (see materials in Supplementary data) were performed using CHAPSO-extracted membranes from dKO PSEN1/2 MEFs stably expressing human wt PSEN1 as source of γ -secretase and $1 \times$ Km substrate concentrations (400 nM APP-C99-3XFLAG or 1μ M Notch-3XFLAG). Structures of the different compounds are displayed. *In vitro*-generated AICD (in black) or NICD (in red) are plotted as percentage of control reaction (DMSO). Error bars indicate s.d. ($n = 3$); except for semagacestat plot (s.e., $n = 5$). (E) Top panel: structures of the GSM tested. Low panel: increasing concentrations of GSM 1–3 did not change *in vitro* AICD generation, neither at 0.4μ M APP-C99 substrate ($1 \times$ Km) nor at saturating conditions (1.75μ M C99-3XFLAG). (F) Effect of increasing concentrations of GSM 1–3 on A β production at $1 \times$ Km APP-C99 substrate (0.4μ M): A β product/substrate ratios show that GSM 1–3 specifically activate the fourth cycle of the γ -secretase complex. In particular, GSM activate the A β 38 product line. Panel shows mean \pm s.e.; statistical significance of the data ($n = 4$) tested with ANOVA and Dunnett's post test, vehicle (DMSO) as control group; * $P < 0.05$, ** $P < 0.01$.

sucrose, 1 mM EGTA, complete PI and 1% CHAPSO. *In vitro* reactions were carried out in 50 mM citric acid, pH 6.7, 0.25 M sucrose, 1 mM EGTA, 1 × EDTA-free complete proteinase inhibitors (Roche), 2.5% DMSO and 0.05% phosphatidylcholine. Reactions were incubated for 4 h at 37 °C unless otherwise mentioned.

Lipids and substrates were extracted by adding 1 volume chloroform/methanol (2:1, v/v). Then, the aqueous fraction (ICD products) was taken and subjected to SDS-PAGE and quantitative western immunoblot. Known amounts of C99-3XFLAG were included as standards for absolute quantifications. ICD-3XFLAG and standards were determined with the anti-FLAG M2 and goat-anti-mouse IR800 antibodies, whereas the AICD₅₀₋₉₉ product was determined with a neo-epitope mAb and a goat-anti-rabbit Alexa680 secondary antibody. Infrared signals were detected using the Odyssey Infrared Imaging System.

Calculation of kinetics constants

Kinetic constants were estimated by nonlinear curve-fitting using GraphPad Prism 4 software. The equation $V = (V_{\max} \times [S]) / (K_m + [S])$ was used to calculate apparent K_m and V_{\max} values for the different enzymes, where V was experimentally determined using a range of substrate concentrations $[S]$. γ -Secretase activities were normalized to PSEN-CTF fragment levels or full-length PS1 levels for the DE9 mutant.

Quantification of soluble A β using sandwich ELISA

Ninety-six-well plates (NUNC) were coated with 1.5 μ g/ml A β capture antibody, excepting A β 43-ab coated at 7.5 μ g/ml, in a final volume of 50 μ l buffer (10 mM Tris HCl, 10 mM NaCl, 10 mM Na₂S₂O₃, pH 8.5). After overnight incubation at 4 °C, the plates were rinsed with PBS + 0.05% Tween 20 and blocked with 100 μ l per well of casein buffer (1 g casein in 1 l 1 × PBS, pH 7.4) for 4 h at room temperature. Samples and standards (synthetic human A β 1-38, A β 1-40, A β 1-42 or A β 1-43 peptides) were diluted in casein buffer. After overnight incubation at 4 °C, plates were rinsed and developed using 50 μ l per well of 100 mM NaAc pH 4.9/TMB (Sigma)/H₂O₂. Reactions were stopped with 50 μ l per well of 2 N H₂SO₄ and read on a Perkin Elmer Envision 2103 multilabel reader at 450 nm.

Urea gels

A β -peptides were analysed by a modified version of the urea-based SDS-PAGE (10% T/5% C instead of 12% T/5% C polyacrylamide

and 0.075 M instead of 0.1 M H₂SO₄ in the separation gel) (Wiltfang *et al*, 2002). Western immunoblot was done using 1E8, amplifying the signal with biotinylated anti-mouse IgG and streptavidin-HRP. Signals were detected using ECL chemiluminescence with an Intas Imager (Intas, Göttingen, Germany).

Supplementary data

Supplementary data are available at *The EMBO Journal* Online (<http://www.embojournal.org>).

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Author contributions: LC-G and BDS designed the study and wrote the manuscript. LB and LZ contributed to the analysis of FAD-APP mutations. AV, IB, JS, FR and KB contributed to A β aggregation assays and characterized the synaptotoxicity of A β peptides on primary neurons. MBo, MBe, SL and LS provided their technical and experimental assistance. SVC contributed to the experimental analysis of FAD-PSEN mutations. HE and JW characterized the A β profiles of FAD-PSEN mutants in urea-based gels. HG and EK synthesized and provided the GSI and GSM. All authors have read the manuscript and provided the input.

Conflict of interest

Harrie Gijssen is an employee of Janssen Pharmaceutica. Bart de Strooper is a consultant for Janssen Pharmaceutica, Envivo Pharmaceutics and Remynd and is supported by research grants from Janssen Pharmaceutics.

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