

The *Uppsala APP* deletion causes early onset autosomal dominant Alzheimer's disease by altering APP processing and increasing amyloid β fibril formation

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One Sentence Summary:

The *Uppsala APP* deletion leads to Alzheimer's disease by modulating α - and β -secretase cleavage of APP and by accelerating A β fibrillization.

Abstract:

Point mutations in the amyloid precursor protein gene (*APP*) cause familial Alzheimer's disease (AD) by increasing generation or altering conformation of amyloid- β ($A\beta$). Here we describe the *Uppsala APP* mutation ($\Delta 690-695$), one of the first reported deletions causing autosomal dominant AD. Affected individuals have an age at symptom onset in their early forties and suffer from a rapidly progressing disease course. Symptoms and biomarkers are typical of AD, with the exception of normal cerebrospinal fluid (CSF) $A\beta_{42}$ and only slightly pathological amyloid-PET signals. Mass spectrometry and western blot analyses of patient CSF and media from experimental cell cultures indicate that the *Uppsala APP* mutation alters APP processing by increasing β -secretase cleavage and affecting α -secretase cleavage. Furthermore, in vitro aggregation studies and analyses of patient brain tissue samples indicate that the longer form of mutated $A\beta$, $A\beta_{Upp1-42\Delta 19-24}$, accelerates the formation of fibrils with unique polymorphs and their deposition into amyloid plaques in the affected brain.

Introduction

Alzheimer's disease (AD) is neuropathologically characterized by a progressive deposition of amyloid β ($A\beta$) in parenchyma and blood vessels of the cerebrum (1). Upon sequential cleavage of the amyloid precursor protein (APP) by β - and γ -secretases, $A\beta$ peptides of 38-43 amino acids are generated. If instead α -secretase cleavage occurs, no $A\beta$ is formed (reviewed in (2)).

Increased generation of the more amyloidogenic $A\beta_{42}$ is seen for several of the *APP* mutations positioned in vicinity of the γ -secretase cleavage site (3-14), whereas the *Swedish* mutation close

to the β -secretase cleavage site results in increased production of both A β 42 and A β 40, as demonstrated in plasma and fibroblasts from mutation carriers (15-17).

Pathogenic *APP* mutations within the A β sequence have been described to result in various disease phenotypes. Patients with the *Dutch* (E693Q) and *Italian* (E693K) mutations display amyloid accumulation in cerebral blood vessel walls and intracerebral hemorrhage (18, 19), whereas carriers of the *Flemish* (A692G) and *Iowa* (D694N) mutations suffer from both intracerebral hemorrhage and progressive dementia (20, 21). The *Arctic* mutation (E693G) leads to an increased formation of protofibrils (22) and other large A β oligomers with particularly neurotoxic properties (23). Clinical examinations and neuropathological analyses confirmed that carriers of the *Arctic* mutation have an AD phenotype (24), although their brains almost only display diffuse parenchymal A β deposits (24-26).

The only protective *APP* variant described to date, the *Icelandic* mutation (A673T), has been shown to decrease β -secretase cleavage resulting in reduced A β production (27) and aggregation (28, 29).

For the only identified disease-causing *APP* deletion (E693 Δ), resulting in a recessive form of familial AD, an increased intraneuronal presence of toxic A β oligomers was suggested as an underlying pathogenic feature. A decreased inhibition of both β - and γ -secretase, with increased enzymatic activities and relative resistance to degradation of mutant A β by neprilysin and insulin-degrading enzyme, have been proposed as other effects of this deletion (30).

Here, we report the pathogenic *APP* deletion (690-695 Δ) that causes a dominantly inherited form of early onset dementia in three mutation-carriers of a family originating from the city of Uppsala,

Sweden. Clinical and neuropathological examinations are compatible with AD and experimental studies indicate that the phenotype is caused by pathological alterations of the β - and α -secretase cleavage of APP, which result in increased A β production in combination with a very rapid aggregation of the longer A β mutant (A β Upp1-42 Δ 19-24) into unique polymorphic structures.

Results

Epidemiological and clinical features of the *Uppsala APP* mutation family

The *Uppsala APP* mutation was detected in two siblings and their cousin, who were all referred to the Memory Disorder Unit, Uppsala University Hospital (for pedigree, see Fig. 1A). The ages of symptom onset were 43 years (sibling 1), 40 years (sibling 2) and 41 years (cousin). All three patients had a manifest cognitive impairment and scored 20-22 points on the MMSE at the time of referral, with word finding difficulties, dyscalculia, apraxia and visuospatial / executive impairment as major symptoms. Sibling 1 developed myoclonus and had a rapid disease progression with severe anxiety and behavioral disturbances. Death occurred six years after onset, at the age of 49. At the initial neuropsychological evaluation sibling 2 had normal scores on episodic memory tests, but featured severe dyscalculia and problems with the clock drawing test. At the four years follow-up examination this patient had become increasingly affected by apathy and mutism. The cousin of the two siblings displayed impaired episodic memory, language and executive functions as dominant symptoms.

One parent of the two siblings had been referred for assessment more than 20 years earlier after having refused to see a physician for several years. This patient had symptom onset at about 47 years of age and the diagnosis of AD was supported by a computerized tomography (CT) scan, showing cortical and central atrophy at the age of 54. Death occurred at the age of 60. One of the

parents of the cousin had onset of symptoms at approximately 45 years of age and was subsequently also diagnosed with AD. This patient became aggressive and spent several years at a geropsychiatric ward before death occurred approximately fifteen years later. In addition, one of the siblings' grandparents developed dementia with onset at approximately 40 years of age and died at the age of 51 (Fig. 1A).

All three cases underwent lumbar puncture and subsequent cerebrospinal fluid (CSF) analyses. CSF concentrations of A β 42 were within the normal range of non-AD controls, whereas concentration of total tau (t-tau) and phospho-tau (p-tau) were pathologically elevated (table S1).

Brain imaging

The two siblings and their cousin underwent CT brain examinations at the time of diagnosis. The scan of sibling 1, who was in a more advanced disease stage at the initial visit, showed medial temporal lobe atrophy (MTA) grade 2 together with a moderate frontoparietal lobe atrophy. Sibling 2 and his cousin had a moderate global cortical and central atrophy, whereas the temporal lobes were well preserved (MTA grade 0-1) (Fig. 1B). Moreover, the siblings and their cousin underwent fluorodeoxyglucose positron emission tomography (FDG-PET), which showed a decreased uptake mainly in the temporal and parietal lobes (Fig. 1C-D). Two of them also underwent amyloid-PET (Pittsburg compound B/PIB), which demonstrated a pathological pattern but only with a slightly increased accumulation of PIB in cortical areas (Fig. 1E-F).

Genetic analyses reveal the *Uppsala APP* mutation

In the three affected cases (two siblings and their cousin), we identified an 18 base pair deletion in exon 17, which leads to the loss of six amino acids (690-695 Δ) within A β . It should be noted that this deletion spans over the region that is affected by previously identified intra-A β mutations

(Fig. 1G). In addition, we have analyzed more than 500 DNA samples from Swedish patients with AD, older unaffected family members, as well as from older healthy control subjects, all of which were negative for this genetic alteration. Furthermore, the two siblings were analyzed for the apolipoprotein E gene (*APOE*) and found to be *APOE* $\epsilon 3$ homozygotes.

The *Uppsala APP* mutation leads to mainly A β Upp1-42 _{Δ 19-24} pathology

One brain of an *Uppsala APP* mutation carrier (sibling 1) has come to autopsy. The weight of the brain was 1480 g and the right hemisphere was subjected to routine analyses. On gross inspection, dilated ventricles were evident. Microscopically, a pronounced gliosis was seen in limbic as well as in neocortical areas (Fig. 2A). Moreover, there was a widespread tau pathology, corresponding to Braak stage VI, as visualized with the AT8 anti-tau antibody (Fig. 2B-C).

Amyloid- β plaques were abundant and widespread, corresponding to Thal stage 5. Tissue sections from occipital (Fig. 2), temporal and parietal (fig. S1) neocortices were analyzed by immunohistochemistry with a panel of different monoclonal anti-A β antibodies and by thioflavin S (ThS) staining. With the 6F/3D antibody, binding to A β amino acids 8-17 (25), abundant pathology was observed in several neocortical regions (Fig. 2D and fig. S1) and a similar pattern could be observed with ThS (Fig 2E). Antibodies directed towards the A β C-terminus revealed abundant A β 42 staining, whereas A β 40 staining was much less intense in all cortical areas investigated (Fig. 2F-I and fig. S1).

Furthermore, brain tissues from fresh frozen frontal, temporal and occipital neocortex, as well as cerebellum of the mutation-carrier, eleven sporadic AD (sAD) and nine non-neurological control brains (table S2) were homogenized and sequentially extracted with tris-buffered saline (TBS) and formic acid (FA) for analysis with MSD electrochemiluminescence-based A β immunoassay

(Meso Scale Discovery) and ELISA. Compared to sAD and control samples, the *Uppsala APP* mutation brain displayed lower concentrations of A β 40 whereas concentrations of A β 42 were elevated, especially in the FA fraction (Fig. 3A), which corresponds to insoluble A β deposits, but also in the TBS fraction, representing more soluble A β , including aggregates (Fig. 3B). In contrast to elevated A β 42 in the FA fraction of all investigated brain regions, concentrations of TBS soluble oligomers were lower in the *Uppsala APP* mutation brain compared to sAD, when analyzed with an ELISA that detects soluble A β aggregates of all sizes (Fig. 3C). Further, when analyzed with an ELISA that preferentially recognizes larger oligomers and protofibrils (31), the *Uppsala APP* mutation brain displayed low concentrations, comparable to those in the controls, whereas the amounts of such A β species in sAD brains were elevated (Fig. 3D).

The composition of amyloid plaques from temporal neocortex was further analyzed by luminescent conjugated oligothiophene (LCO) staining and matrix-assisted laser desorption/ionization (MALDI) imaging mass spectrometry (IMS). Similar to those that were positive for ThS, plaques stained with the LCOs q-FTAA and h-FTAA (Fig. 3E) showed a distinct core surrounded by a diffuse halo of fibrillar A β (Fig. 3F). The MALDI-IMS analyses suggested that the plaques mainly consist of A β Upp42 Δ 19-24, either in its full-length version or as N-terminally truncated peptides, which mainly start at positions 3 (pyroglutamate), 4, 5 or 8. The contribution by A β Upp1-40 Δ 19-24, A β wt1-40 and A β wt1-42 to the formation of amyloid plaques in the *Uppsala APP* mutation brain seemed to be minor (Fig. 3G-H). These results, as well as the peptide sequence identity were confirmed by immunoprecipitation (IP) and mass spectrometry (MS) analyses of pooled material from 50 individually laser microdissected plaques, identified with LCO staining (table S3).

In order to investigate the contribution of A β Upp and A β wt in CSF from *Uppsala APP* mutation cases and thereby understand why A β 1-42 CSF concentrations were within the normal range in these patients, we performed IP-MS analyses using 6E10 (A β amino acid 5-10) and antibodies targeting the A β 40 and A β 42 C-terminus for immunoprecipitation. Cerebrospinal fluid from the three mutation carriers (sibling 1, sibling 2 and cousin) was analyzed and compared to CSF from eleven sAD cases and ten healthy control subjects (Fig 4A) (demographic information, table S1). Mutation carriers displayed higher CSF amounts of A β 1-40, A β 1-42 and total A β (sum of all detected A β variants) as compared to sAD cases and control subjects (Fig. 4B). The amounts of A β wt1-40 in CSF, produced from their non-mutated *APP* allele, was lower in patients with the *Uppsala APP* mutation than in sAD cases and control subjects, whereas A β wt1-42 was not different in patients with the mutations compared to controls (Fig. 4C). The *Uppsala APP* mutation carriers displayed a relative increase of both A β Upp1-40 $_{\Delta 19-24}$ and A β Upp1-42 $_{\Delta 19-24}$ (Fig. 4D). Thus, the expression of A β from the mutated allele probably accounts for the unexpectedly high A β 1-42 CSF concentrations in the routine analysis of patients with the *Uppsala APP* mutation. These measurements were performed with an immunoassay that should detect A β wt1-42 and A β Upp1-42 $_{\Delta 19-24}$ equally well. However, a comparison of A β 1-42 measurements performed with the routine immunoassay and IP-MS (fig. S2A) showed that whereas values from control and sAD samples correlated well between the two methods, those from the three *Uppsala APP* mutation cases did not (fig. S2B). Ion spectra from A β Upp1-42 $_{\Delta 19-24}$ and A β Upp1-40 $_{\Delta 19-24}$ are shown in fig. S3.

The Uppsala APP mutation alters APP processing, resulting in increased A β production

To study the potential effects of the *Uppsala APP* mutation on APP processing, conditioned media of HEK293 cells transfected with *APP* carrying the *Uppsala* mutation (*APP^{Upp}*) or wild-type *APP* (*APP^{wt}*) were analyzed with MSD immunoassays to determine sAPP α and sAPP β

concentrations, soluble APP fragments resulting from α - and β -cleavage, as well as A β 40 and A β 42.

Only background amounts of sAPP α were detected in media from the *APP^{Upp}* culture, whereas high concentrations were found in media from *APP^{wt}* expressing cells (Fig. 5A). In contrast, concentrations of sAPP β were higher in media from *APP^{Upp}* than *APP^{wt}* cells (Fig. 5B), which was also reflected in a higher concentration of both A β 40 and A β 42 (Fig. 5C).

To study whether the *Uppsala APP* mutation causes an ablation of α -cleavage or whether the cleavage site is shifted towards the N-terminus due to the six amino acid deletion, western blot was performed on the same cell media as used for the MSD analyses. Total soluble APP (detected with 22C11, binding to an N-terminal epitope of APP) was similar in cell media from *APP^{Upp}* and *APP^{wt}* transfected cells. Similar to the MSD analyses, no sAPP α could be detected in the media from the *APP^{Upp}* culture when probing with 6E10 (directed against A β amino acid 5-10 and usually present on sAPP α) ($P=0.0004$) or 2B3 (specific for the C-terminal end of sAPP α) ($P=0.0045$). However, when detected with mAb1C3 which binds to A β amino acid 3-8, which closer to the APP N-terminus than 6E10 (fig. S4), a faint sAPP α band was observed ($P=0.0291$) (Fig. 5D-E).

Next, the same cell media were analyzed with two different A β sandwich ELISAs: A β 1-40 (using A β N-terminal specific antibody 3D6 for detection) and A β x-40 (using 6E10 for detection). Since the MSD and Western blot-based results suggested an additional cleavage site in *APP^{Upp}* resulting in mAb1C3 positive sAPP fragment, we expected that the C-terminal side of this cleavage site would be detectable with the A β x-40 ELISA (Fig. 5F). Indeed, cell media from the *APP^{Upp}* culture showed a significantly higher ($P=0.0016$) A β x-40/A β 1-40 ratio compared to

APP^{wt} transfected cells, indicating that, in addition to A β , an extra N-truncated A β fragment was present in the *APP*^{Upp} cell media (Fig. 5G).

To confirm if the reduction of sAPP α in *APP*^{Upp} cell media was α -secretase cleavage specific, we performed western blot with the same constructs as in the other cell culture-based experiments. When probing with the sAPP α specific antibody 14D6, sAPP α was found to be significantly ($P<0.0001$) reduced in *APP*^{Upp} compared to *APP*^{wt} cell media and was markedly reduced ($P<0.01$) in media from both cell cultures treated with a metalloprotease inhibitor (GI) that blocks ADAM 10, the major α -secretase (32) (Fig 5 H-I). In addition, and in line with the MSD results (Fig. 6B), sAPP β was increased ($P<0.01$) in media from *APP*^{Upp} transfected cells (Fig. 5 H-I). A mild reduction in total sAPP was observed (Fig. 5 H-I).

Moreover, we performed western blot on the cell lysates using a γ -secretase inhibitor (DAPT) to detect C-terminal fragments (CTF) resulting from α - and β -secretase cleavage of APP and found a slight reduction in CTF α and a more prominent increase in 6E10-positive CTFs (fig. S5).

In addition to the western blot analyses, we applied MS to investigate the altered APP processing. With respect to *APP*-transfected cells, subjected to immunoprecipitation with 6E10, anti-A β 40 and anti-A β 42, the most prominent forms of A β were A β 1-40 and A β 17-40 in media from *APP*^{wt}-transfected cells (Fig. 6A), whereas A β ^{Upp}1-40 Δ 19-24 and A β ^{Upp}5-40 Δ 19-24 were the dominating species in media from *APP*^{Upp}-transfected cells (Fig. 6B), suggesting a new major cleavage of *APP*^{Upp} between amino acids 4 and 5 in the A β sequence.

Furthermore, affinity purified sAPP was digested with the protease LysN, which cleaves proteins at the N-terminal side of lysines, followed by liquid chromatography tandem MS analyses of

peptides that are specific for the cleavage sites of α -secretase and β -secretase. Two different peptides in the N-terminal part of APP, upstream of the α -secretase and β -secretase cleavage sites, respectively, could then be found at similar concentrations in *APP^{Upp}* and *APP^{wt}* cell media (Fig. 6C-D). Analysis of the α -secretase cleavage site-specific peptide KMDAEFRHDSGYEVHHQK (595-612 in wt hAPP695) showed that cleavage at this site was strongly reduced in *APP^{Upp}* compared to *APP^{wt}* media (Fig 6E). In contrast, the intensity of the β -secretase cleavage site-specific peptide KTEEISEVKM(ox) (587-596 in wt hAPP695) was strongly increased in *APP^{Upp}* in comparison to *APP^{wt}* media, indicating increased cleavage by BACE1, the major β -secretase (Fig. 6F). Moreover, the semi specific peptide (N-terminus specific for LysN, C-terminus unspecific) KMDAEFRHDSGY (595-606 in wt hAPP695) was identified. It ends at amino acid 10 of the A β sequence (same as the β' -secretase cleavage site), and was found to have a much higher intensity in media from *APP^{Upp}* compared to *APP^{wt}* cells (Fig. 6G). In addition, a peptide generated upon cleavage between amino acid 4 and 5 of the A β sequence, KTEEISEVKMDAEF (587-600 in wt hAPP695), was strongly increased in *APP^{Upp}* media (Fig. 6H). These MS-based results thus demonstrate activity at two major cleavages sites within the N-terminal part of the A β domain of *APP^{Upp}*. The effects on APP processing revealed by MS analyses are summarized in Fig. 6I. MS sequencing results are shown in fig. S6.

A β 1-42^{Upp} _{Δ 19-24} is prone to form amyloid fibrils in vitro and displays a unique structural polymorphism

In order to investigate the aggregation behavior of A β ^{Upp} we performed in vitro aggregation experiments. Monomeric A β was extracted from solubilized lyophilized synthetic A β ^{Upp}1-42 Δ 19-24, A β ^{wt}1-42, and A β ^{Arc}1-42 peptides with HPLC-SEC and analyzed with three different methods upon aggregation at 37°C without shaking.

292

293 First, we analyzed fibril formation with the Thioflavin T assay (ThT), which revealed that
294 A β Upp1-42 Δ 19-24 fibrillizes very rapidly, reaching half of its maximum ThT signal after 0.93 h,
295 compared to 8.3 h for A β wt1-42 and 1.3 h for A β Arc1-42 $E_{22}G$. A β Upp1-42 Δ 19-24 thus aggregated
296 significantly faster than A β wt1-42 ($p < 0.0001$), and with a similar rate as A β Arc1-42 $E_{22}G$, albeit
297 apparently with a somewhat less prominent lag phase (Fig. 7A). The A β Upp1-40 Δ 19-24 peptide
298 did not display any fibril formation and was therefore not included in the analysis. Next, we
299 applied an ELISA that selectively measures soluble A β oligomers/protofibrils (33) and found that
300 the concentration of such A β species decreased for A β Upp1-42 Δ 19-24, whereas they increased with
301 time for A β wt1-42 and A β Arc1-42 (Fig. 7B). In the same samples (isolated at 1 h and 8 h), the
302 distribution of protofibrillar and monomeric A β was qualitatively visualized with size exclusion
303 chromatography (SEC). Prior to SEC analysis, fibrillar A β was pelleted with centrifugation and,
304 hence, also fibril formation was indirectly monitored as a decreased size of the protofibril and
305 monomer peaks. Overall, SEC largely confirmed the results from the ThT assay and ELISA, with
306 a prominent decrease of monomeric A β Upp1-42 Δ 19-24 over time (from 1 h to 8 h). In addition,
307 similar to the ThT assay and ELISA data and in contrast to A β wt1-42 and A β Arc1-42, no increase
308 in the protofibril peak could be observed over time for A β Upp1-42 Δ 19-24 (Fig. 7C).

309

310 To further investigate the structural polymorphism of A β formed as a result of the *Uppsala APP*
311 mutation, A β Upp1-42 Δ 19-24 was fibrillized under a low pH condition that has previously been
312 shown to yield slow-growing and well-ordered A β wt1-42 fibrils (34). Electron microscopy (EM)
313 imaging of negatively stained fibrils revealed the presence of long and well-ordered fibrils with
314 at least four different polymorphs (Fig. 7D). For higher resolution examination of these fibrils,
315 cryo-EM experiments were performed. For the two most dominant polymorphs we obtained 3D
316 density reconstructions at resolutions of 5.7 Å and 5.1 Å for polymorph 1 and polymorph 2,

respectively (Fig. 7D). Both fibrils were found to consist of two symmetric protofilaments. The density allowed for building a tentative backbone trace but since no side-chain density was visible, the amino acid sequence and also the N- and C- termini could not be assigned to the backbone trace. For both polymorphs, all 36 residues could be accommodated by the density and thus seem to be part of the folded A β Upp1-42 Δ 19-24 fibril.

Polymorph 1 (Fig. 8D, middle) resembles an A β wt1–42 fibril structure that has been earlier described (35) and possibly shares the same protofilament interface in the fibril core (fig. S7). Polymorph 2 has a vague similarity to the previously described A β wt1–42 structure (34), where the prominent salt bridge between the N-terminal Asp1 and Lys28 residues could be present also in A β Upp1-42 Δ 19-24 fibrils (fig. S7). However, the C-terminal protofilament interface in the core of the fibril is instead very similar to the interface in two solid-state NMR structures of A β wt1–42 (36, 37) (fig. S8).

Discussion

We here describe the *Uppsala APP* mutation, an *APP* deletion causing a dominantly inherited form of AD. This pathogenic deletion, resulting in a loss of six amino acids in the mid-region of A β , was found in three affected family members and not in 500 other subjects, including older non-affected family members, older healthy controls and sporadic AD cases.

Most of the AD-causing *APP* mutations lead to symptom onset between 40 and 65 years (38), although cases with an even earlier onset have been reported for some mutations (9, 12, 39). The clinical effects of the *Uppsala APP* mutation are severe, insofar that mutation carriers develop symptoms already in their early forties and have an aggressive disease course. The clinical picture involves severe dementia, characterized by wide-spread parieto-temporal lobe involvement,

leading to death from dementia-related illnesses within 5-11 years. These clinical characteristics are thus similar to AD in general and to what has been reported for other familial disease variants.

In terms of structural brain imaging, the CT scans displayed the expected symmetrical pattern of global cortical atrophy and mild medial temporal lobe atrophy. As for PET, investigations with the [^{18}F]FDG ligand showed a disease-characteristic hypometabolism of posterior parietal and temporal lobes whereas analysis with [^{11}C]PIB, which selectively binds to amyloid plaques, only showed a slightly positive pattern. Analyses of *post mortem* brain tissue from one of the affected cases resulted in several important observations. Firstly, the pathological picture was compatible with AD, including abundant deposition of extracellular A β -positive plaques as well as intracellular tau-positive tangles and neurites accompanied by pronounced gliosis. The regional distribution of A β aggregates was extended from neocortex to cerebellum, corresponding to Thal phase 5 and p-tau pathology was observed from locus coeruleus to neocortex, corresponding to Braak stage VI. Secondly, upon a more detailed examination of tissues from different cortical areas, it became evident that the A β pathology of the *Uppsala APP* mutation carriers mainly consists of A β 42. This observation was corroborated by MALDI-IMS analyses of A β plaques from the temporal cortex, which in addition suggested that almost only mutated A β was present, either in its full-length version or as N-terminally truncated forms. Thirdly, A β plaques were positive for staining with the amyloid dye ThS, which is structurally similar to PIB, raising the question why patients were only slightly [^{11}C]PIB-PET positive, despite high total A β concentrations in the *post mortem* brain tissue analysis. The PET results are displayed as a standard uptake value ratio (SUVR), which is a ratio of the PET signal in the region of interest to the signal from a reference region, in this case the cerebellum. Hence, a low SUVR could have been explained by a high reference region signal but this interpretation could be ruled out as ThS staining of cerebellum in the patient with *Uppsala APP* revealed a low amyloid burden in this

367 brain region. Although amyloid plaques generally reach the plateau phase rather early in the
368 disease course, we cannot rule out that the time between the scan and the *post mortem* analyses
369 could not explain these difference in patient with *Uppsala APP*. Subtle changes in the fibrillar
370 structure of A β Upp Δ 19-24 could be another potential explanation for the low PIB retention signal
371 seen for the patients with *Uppsala APP*.

372
373 Biochemical analyses of AD CSF biomarkers revealed the expected pathological increase of t-
374 tau and p-tau whereas, unlike other *APP* mutation cases (40), concentrations of A β 42 were
375 normal for all three *Uppsala APP* mutation carriers investigated. The IP-MS-based CSF analyses
376 suggest an explanation for this unexpected finding, as they demonstrate that the amounts of
377 A β Upp1-40 Δ 19-24 and, especially, A β Upp1-42 Δ 19-24 produced by the mutated allele were
378 substantially higher than A β wt1-40 and A β wt1-42 generated from the non-mutated allele. Thus,
379 an increased generation of A β from the allele with the *Uppsala APP* mutation seems to result in
380 higher total CSF concentrations of A β 1-40 and A β 1-42 as compared to both sporadic AD cases
381 and controls. Comparison the IP-MS and routine ELISA based CSF A β data revealed a good
382 correlation for all control and sAD samples, whereas for the *Uppsala APP* mutation carriers IP-
383 MS-generated CSF A β concentrations were relatively higher. The ELISA measurement displayed
384 normal concentrations of A β 1-42 in CSF from *Uppsala APP* mutation cases, which is higher than
385 in sAD cases, but inaccurately in the same range as control subjects. We speculate that this
386 discrepancy may be related to a difference in conformation between A β Upp Δ 19-24 and A β wt.
387 Whereas ELISA detection of A β 1-42 relies on the simultaneous binding of the assay antibodies
388 to the C- and N-terminus of A β , the IP-MS method is only dependent on one antibody-A β
389 interaction at a time, which could facilitate its detection of A β Upp Δ 19-24. We therefore believe
390 that the IP-MS results in this case better reflect the true amounts of A β 1-42 in CSF, which in turn

suggests a substantially increased production of A β in the brain of *Uppsala APP* mutation carriers.

We next performed cell-based experiments to seek a molecular explanation for the difference in A β production from the mutant and wild-type alleles in the mutation carriers. The MSD and western blot-based analyses of cell media from *APP* transfected cells, together with analyses of CTF fragments with the 2C11 (C-terminal APP) and 6E10 (A β 5-10) antibodies on the same cell model, demonstrated an increased production of sAPP β accompanied by higher concentration of A β in medium from cells transfected with *APP^{Upp}*. The MS analyses of the same cell culture media confirmed an increased β -secretase cleavage, thereby providing an explanation to the elevated amounts of A β Upp Δ 19-24 observed with IP-MS in CSF from the *Uppsala APP* mutation carriers. The detection of an increase in both A β and sAPP β by different methods confirms that the increased CSF A β Upp Δ 19-24 detected by IP-MS was not a method related artifact.

Since the N-terminal start of the *Uppsala APP* mutation is located only two amino acids from the α -secretase cleavage site, and this enzyme is dependent on the distance from the membrane and not exclusively on a determined cleavage site (41-43), we reasoned that the mutation may also affect α -secretase activity and/or the location of the cleavage site itself. In line with this, we could demonstrate that α -secretase-related APP processing is indeed altered by the *Uppsala APP* mutation. MSD and western blot analyses of conditioned media from HEK293 cells transfected with *APP^{Upp}* suggested a strongly decreased α -secretase cleavage at position A β 16-17. This was further strengthened by analysis of corresponding CTF fragments with the C-terminal APP (2C11) antibody in the same cell model. Moreover, with enzyme inhibition experiments we could demonstrate that the decreased cleavage indeed was specific to α -secretase. Furthermore, western blot analysis revealed an additional APP fragment that was faintly detected with mAb1C3 (which

binds to A β amino acids 3-8), but not with 6E10 (which binds to A β amino acids 5-10) or 2B3 (epitope to the C- terminal end of APP α) antibodies, suggesting possible alternative cleavage sites.

To identify the alternative cleavage sites, IP-MS analyses of sAPP fragments secreted in media from *APP^{Upp}* or *APP^{wt}* transfected cells indicated that the mutation results in a new major cleavage site located twelve amino acids N-terminally of the conventional α -secretase cleavage site, between amino acid 4 and 5 of the A β sequence. This site was also detected with MS analysis of LysN digested peptides from media of *APP^{Upp}* transfected cells. The resulting peptide, A β Upp5-40 Δ 19-24, was present in media from *APP^{Upp}* cells to a similar extent as A β wt17-40 (also known as p3) was in media from *APP^{wt}* cells. Further, A β Upp5-40 Δ 19-24 was identified by IP-MS in CSF from *Uppsala APP* mutation carriers, albeit less abundantly than in media from *APP^{Upp}* transfected HEK293 cells, a cell type which usually has a much higher activity of α -secretase than β -secretase as we could observe with the MSD analyses of the cell culture media. In addition, A β Upp5-42 Δ 19-24, likely resulting from the same enzymatic cleavage, was detected by MALDI-IMS in the brain, suggesting that it co-aggregates with A β Upp1-42 Δ 19-24 in plaques.

Whether A β Upp5-40 Δ 19-24 seems to be consistently present as a result of the *Uppsala APP* mutation, it is at this point unclear whether it is generated as a result of cleavage at an alternative α -secretase site or by some other protease. Several additional proteolytic cleavages of APP may occur within or just outside of the A β sequence, for example by BACE2 or proteases referred to as delta- and eta-secretases, or by alternative β -secretases, such as meprin-b (44-52). We speculate, however, that A β Upp5-40 Δ 19-24 could be an alternative version of p3 resulting from a shifted α -secretase cleavage, which co-aggregates with A β Upp1-42 Δ 19-24 to form plaques and thereby contributes to the pathogenesis in mutation carriers. As additional support of its potential

pathogenic significance, A β wt5-42 has in a previous study been found to have similar toxicity as A β wt1-42, but with an even higher propensity to aggregate (53). Furthermore, two previous studies have shown that treatment with a BACE1 inhibitor resulted in increased concentrations of A β 5-40/42 (54, 55), indicating that the cleavage of *APP*_{wt} between A β 4 and A β 5 indeed is independent of β -secretase. Irrespective of the nature of the secretases involved our data, together with these previous observations, thus suggest that both the β -secretase and the new cleavage site N-terminally of the α -secretase cleavage site are altered by *APP*_{Upp}.

An additional major cleavage site between amino acid 10 and 11 of the A β sequence was found to be increased in *APP*_{Upp} compared to *APP*_{wt} cell media. Accordingly, A β 11-40 Δ 19-24 and A β 11-42 Δ 19-24 were identified in CSF from patients with the *Uppsala APP* mutation and in *APP*_{Upp} cell media. Moreover, A β 11-42 Δ 19-24 was abundant in the plaques of the mutation carrier brain, likely as a consequence of increased cleavage at this site. This additional cleavage occurs at the β' cleavage site and it is at this point uncertain whether the observed activity with the *Uppsala APP* mutation is due to a general increase of BACE1 activity that affects both β -secretase sites in *APP*_{Upp}, or if it represents an additional shifted α -secretase cleavage site resulting from the deletion of six amino acids in the *APP*_{Upp} sequence.

Thus, the *Uppsala APP* mutation seems to abolish the non-amyloid generating pathway of APP processing, which may further contribute to the pathogenesis in affected individuals. However, in order to confirm the responsible protease(s) of the additional cleavage sites, additional experimental studies using different protease inhibitors are needed.

In order to investigate the inherent properties of A β , we next performed in vitro studies that examined the aggregation behavior of the mutated peptides. Upon analyzing results generated by

the ThT assay it became evident that A β Upp1-42 Δ 19-24 was forming *bona fide* fibrils very rapidly. With respect to oligomers or protofibrils, both A β wt1-42 and A β Arc1-42 formed such intermediately sized soluble aggregates, that increased with time, which is in line with what has been proposed as the pathogenic mechanism for the *Arctic* mutation (22). However, A β Upp1-42 Δ 19-24 oligomer/protofibril concentrations decreased over time, probably because the in vitro fibril formation was so rapid and complete that intermediate species were immediately fibrillized. This theory may be supported by the finding that, compared to sAD cases, TBS extracts of the *Uppsala APP* mutation carrier brain displayed lower concentrations of oligomers/protofibrils, especially larger variants, which may suggest that A β Upp Δ 19-24 aggregates into smaller-sized oligomers which rapidly fibrillize and deposit into plaques. In spite of this, oligomers/protofibrils of A β are likely to be of relevance for the pathogenesis of AD caused by the *Uppsala APP* mutation.

Structural analyses of two different polymorphs of the *Uppsala APP* mutation revealed that they share some features of previously published A β 1–42 fibril structures, but generally differ from all A β fibril structures that have been described to date. Further studies will be required to determine the effects of these structural polymorphs with respect to how they interact with amyloid dyes, such as PIB, and how they may contribute to the formation of toxic A β oligomers. Such oligomer formation could be driven by secondary nucleation, which has previously been reported to depend on the structure of the fibrillar surface where such a process is believed to occur (57).

At the best of our knowledge, the only previously described intra-A β *APP* deletion is the *Osaka* mutation (30). Whereas this mutation has been reported to have a recessive character, the *Uppsala APP* mutation is inherited in a dominant manner. Overall, A β Upp1-42 Δ 19-24 seems to be forming

amyloid fibrils much more aggressively than the corresponding form of A β _{Osaka}. For example, an *Osaka APP* mutation knock-in mouse model was reported to display brain pathology only when the inserted gene was expressed homozygously (58). Interestingly, in the *Osaka APP* mutation mouse model a reduced α -secretase cleavage could be observed, similar to what we here report for the *Uppsala APP* mutation (59).

Taken together, we have identified an *APP* mutation, which is an intra-A β deletion causing dominantly inherited AD. The loss of six amino acids results in an increase of the A β promoting β -secretase cleavage, leading to an elevated generation of A β Upp Δ 19-24 with a concomitant suppression of the regular α -secretase cleavage. Thus, the non-amyloid generating pathway is seemingly abolished with the mutation. Instead, two other A β species, A β Upp5-40/42 Δ 19-24 and A β Upp11-40/42 Δ 19-24, are formed, possibly as a result of a shift of the α -secretase cleavage site, and these may contribute to disease development in mutation carriers. The facts that A β wt5-42 has previously been reported as pathogenic and that A β Upp5-42 Δ 19-24 was found to be present in plaques from the investigated *Uppsala APP* mutation brain, support that at least this species may be contributing to the pathogenesis. Moreover, the mutation also renders unique properties to A β Upp1-42 Δ 19-24, which accelerates its fibrillization into distinctive polymorphs and promotes plaque deposition in the affected brains. Thus, the combined effect of three putative pathogenic mechanisms by the *Uppsala APP* mutation may well explain why affected carriers develop an aggressive form of disease with a very early age at symptom onset.

Although the study has clearly identified that the *Uppsala APP* mutation causes AD by a combination of three mechanisms, all related to APP, it is based on a limited patient material from *Uppsala APP* mutation carriers (CSF, n=3 and brain, n=1), which limits the statistical power of certain analyses. Therefore, the exact quantitative impact of the mutation on development of

A β and downstream pathologies is difficult to assess, as a certain individual variation between patients is to be expected. Further, although we have identified alterations in APP processing, at both the α - and β -cleavage sites, it remains to be confirmed by which enzymes the cleavage occurs. Future studies will also be needed to elucidate the impact of each of the three disease mechanisms presented here, as well as the temporal and structural aspects of the development of A β pathology. Some of these future studies could be performed in genetically modified mice carrying the Uppsala APP mutation.

Material and Methods

Study design

This study was designed to characterize the clinical and mechanistic features of the herein identified *Uppsala APP* mutation, which results in early onset familial AD. Three members of the ‘Uppsala family’ showing manifest of AD symptoms were identified as mutation carriers and subjected to clinical evaluation, structural and amyloid brain imaging and lumbar puncture for analysis of CSF biomarkers. Further, brain tissue from one of the mutation carriers was analyzed *post mortem* to assess a range of pathological markers – A β , tau and neuroinflammation – and stage the pathology according to established criteria. Brain tissue was also analyzed with MALDI imaging and immunoassays to investigate the nature of A β pathology in comparison with groups of sporadic AD brains (n=11) and neurologically normal control brains (n=9). To study the mechanistic properties of the *Uppsala APP* mutation, mass spectrometry and immunoassays were used to analyze A β and APP fragments resulting from APP processing. Such studies were performed in a) CSF from the three *Uppsala APP* mutation carriers in comparison with CSF from sporadic AD (n=10) and control (n=10) and b) medium and lysate from cell cultures transfected with APP harboring the *Uppsala APP* mutation in comparison with wild type APP. Finally, the

aggregation behavior and structure of A β aggregates were studied with ThT aggregation assay and cryo-electron microscopy. Sample sizes for brain tissue and CSF studies were determined to achieve a statistical power of 80%, based on group differences and variability from previous experience of measurements of A β concentrations. Researchers were blinded to sample identity where possible. Figure legends contain sample sizes, replicate information, and statistical tests used.

Statistics

Statistical analyses were performed using GraphPad Prism (version 6 and 7). Differences between two groups were evaluated for significance with two-tailed Student's *t* test and multiple *t* test when comparing two treatments. Comparisons of three or more groups on a single dataset were performed by one-way ANOVA, followed by Tukey's post hoc test. A *p* value threshold of 0.05 was used for assessment of the statistical significance. Values are shown as means \pm SD. Individual subject-level data are reported in data file S1 and data file S2.

List of supplementary materials

Material and Methods

Fig. S1. *Immunohistochemistry of different regions of the Uppsala APP mutation brain with A β 40, A β 42 and GFAP antibodies.*

Fig. S2. *A β 42 in CSF measured by two different methods.*

Fig. S3. *Sequence verification of A β peptides.*

Fig. S4. *6E10 and 1C3 binding epitopes.*

Fig. S5. *CTFs fragments in cell lysate from HEK293 cells transfected with APPwt and APPUpp.*

Fig. S6. *Fragment ion spectra of selected APP derived peptides from Figure 7.*

Fig. S7. *Comparison of the most common A β 1-42 Δ 19-24 polymorphs with existing A β fibril structures.*

Fig. S8. *Fourier shell correlation (FSC) of the most common A β 1-42 Δ 19-24 polymorphs.*

Table S1. *CSF samples from the Uppsala BioBank (UBB), patient information.*

Table S2. *Brain samples from the Uppsala BioBank (UBB) and Netherlands Brain Bank (NBB), patient information.*

Table S3. *Assignment of observed peaks in human brain tissue using MALDI imaging MS and LC-MS/MS based verification from extracts of individually laser microdissected plaques.*

Data file S1. *Individual-level data figures 4-8.*

Data file S2. *Individual-level data supplementary figures.*

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Acknowledgments

We thank Christine Röder and Lothar Gremer for help with sample preparation for cryo-EM, discussions and advice. We would also like to thank Per Hammarström and Peter Nilsson for providing fluorophores q-FTAA and h-FTAA. We also thank the Uppsala Genome Center for the help with targeted exome sequencing, Linn Streubel-Gallasch for statistical advice and Joel Watts for critical proof-reading of the manuscript. Finally, figure 2 was made in part using Biorender. **Funding:** This work was supported by the Swedish Research Council (#2016-02120 LL, #2018-02181 JH); the Swedish Alzheimer Foundation (LL, JH); the Swedish Brain Foundation (LL); Åke Wibergs Stiftelse (JH); Åhlén Stiftelsen (JH); the German Research Foundation within the framework of the Munich Cluster for Systems Neurology (EXC 2145 SyNergy, project ID 390857198) and the research unit FOR2290, the BMBF within project CLINSPECT-M and by JPco-fuND2 2019 Personalized Medicine for Neurodegenerative

Diseases 01ED2002B. **Author Contributions:** MP, VG, LL, DS and MI designed the study. LL obtained the funding. These authors performed the experiments: MP (genetic analyses, immunostainings, ELISAs, ThS staining, cell transfection, Western blot, Mesoscale electrochemiluminescence, Thioflavin T assay), VG (genetic analyses, plasmid designs) WM (MALDI imaging, mass spectrometry and LCO stainings), GG (cell transfection, immunoblotting with protease inhibitors and CTF fragments), MZ (Transmission Electron Microscopy), LS (Size Exclusion Chromatography) and SAM (Mass Spectrometry for APP^{Upp} cleavage site determination of transfected cell culture). MP, VG, WM, TD, LS, IA, LNGN, AE, DW, GFS, JH, SFL, LL, DS and MI analysed the data. LK, RMB, ML and MI contributed to sample collection. MP, VG, DS and MI wrote the first draft of the paper. All authors contributed to the final version of the paper. **Competing Interests:** Dr. Nilsson has received an honorarium from BioArctic AB, and has a research collaboration with this company, outside the submitted work. Dr. Lannfelt is a co-founder of BioArctic AB. Dr. Ingelsson is a paid consultant for BioArctic AB. **Data and materials availability:** All data associated with this study are present in the paper or the Supplementary Materials. Materials in this study will be made available by contacting the corresponding author and completion of a material transfer agreement. Exon 17 nucleotide sequence for Uppsala mutation submitted to GenBank (MW892394). Density maps of the A β ^{Upp1-42 Δ 19-24} obtained by cryo-EM were deposited in the EMDB for polymorph 1 (EMD-12592) and for polymorph 2 (EMD-12593).

Figures

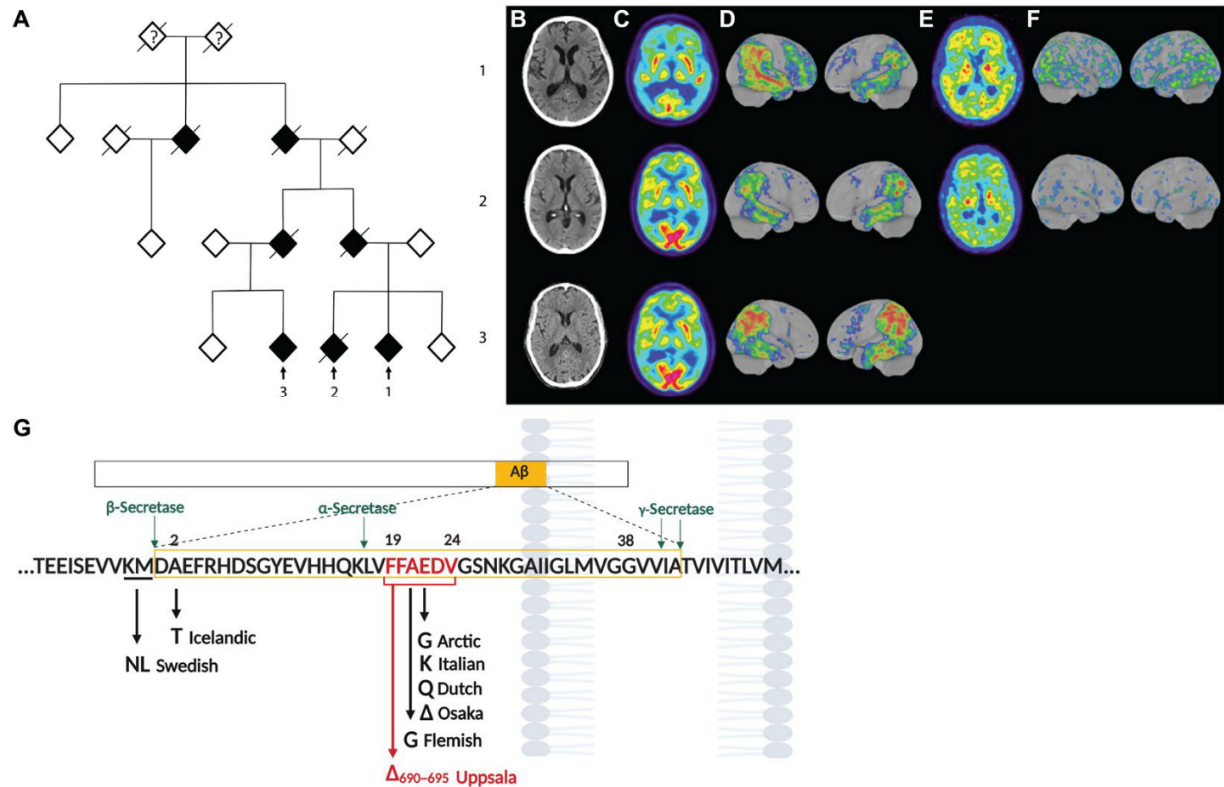


Fig. 1: Epidemiological and clinical features of the *Uppsala APP* mutation family. Pedigree of the *Uppsala APP* mutation family (A). Filled symbols are affected family members. Slashed symbols are deceased individuals. Index cases are indicated by arrows. Several healthy individuals in the latest generation have been omitted on purpose. CT and PET images of the three affected patients (sibling 1, sibling 2 and cousin 3, from upper to lower rows). Axial CT (B), FDG-PET (C-D) and PIB-PET images (E-F). 3D-surface projection showing areas with pathological cortical tracer uptake (D) and (F). For (C) and (D) the color scale represents decreased tracer uptake with Z-score between 0 (blue) and -7 (red). For (E) and (F) the color scale represents increased tracer uptake with Z-scores between 0 (blue) and 8 (red). The location of the *Uppsala APP* mutation and its relation to other intra-Aβ *APP* mutations (G).

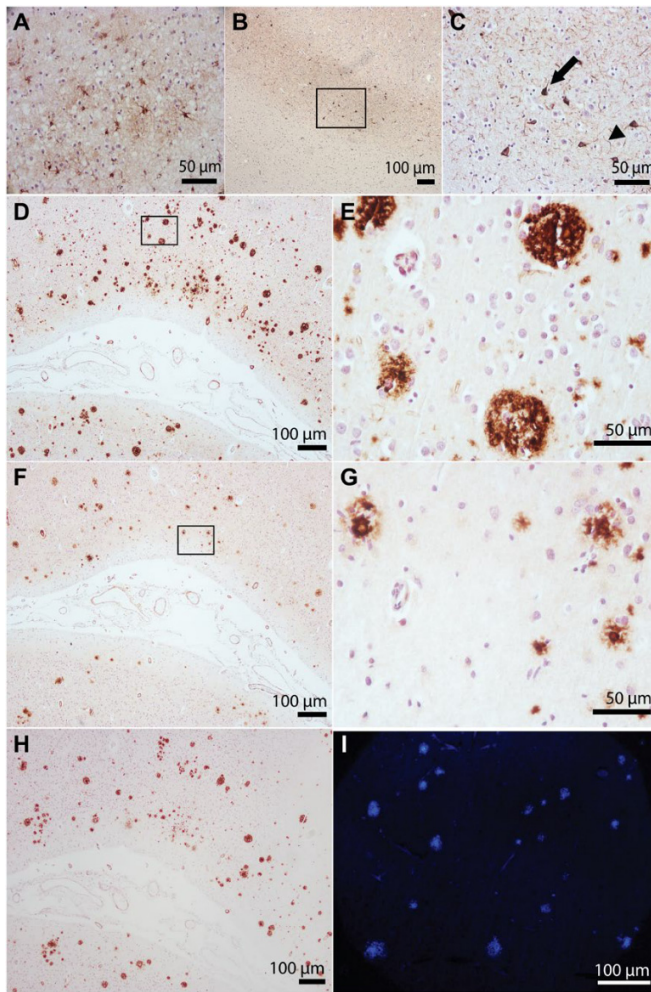


Fig. 2: Pronounced A β and tau pathology in the *Uppsala APP* mutation carrier brain.

Immunohistochemistry of tissue sections from the *Uppsala APP* mutation brain, against GFAP (anti-GFAP) (A), tau (AT8) (B-C), total A β (6F/3D) (D), A β 42 (anti-A β 42) (E-F) and A β 40 (G-H). Staining of amyloid plaques with Thioflavin S (I). A-D: temporal cortex, E-I: occipital cortex. Squares in B, E and G: regions displayed in the higher magnification images C, F and H. Arrow in C points to a tangle, arrowhead to a dystrophic neurite.

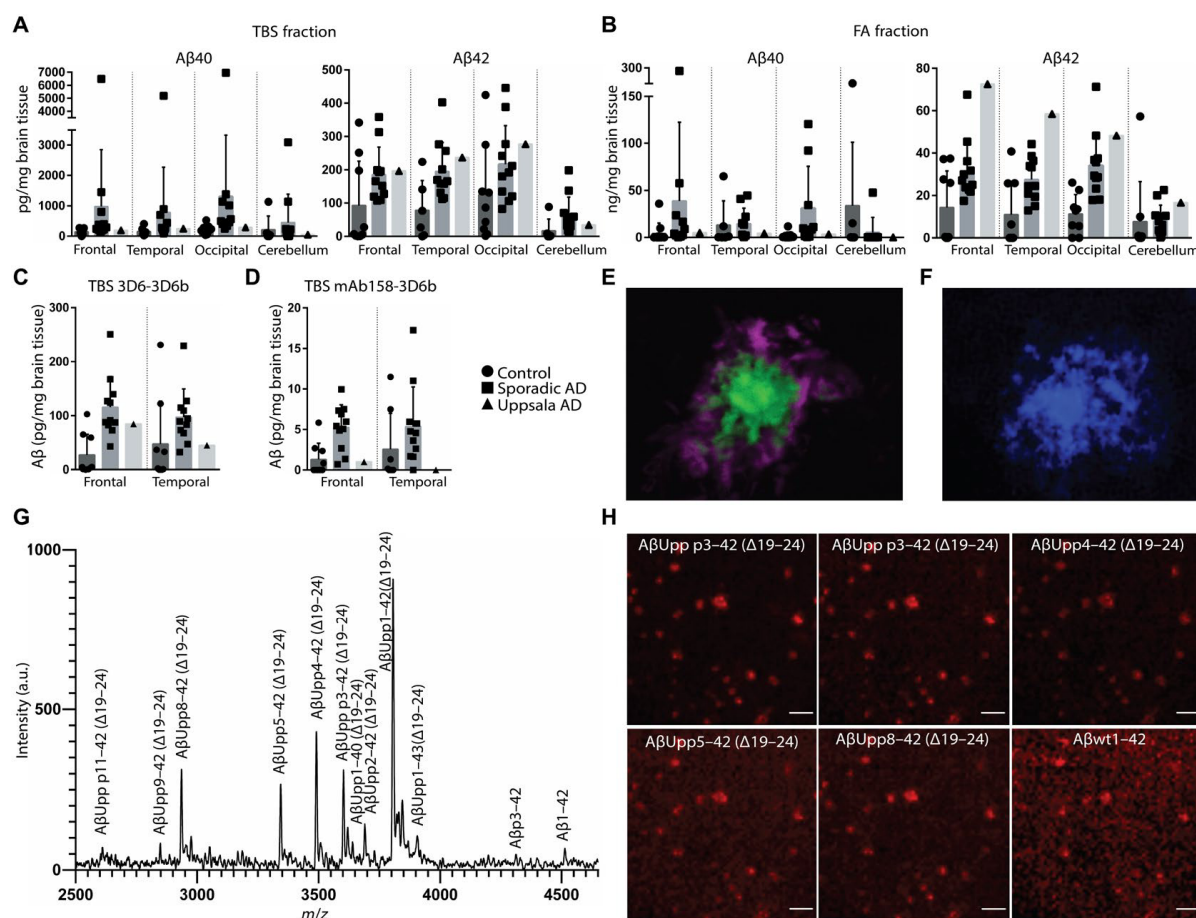


Fig. 3: Plaques in the Uppsala APP mutation brain mainly consist of AβUpp42_{Δ19-24}

Electrochemiluminescence (MSD) immunoassay analyses of human brain extracts (A-D). Concentrations of Aβ42 in the TBS (A) and FA (B) fractions. Concentrations of TBS soluble Aβ aggregates (C) and large Aβ oligomers (D). Staining of Aβ plaques on fresh frozen temporal neocortical tissue of the Uppsala APP mutation brain with the luminescent conjugated oligothiophenes (LCOs) q-FTAA (green) and h-FTAA (purple) (E) and with ThS (blue) (F). MALDI-IMS spectrum (G) and images of plaques from the Uppsala APP mutation brain (H). Error bars represent standard deviation (SD). For controls $n=9$ and for sAD $n=11$. Due to $n=1$ for the Uppsala AD group, no statistical analysis was performed.

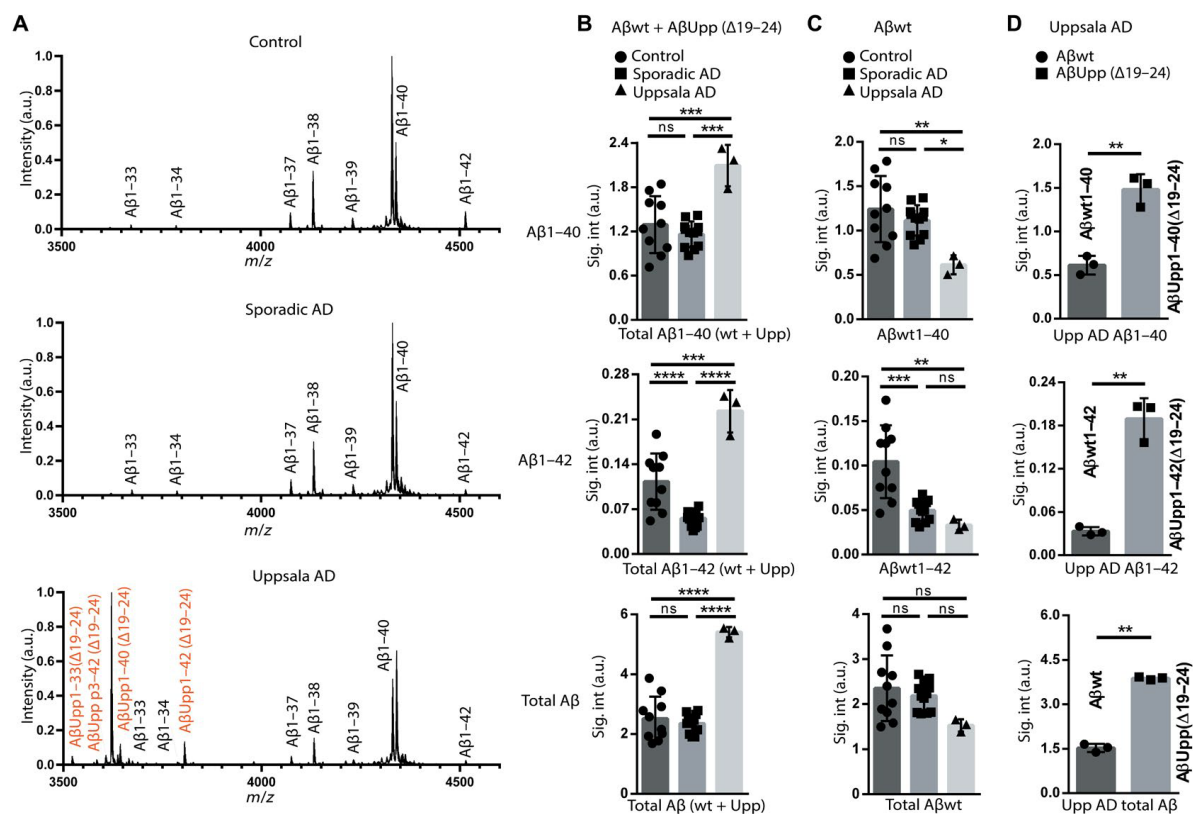
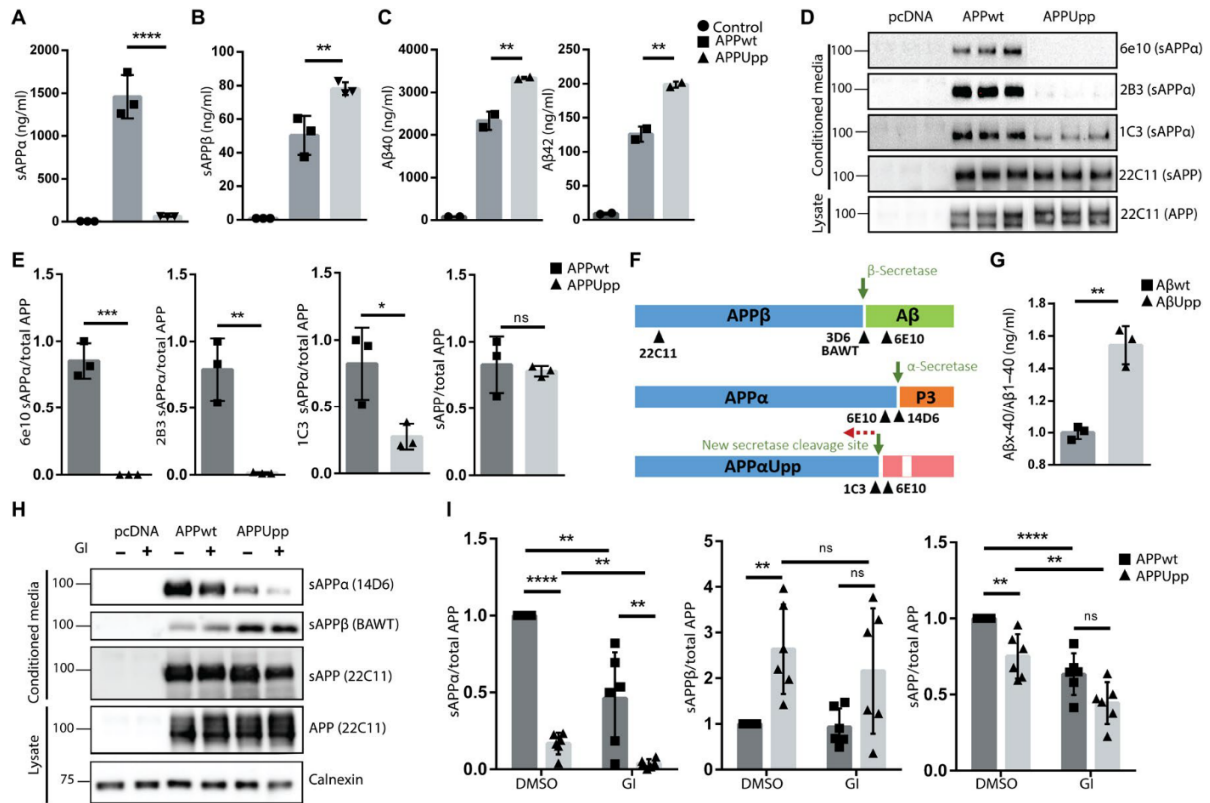


Fig. 4: Uppsala APP mutation carriers present increased concentration of Aβ in CSF. IP-MS analyses of CSF from ten healthy controls, eleven sporadic AD cases and the three *Uppsala* APP mutation patients (A). IP-MS based quantitation of CSF Aβ1-40, Aβ1-42 and total Aβ concentrations in CSF (B-D). All data are represented as group means and error bars represent standard deviation (SD). The arbitrary units in the y-axes represent ratios of intact mass peak area of individual Aβ peptide signals normalized to the spiked Aβ1-40 internal standard. For control CSF $n=10$, for sAD $n=11$, for Uppsala CSF $n=3$. Analyses of individual peptide signals and comparisons between the groups were performed by one-way ANOVA in B) ($P=0.0002$, $P<0.0001$ and $P<0.0001$, for Aβ1-40, Aβ1-42 and total Aβ respectively) and C) ($P=0.0069$, $P=0.0001$ and $P=0.0767$ for Aβ1-40, Aβ1-42 and total Aβ respectively) followed by Tukey's post hoc test. For D) paired t-test was performed ($P=0.0029$, $P=0.0081$, $P=0.0007$; for Aβ1-40, Aβ1-42 and total Aβ respectively). Non-significant (ns) * $P<0.05$, ** $P<0.01$, *** $P<0.001$, **** $P<0.0001$.



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933 **Fig. 5: The Uppsala APP mutation alters APP processing**

934 Electrochemiluminescence (MSD) immunoassay analyses of sAPPα (A), sAPPβ (B) and Aβ40
 935 and Aβ42 (C) in conditioned media from HEK293 cells transfected with *APPUpp* as compared
 936 to *APPwt* ($n=3$, $N=1$). Western blot of *APPUpp* conditioned media, with the sAPPα antibody 2B3
 937 (C-terminal), 6E10 (Aβ5-10), mAb1C3 (Aβ3-8) ($n=3$, $N=3$) (D). Results from (D) quantified as
 938 a ratio of sAPPα (detected with 6E10, 2B3 or 1C3) or total sAPP (detected with 22C11) in cell
 939 medium over total APP in cell lysate (E). Schematic image indicating β- and α-cleavage sites of
 940 *APPwt* and *APPUpp*, with antibody binding epitopes indicated (F). Ratio of Aβx-40 and Aβ1-40
 941 in *APPwt* and *APPUpp* medium quantified by ELISA ($n=3$, $N=3$) (G). Western blot analyses of
 942 media and lysates from HEK293 cells transiently transfected with *APPwt* or *APPUpp*, with or
 943 without the ADAM10-preferring inhibitor GI254023X (GI) using specific antibodies for α- and
 944 β-APP ($n=6$, $N=4$) (H). Results from (H) quantified as a ratio of sAPPα, sAPPβ or total sAPP

945 in cell medium over total APP in cell lysate (I). Statistical significance was determined by one-
946 way ANOVA (A-C) (for A) and B) $P<0.0001$; for C) A β 40 and A β 42 $P=0.0003$) followed by
947 Tukey's post hoc test; two tailed unpaired t test ($P= 0.004, 0.0045, 0.0291, 0.7167$) (E) and
948 ($P=0.0016$) (G); and multiple t-test (H). All data are represented as group means and error bars
949 represent standard deviation (SD). Non-significant (*ns*), $*P<0.05$, $**P<0.01$, $***P<0.001$,
950 $****P<0.0001$. All results were normalized to total APP.

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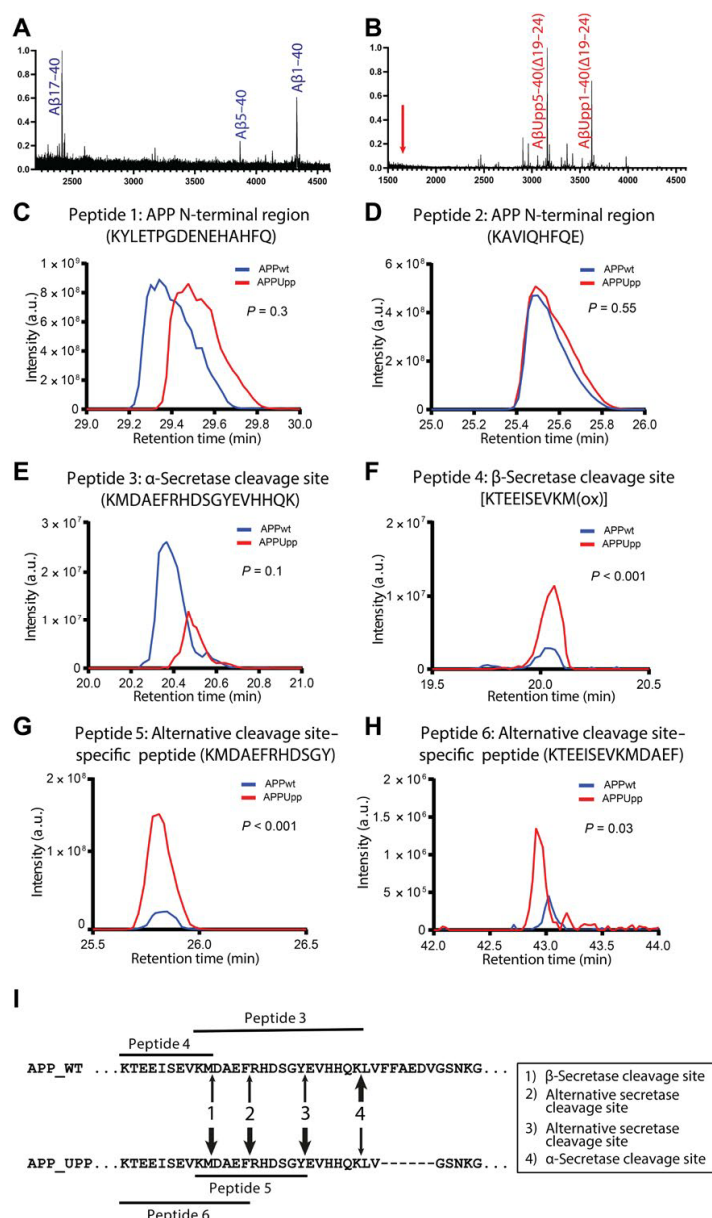


Fig. 6: The *Uppsala* APP mutation increases β-secretase cleavage and alters α-secretase cleavage

MS spectra of the *APPwt* (A) and *APPUpp* (B) transfected HEK293 cells that showed the most prominent peptides present in cell media from cells transfected with either *APPUpp* or *APPwt*. The absence of Upp17-40Δ19-24 is indicated by the red arrow (B). Extracted ion chromatograms of different peptides of APP (blue: *APPwt*, red: *APPUpp*). Chromatogram of the N-terminal APP peptides, Peptide 1, KYLETPGDENEHAHFQ (302-317 in wt hAPP695) (C) and Peptide 2,

960 KAVIQHFQE (354-362 in wt hAPP695) (D). Shifts in retention times between *APP_{wt}* and
961 *APP_{Upp}* were within the normal range of shifts between runs. Chromatogram of the α -secretase
962 cleavage site-specific peptide 3, KMDAEFRHDSGYEVHHQK (595-612 in wt hAPP695,
963 containing one missed LysN cleavage site) (E). Chromatogram of the β -cleavage site-specific
964 peptide 4, KTEEISEVKM(ox) (587-596 in wt hAPP695, also containing one missed LysN
965 cleavage site) (F). Chromatogram of the semi-specific peptides (N-terminus specific for LysN,
966 C-terminus unspecific) peptide 5, KMDAEFRHDSGY (595-606 in wt hAPP695) (G) and peptide
967 6, KTEEISEVKMDAEF (587-600 in wt hAPP695) (H). (I) Cleavage sites of α - and β -secretase
968 are indicated for the sequences of *APP_{wt}* and *APP_{Upp}*. Thick arrows indicate increased and thin
969 arrows indicate decreased cleavage of the two *APP* sequences.

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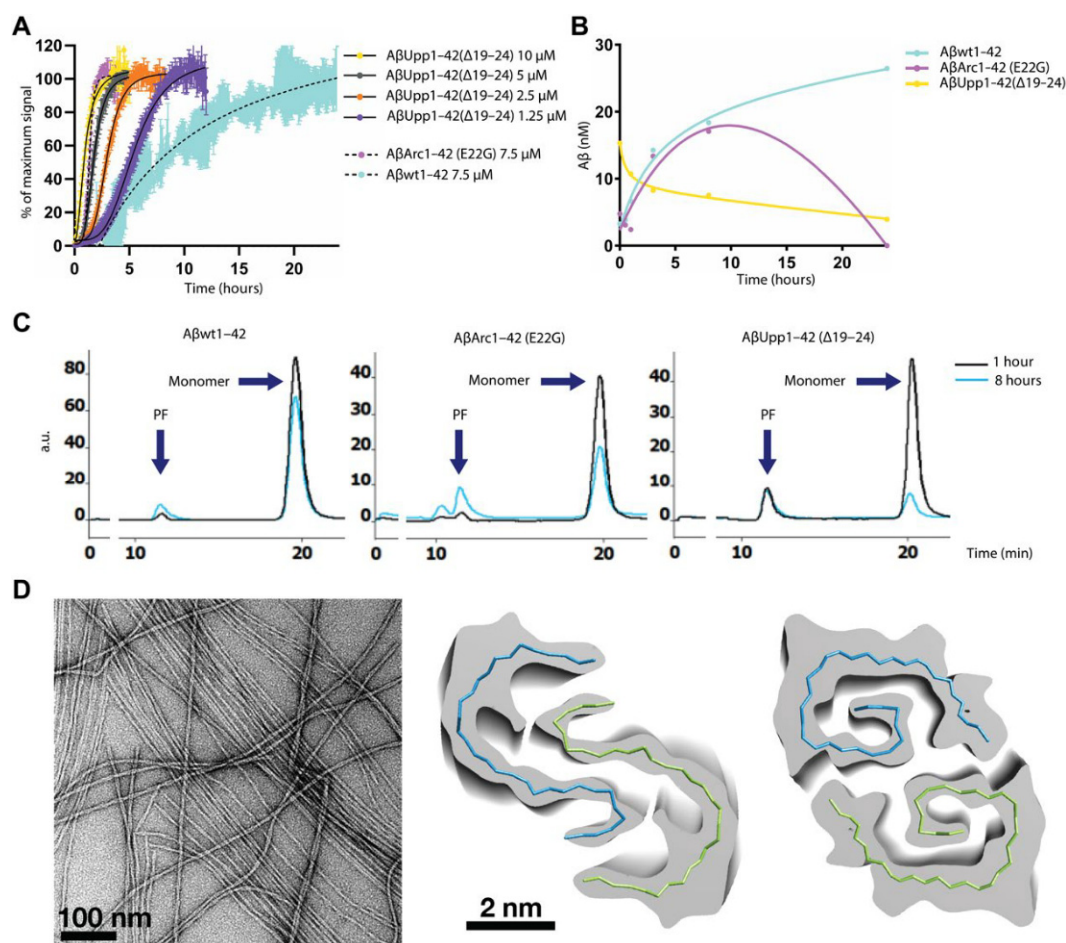


Fig. 7: The Uppsala APP mutation accelerates the formation of fibrils

ThT assay of different AβUpp1-42 Δ 19-24, AβArc1-42 $E22G$ and Aβwt1-42 (A). Concentrations of soluble Aβ oligomers/protofibrils for incubated AβUpp1-42 Δ 19-24, AβArc1-42 $E22G$ and Aβwt1-42 samples, as measured by ELISA (B) and size exclusion chromatography (C). Negative stain EM image of AβUpp1-42 Δ 19-24 fibrils formed in low pH resulted in long, well-ordered fibrils (D, left). Cryo-EM reconstructions (D, middle and right). For ThT, 4 replicates of each peptide were aggregating simultaneously ($n=4$) for each of the three experiments ($N=3$). Error bars represent standard deviation (SD) of the replicates and black and dashed lines represent curves fitted to the ThT data points. For ELISA, from the same monomeric fraction used for ThT, 2 replicates ($n=2$) for each experiment ($N=3$). Size exclusion chromatography was performed one time ($N=1$).