Expression, purification, and preliminary characterization of human presenilin-2

Ge Yang\textsuperscript{a,b,1}, Kun Yu\textsuperscript{a,b,1}, Jan Kubicek\textsuperscript{b}, Jörg Labahn\textsuperscript{a,b,*}

\textsuperscript{a} Centre for Structural Systems Biology (CSSB), CSSB-FZJ, Notkestr. 85, 22607 Hamburg, Germany
\textsuperscript{b} Institute of Complex Systems-Structural Biochemistry (ICS-6), Forschungszentrum Jülich, Wilhelm-Johnen-Str., 52425 Jülich, Germany
\textsuperscript{c} Cube Biotech GmbH, Alfred-Nobel-Str. 10, 40789 Monheim, Germany

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ABSTRACT

Presenilins (PS1 and PS2) exhibit similar γ-secretase-dependent and –independent functions with subtle variations. In this study, we established a cost-effective process to overexpress and purify full-length human PS2 in sufficient quantities and quality for structural studies. Upon optimization, milligram quantities of homogeneous trimeric hisPS2 were purified, which enabled the preliminary characterization of human hisPS2 zymogen. Far-UV and near-UV CD as well as fluorescence spectroscopy revealed that purified hisPS2 contained the expected secondary structure and was folded into a defined tertiary structure. Thermal stability analysis revealed a T\textsubscript{m} value of \textasciitilde55 °C for secondary structure while cholesterol significantly increased the stability. The low melting temperature of \textasciitilde34 °C for the tertiary structure was able to explain the purity and aggregation problems observed during purification. Additionally, the occurrence of calcium ions induced structural changes to different extents for PS2WT and PS2-D263A/D366A was observed, which is consistent with previous studies.

1. Introduction

Full-length presenilin zymogens (PS1 and PS2) are short-lived multi-transmembrane proteins found in the endoplasmic reticulum (ER) \cite{1}. During ER-to-Golgi recycling \cite{2}, PS zymogens are activated by autocleavage into stable N- and C-terminal fragments that associate with three other components – nicastrin, Aph1, and Pen-2–to form the γ-secretase complex \cite{3,4}. In contrast to endogenous PS1, PS2 was mainly found as the full-length protein in human and mouse brains \cite{5,6}. During embryonic development \cite{7}, neuron maturation \cite{6}, and aging \cite{8}, PS1 expression was either constant or decreased while PS2 expression increased, thus indicating their distinct regulatory functions.

Due to the existence of two PSs and – in humans – two APH1 isoforms, at least four complexes co-exist in the cell \cite{9,10}, which process distinct substrates depending on their subcellular localization \cite{9}. The PS1-containing complexes reside predominantly at the recycling endosomes and the plasma membrane \cite{11,12}, while PS2-containing complexes are restricted to late endosomes and lysosomes \cite{13}. PSs exhibit similar but subtly distinctive biological functions in γ-secretase-dependent and –independent pathways \cite{14}.

The most widely studied γ-secretase substrates are Notch and C-terminal fragments of β-amyloid precursor protein (APP-CIFs). The cleavage of Notch by γ-secretase releases the intracellular domain to the nucleus, which in turn affects the development of cells and cancer \cite{15}. The cleavage of APP-C99 by γ-secretase generates amyloid-beta peptides (Aβ) of varying lengths, among which the longer peptides – particularly Aβ42 and Aβ43–are aggregation-prone and are the initial and key components of Aβ plaques \cite{16}, a hallmark of Alzheimer’s disease (AD). According to the amyloid hypothesis of AD, mutations in APP or PSs induce the relatively elevated production of Aβ42 peptides. The abnormal deposition of Aβ42 was proposed to trigger the AD pathogenesis \cite{17,18}. Of all the PS-related AD pathogenic mutations, about 150 are found in PS1 and about 13 in PS2 \cite{19}. PS1-containing γ-secretase was reported to produce more Aβ than PS2-containing γ-secretase \cite{20} and is responsible for the production of the majority of CNS Aβ \cite{21}. However, familial AD (FAD) mutations in PS2 increase the production of intracellular toxic Aβ42 in late endosomes and lysosomes. Some FAD-PS1 mutations lead to a change in the subcellular location of γ-secretase and phenocopy the FAD-PS2 mutations \cite{13}. The amyloid hypothesis is challenged by the heterogeneous clinical phenotypes of PS mutations and is yet to be experimentally proven \cite{22–24}. The presenilin hypothesis of AD proposes that the loss of
essential functions of PSs alone might be a more potent cause of AD-related phenotypes than the accumulation of toxic Aβ peptides [25,26]. It is still debated as to whether PS mutations induce AD through gaining a toxic function of γ-secretase or through the loss of an essential function of PSs.

PSs are also proposed to be involved in many other biological processes independent of γ-secretase, including the modulation of intracellular calcium homeostasis although the exact site(s) and mechanism of this function requires further investigation. It was suggested that PS holoproteins form ER channels [27,28], but contradicting results have been reported [29–31]. This calcium leakage function is independent of γ-secretase activity: PS1-D257A, which lacks proteolytic activity and therefore remained uncleaved, still exhibited the channel function [28]. PSs physically interact with ER calcium channels (e.g. InsP3R [29,32] and RyR [33,34]) and up-regulate calcium release. Furthermore, PSs affect the calcium influx by interacting with the ER SERCA pump [35,36]. For PS2, it is only the full length protein and not the cleaved fragments that exhibits depleting effects on the intracellular calcium stores [36]. The preferential interaction of PS2 with calcium binding proteins (e.g. calmyrin [37], μ-calpain [38], and sorcin [39]) were also observed. PS2– but not PS1—was also reported to modulate ER-mitochondria tethering and calcium cross-talk between these two organelles [40,41]. PSs are also essential in regulating mitochondrial calcium homeostasis [42]. Moreover, PSs also alter lysosomal calcium release/storage [43,44]. Many familial AD mutations in PSs cause disturbed calcium signaling, which may indicate an important role for abnormal calcium signaling in AD pathogenesis [45–47].

The PS-related structural information reported were the crystal structure of the presenilin archaeal bacterial homolog (PSH) [48] and the nuclear magnetic resonance (NMR) structures of the C-terminal domains of PS1 (PS1-CTD) [49] and the cryo-electron microscopy (cryo-EM) structures of PS1-containing γ-secretase [50–52]. These analyses are all based on activated PS1. The low expression level and poor solution behavior of full-length eukaryotic PSs [48] directed research towards bacterial homologues. Structural information on the PS2 zymogen alone may help to elucidate the different physiological and pathological functions of PS1 and PS2, and might also shed light on the auto-inhibition and activation mechanisms for both the proteolytic activity of γ-secretase and the suggested passive calcium leakage channel.

The present study reports on the detailed process for large-scale expression, purification, and preliminary structural characterization of human PS2 by CD spectroscopy. This expression and purification approach allows milligram quantities of well-folded, homogeneous full-length PS2 zymogens to be obtained for further structural analysis.

2. Materials and methods

2.1. Materials

The E. coli strain BL21-CodonPlus (DE3)-RP was obtained from Agilent Technologies (Hamburg, Germany), BL21 Star™ (DE3) from Invitrogen (Darmstadt, Germany), C41(DE3) and C43(DE3) from Lucigen (Heidelberg, Germany), detergents and cholesterol from Anatrace (Maumee, USA) and Glycon (Luckenwalde, Germany), Ni-NTA resin from Qiagen (Hilden, Germany), and the EDTA-free protease inhibitor cocktail from Roche (Basel, Switzerland). Furthermore, the HiLoad 16/60 Superdex 200 pg column was obtained from GE Healthcare (Freiburg, Germany). All other reagents were purchased from either Sigma-Aldrich or Merck.

2.2. Plasmids construction

A codon usage-optimized human PS2 gene was synthesized by GENART (Regensburg, Germany) and cloned into the pQE2 vector (Qiagen, Hilden, Germany) with an N-terminal hexa-histidine (hisPS2).

PS2 active site mutations (hisPS2-D263A/D366A) were performed using the QuikChange Site-Directed Mutagenesis Kit (Agilent Technologies, Hamburg, Germany). The correctness of resulting plasmids was confirmed by sequencing.

Initially, the possibility of expressing and purifying the entire PS2 in E.coli was evaluated using a PS2 construct with N-terminal hexa-histidine and a C-terminal strep tag (hisPS2strep), which allowed protein integrity to be monitored by western blot.

2.3. Expression and membrane preparation

An overnight pre-culture (supplemented with 50 µg/ml kanamycin and 2% (w/v) glucose) was diluted into 300 ml Terrific Broth (TB) or Double Yeast Tryptone (DYT) media in a 2 l flask to an OD600 of ~0.2 and grown at 30 °C to an OD600 of ~0.8 for induction with 0.2 mM IPTG. Cells were then grown at 18 °C for 16 h before being pelleted. The removal of the periplasmic fraction was performed after harvesting [53].

The obtained pellet was resuspended with 10 ml ice-cold lysis buffer (20 mM Tris HCl; pH 8.0; 10% (v/v) Glycerol; Protease inhibitor 1 tablet/50 ml buffer; 1 mM PMSF; 1 mg/ml Lysozyme; DNAlase 5 mg/50 g cell pellet; 1 mM TCEP) per gram pellet and incubated at 4 °C for 1 h under gentle stirring. The cell suspension was passed through a cell disruptor (EmulsiFlexC3, Avestin, Mannheim, Germany) five times at a pressure of ~20, 000 psi. After lysis, EDTA (10 mM) and NaCl (300 mM) were added to the lysate. This suspension was subjected to three steps of sequential centrifugations at 4 °C: 900g (15 min), 10, 000 × g (30 min), and 100, 000 × g (1 h). The pellet fraction from the second (inclusion body fraction) and third centrifugation step (membrane fraction) were analyzed by western blot.

2.4. Solubilization screening

Membrane from 0.1 g cell pellet was suspended in 0.5 ml solubilization buffer (20 mM Tris HCl; pH 8.0; 10% (v/v) Glycerol; 300 mM NaCl; 1 mM PMSF; Protease inhibitor 1 tablet/50 ml buffer; 1 mM TCEP; 1–2% (w/v) detergent). The detergents used in the study are indicated in Table 1 in Ref. [54]. After solubilization, the supernatant and pellet were separated by centrifugation at 100,000g for 1 h and analyzed by western blot.

2.5. Ni-NTA purification

Membrane from 20 g cell was solubilized in 100 ml solubilization buffer (1% (w/v) Fos-choline-14 (FC14)). After centrifugation, imidazole was added to the supernatant to 15 mM. This solution was loaded onto a 5 ml pre-equilibrated Ni-NTA column. The flow-through was collected and reapplied to the column twice. Five column volumes (CV) of the solubilization buffer supplemented with 15 mM imidazole (wash 1) were then passed through the column. After this step, the detergent concentration was decreased to 0.014% (w/v) FC14. Other components of the buffer were kept constant except for changes in salt and imidazole concentrations: wash 2 (500 mM NaCl, 5 CV), wash 3 (1 M NaCl, 5 CV), wash 4 (20 mM imidazole), and wash 5 (25 mM imidazole) were carried out sequentially. Elution of the bound protein was achieved by the buffer being supplemented with 300 mM imidazole. For solubilization and purification of hisPS2 in mixed micelles of FC14 and cholesterol, cholesteryl hemisuccinate tris salt (CHS) was added to the corresponding buffers (CHS: FC14 = 1:10 (w/w)).

2.6. Mass spectroscopy (MS)

Bands of interest were cut from SDS-PAGE, desaturated, dehydrated, and digested as described in the Trypsin Profile IGD Kit (Sigma-Aldrich, Hamburg, Germany). 0.1% TFA was added to the extraction for acidification prior to matrix addition. The digested peptides were then
analyzed by matrix-assisted laser desorption/ionization time-of-flight (MALDI-TOF) and MS/MS using a Bruker Daltonics Ultraspec III TOF/TOF mass spectrometer (Bruker Daltonics, Billerica, USA). Baseline correction and mass peak detection were performed using the software flex Analysis Version 3.0 (Build 92, Bruker Daltonics, Billerica, USA). Finally, both the MS and tandem MS data were searched against Swiss-Prot protein databases for protein identification.

2.7. Size exclusion chromatography (SEC)

2 ml of concentrated protein sample (30 KDa Millipore concentrator, Millipore, Darmstadt, Germany) was filtrated (0.2 µM centrifugation filter, Nanospec®, Dreieich, Germany) and subjected to SEC on a superdex 200 pg column calibrated by a SEC calibration kit (Sigma-Aldrich, Hamburg, Germany). The flow rate used was 0.3 ml/min. Detergent micelles were visualized by staining FC14 micelles by rhodamine B (Avanti Polar Lipids, Hamburg, Germany) at a monomer mole ratio of ~20:1. The apparent molecular weight and radius of the eluted protein–detergent complex were obtained from the calibration curve.

2.8. Circular dichroism (CD) and fluorescence spectroscopy

The CD and fluorescence spectra were recorded using an Aviv 425 circular dichroism spectrometer (Aviv Biomedical, Lakewood, USA) equipped with fluorescence emission scanning monochromator which enables the simultaneous collection of CD and fluorescence data. Purified hisPS2 was buffer exchanged by a PD-10 desalting column (GE Healthcare, Freiburg, Germany) into CD buffer (10 mM sodium phosphate, pH 7.4, 0.014% (w/v) FC14) for data collection. For far-UV CD, spectra between 260 nm and 185 nm were collected with 1 nm step size, 1 nm bandwidth, and 9 s averaging time in 0.1 cm path length Suprasil cuvettes. For near-UV CD, data from 350 nm to 250 nm were obtained with 0.25 nm step size, 1 nm bandwidth, and 30 s averaging time in 1 cm path length cuvettes. Baselines were measured from buffer solutions under the same experimental conditions. Baseline subtraction and spectra smoothing were carried out using the Aviv CDS software. All spectra were recorded in triplicate and collected at 4 °C except the thermal unfolding experiments. The protein concentrations used for far- and near-UV CD were ~3.3 µM and ~20.8 µM, respectively. Protein concentrations were determined from the absorbance at 280 nm using the calculated extinction coefficient 76780 M⁻¹ cm⁻¹ obtained by ExPASy ProtParam [55]. The baseline-subtracted spectra were scaled to obtain the mean residue ellipticy (MRE) using a mean residue weight of 112.54 Da. The CDSSTR algorithm and the reference data sets 4, 7, and SMP 180 were used for the deconvolution of spectra [56, 57].

During thermal unfolding, the temperature was raised in 2 °C steps with a heating rate of 0.5 °C/min and 1 min equilibration time before data collection. For far-UV CD, the signal at 221 nm was collected with an averaging time of 9 s between 4 °C and 98 °C. A temperature reverse scan was carried out in order to investigate the reversibility. For near-UV CD, the 292 nm signals were recorded with an averaging time of 30 s from 4 °C to 80 °C. The measured signals were corrected for buffer contribution and fitted with a Boltzmann sigmoidal equation (Origin 9.0) to obtain the melting temperature (Tm).

The fluorescence emission spectra (450 nm to 270 nm) were recorded at different excitation wavelengths with a bandwidth of 2 nm, a photomultiplier tube (PMT) voltage of 900, an averaging time of 1 s, and an emission slit setting of 2 nm.

3. Results

3.1. Overexpression of hisPS2

Heterologous overexpression of human membrane proteins often result in cell death, protein misfolding, and aggregation due to the saturation of the host sorting and translocation system and cell membranes [58, 59]. We therefore applied codon usage-optimized DNA and screened four different strains in DYT and TB media for their ability to overexpress human PS2.

As listed in Fig. 1a, hisPS2 exhibited toxicity to STAR and RP strains, resulting in cell death (OD600 lower than induction) after 16 h expression when the DYT medium was used – but not for the TB medium. This is probably due to the acidic stress upon overexpression [59]. For the DYT medium, an acidic pH was observed upon cell harvesting. Regardless of the medium used, C41 and C43 strains yielded relatively higher cell densities than the other two strains. These strains contain mutations in the lacUV5 promoter which slows down the speed of protein translation [60]. Meanwhile, a lower temperature (18 °C) after induction was used to further slow down the protein expression rate, which allows the translation system to optimally adapt to induction stress [61]. It was revealed by means of Western blot (Fig. 1b) on the N-terminal his-tag that most protein was present in the inclusion body fraction when the DYT medium was used. For the TB medium, an equal distribution of the protein between the inclusion bodies and the membrane fraction was found. In order to obtain the correctly folded protein, only the isolated membrane fraction was used for extraction of hisPS2. Large-scale overexpression of hisPS2 was achieved in the C43 (DE3) strain with the TB medium. This combination yielded the highest cell density and cell pellet weight. Typically, after 16 h of expression, an OD 600 of 8.3 ± 0.3 and a cell pellet wet weight of 12.9 ± 0.3 g/L was obtained.

3.2. Fos-choline detergents are most efficient in solubilizing hisPS2

After overexpression, we investigated the ability of 13 detergents with different soluble groups, charge, and chain length to solubilize hisPS2 (Table 1 in Ref [54]). After detergent extraction (Fig 2a), clear, transparent residual pellets were observed for FC12, FC14, FC16, and NLS. The other detergents exhibited pellets that were brown in color. Western blots (Fig. 2b) revealed that the signal in the supernatant (SN)
after centrifugation was comparable to the total signal found for the suspension (S) before centrifugation for FC12, FC14, FC16, and NLS. No protein signal was detected in the pellets for these four solubilizers. Other protein investigate solubilized hisPS2 either poorly or not at all. It was generally observed that the less brown the color of the pellet after detergent extraction, the more efficient the solubilization was. The common denominator for complete solubilization was detergents with an aliphatic chain length of at least 12 carbons and the presence of negative charge at the head group.

3.3. Purification of hisPS2 from membrane fraction by Ni-NTA and SEC

The small-scale purification of hisPS2 by Ni-NTA or a combination of Ni-NTA and Strep-Tactin purification for hisPS2strep was performed in the presence of different detergents (data not shown). It was observed that Ni-NTA binds hisPS2 more tightly than Strep-Tactin. Ni-NTA resin was therefore used for preparing the setup of purification. FC14 was chosen as the purification detergent. Different amounts of FC14 micelles and Ni-NTA matrix were used with a constant amount of cell pellets to investigate the yield and purity (data not shown). 5 mL 1% (w/v) FC14 and 0.25 mL Ni-NTA matrix per gram of cell pellet gave the best result. SDS-PAGE exhibited two dominant bands of PS2 and impurities in the elution fractions (Fig. 3a). The two dominant bands were found to contain PS2 by MS based on five identified peptides (Fig. 1 in Ref [54]) with 14% sequence coverage (Fig. 3b). All the peptides detected belong to extra-membrane segments (Fig. 2 in Ref [54]), which is similar to the MS result reported for PS1 [62]. The low sequence coverage in MS is probably a result of the inaccessibility of the cleavage sites due to bound detergent. Despite repeated efforts to improve the sequence coverage, only a segment of TM7 was detected (data not shown). ImageJ was used to quantify the intensities of the Coomassie-stained bands. The purity after Ni-NTA was calculated at ~ 81% (Fig. 3a).

After Ni-NTA chromatography, minor impurities and aggregated hisPS2 were removed by SEC. HisPS2 was distributed into two major peaks with a shoulder on the left side of the first peak (Fig. 4a). The pooled fraction of the first peak not only contained hisPS2 but also an E.coli protein of ~75 kDa, as detected by SDS-PAGE (Fig. 4a, inset, lane1). The purity of the pooled fractions from the second peak was estimated at ~95% via ImageJ by quantifying the intensity of blue-silver-stained bands (Fig. 4a, inset, lane2). Fractions from the second peak were collected and again subjected to SEC. The Gaussian-shaped peak indicated a monodisperse protein detergent complex (Fig. 4b) at an elution volume of 67.9 mL, which corresponded to a radius of 5.3 nm (Fig. 4c). This radius indicated the insertion of protein into the detergent micelles. The elution volume of rhodamine B-stained empty FC14 micelles at 79.9 mL corresponded to a radius of 3.7 nm (Fig. 4c), whereas FC14 micelles were reported as having a hydrodynamic radius of 3.2 nm [63]. The value found here is in agreement with the reported value considering the differences in methods, conditions, and the staining of micelles. By correcting the apparent molecular weight of the hisPS2-FC14 complex (185.3 kDa, Fig. 4d) for the empty FC14 micelles (47 kDa, Anatrace), hisPS2 was calculated to form trimers in FC14 micelles. Therefore, from 11 culture, 1.05 ± 0.12 mg homogeneous trimeric hisPS2 was obtained by a combination of Ni-NTA and SEC (Fig. 4e).

Before delipidation of the solubilized membrane proteins by the washing steps during Ni-NTA chromatography, the detection of hisPS2 was poor in western blots. It was thus not possible to establish reliable estimates of the expression level. We therefore used a relative recovery that linked yields of hisPS2 under native purification conditions to yields from the same procedure, but using denaturing conditions throughout (6 M urea, 1% NLS). A relative recovery of 90% was calculated for the ratio between the yields from native and denaturing purification under the final protocol.

On 12% SDS-PAGE (Fig. 3a), hisPS2 exhibited an apparent Mw of ~50 kDa and ~145 kDa for the oligomer band, whereas on 10%, SDS-PAGE (Fig. 4a, inset), the higher Mw band appeared reproducibly at ~100 kDa. This type of Mw shift for helical membrane proteins dependent on the acrylamide concentration has been previously studied and reported [64]. At ~50 kDa, two diffuse bands were observed on 10% SDS-PAGE (Fig. 4a), but not on 12%. The three bands in Fig. 4b...
show different states of the SDS complex of the protein. The appearance of two monomer bands is a typical SDS artifact for membrane proteins when samples are only heated to 46 °C in order to avoid the formation of SDS insoluble aggregates. They are thought to be caused by variations of SDS binding due to the different degrees of denaturation [65].

3.4. Purified hisPS2 is well-folded in FC14 micelles

In order to investigate the secondary structure of purified hisPS2, the protein was subjected to far-UV CD. As shown in Fig. 6a, the CD spectrum at 4 °C displayed one positive peak with a maximum at ∼192 nm and two negative peaks with minima at ∼209 nm and ∼221 nm, which are characteristic for α-helical structures. The spectrum of hisPS2 in the SEC buffer exhibited a similar shape and intensity to the spectrum in the CD buffer above 200 nm. Deconvolution revealed that purified hisPS2 zymogen contained 51.0 ± 1.0% helix and 10.7 ± 0.6% β-strand, which is in agreement with the secondary structure content predicted by RaptorX (Table 1). The addition of cholesterol does not influence the shape of the CD spectrum or the
secondary structure content (Table 1). After validating the presence of the expected helical content, the tertiary structure was investigated by means of near-UV CD. The near-UV CD spectrum at 4 °C showed two positive defined peaks for phenylalanine centered at 259 nm and 265 nm, respectively. The strong positive peak centered at 292 nm is due to tryptophan absorption. Smaller positive peaks between 270 nm and 282 nm represented the tyrosine absorption (Fig. 7a). The presence of the near-UV CD signal indicated that the secondary structure of purified hisPS2 was folded into a well-defined tertiary structure [66].

Fluorescence emission spectra were recorded to investigate the microenvironment of the aromatic amino acids of hisPS2. As shown in Fig. 5a, the fluorescence spectra with different excitation wavelengths displayed varied intensities with distinctive emission maxima. For excitation wavelengths lower than 285 nm, both tyrosine and tryptophan were excited, while for excitation wavelengths higher than 290 nm, only tryptophan was excited and therefore the intensity was somewhat lower. The fluorescence emission maximum was centered at ~326 nm for excitation wavelengths lower than 285 nm. However, for excitation above 290 nm, the fluorescence emission maximum shifted to ~333 nm (Fig. 5b), which corresponded to tryptophan fluorescence.

3.5. Thermal stability of the secondary and tertiary structure of hisPS2

The thermal stability of the hisPS2 structure was investigated by investigating the structural stability by means of CD-spectroscopy, which is much more sensitive to structural changes than UV/Vis absorption spectroscopy (Fig. 6a, d). Upon heating the secondary and tertiary structure changes can be monitored by CD in the far-UV and the near-UV range, respectively. Unlike the far-UV CD spectrum at 4 °C, the 98 °C spectra displayed a peak minimum at 217 nm, which is characteristic for beta-strand-rich structures (Fig. 6a). The 209:217 nm peak ratio decreased from 1.11 at 4 °C to 0.83 at 98 °C, indicating a loss of helical structure and a gain in beta-strand structure. The 209:217 nm peak ratio from the spectrum after cooling back to 4 °C was 0.90. The presence of an isodichroic point at 200 nm indicated that thermal unfolding is a two-state process. Deconvolution of these spectra showed that the helical content decreased from 51% at 4 °C to 22% at 98 °C, while the beta-strand content increased from 10% to 29% (Table 1). The helical content after cooling back to 4 °C increased only slightly to 28%. Accordingly, a decrease in helical segments or average helix length was accompanied by an increase in strand segments or average strand length (Table 2 in Ref [54]).

The temperature reverse scan revealed that the CD signal at 221 nm did not return to the initial value of the signal at 4 °C when cooling back from 98 °C to 4 °C (Fig. 6b). Fitting the CD signals at 221 nm into a two-state transition revealed Tm values of 53.7, 55.2, and 52.9 °C, respectively, for hisPS2 in CD buffer, SEC buffer, and SEC buffer supplemented with CaCl2. It is interesting to note that although hisPS2 adopted a similar secondary structure in the absence and presence of CHS, FC14-CHS-purified hisPS2 exhibited a Tm value more than 10 °C higher than with FC14 micelle alone (Fig. 6c).

The reliability of the deconvolution depends on the sample concentration. The absorption spectrum (Fig. 6d) before and after thermal unfolding exhibited no significant difference, thus indicating that there was no loss of sample during thermal unfolding due to the precipitation of aggregated protein. Therefore, far-UV CD spectra were deconvoluted reliably to high temperatures. Interestingly, the formation of soluble SDS-resistant oligomers was observed. SDS-PAGE analysis (Fig. 6d, inset) showed that after thermal unfolding, hisPS2 formed SDS-resistant aggregates that did not enter the separation gel. This kind of thermal unfolding-induced aggregation has also been reported for other membrane proteins [67,68].

The tertiary structure stability was investigated by means of near-UV CD spectra. Spectra showed a progressive loss of the CD signal for temperatures from 4 °C to 80 °C (Fig. 7a). The change of the CD signal at 292 nm agreed with a two-state transition model for unfolding with a...
$T_m$ of 34.2 °C (Fig. 7b). This indicated that the tertiary structure of hisPS2 unfolds earlier than the secondary structure.

3.6. Calcium induces structural change for hisPS2-WT

It was proposed that PSs form passive calcium leakage channels, a function which is independent of γ-secretase activity [28,69,70]. We therefore investigated the effect of calcium on the secondary structure and local differences of hisPS2-WT and hisPS2-D263A/D366A by far-UV CD and intrinsic fluorescence.

As shown in Fig. 8a and b, hisPS2-WT and hisPS2-D263A/D366A exhibited similar far-UV CD spectra. In the presence of calcium, a slight
decrease of the “w” shape intensity was detected for both hisPS2-WT and hisPS2-D263A/D366A compared to the corresponding spectra without calcium. Deconvolution (Fig. 8c) revealed that in the presence of calcium, the helical content decreased from 46.7% to 43.3% while the beta-strand content increased from 12.0% to 14.3% for hisPS2-WT. HisPS2-D263A/D366A adopted similar structural changes. Although hisPS2-D263A/D366A also displayed a secondary structure change in the presence of calcium, the microenvironment of tryptophan residues was evidently different from the wild type. As shown in Fig. 8d and e, hisPS2-D263A/D366A exhibited 10.8% lower fluorescence intensity than hisPS2-WT. In the presence of calcium, the fluorescence was quenched by 11.1% for hisPS2-WT but only 4.2% for hisPS2-D263A/D366A, which demonstrated the different effects of calcium on these proteins.

4. Discussion

Structural investigations of PSs have been mainly focused on cleaved PS1 as the proteolytic subunit of γ-secretase. The investigation of isolated PSs had been limited to the crystal structure of a bacterial homolog and the NMR structure of the PS1 C-terminal fragment due to the low expression level and poor solution behavior of full-length PSs. Fig. 8. Effect of calcium ions on the secondary structure and fluorescence of hisPS2.

Purified hisPS2 wild type (PS2 WT) or hisPS2 double mutation (PS2 D263A/D366A) was incubated in the presence or absence of CaCl2 (pro: Ca2+ = 1:100 (mol:mol)) at 22 °C for 30 min before being subjected to CD and fluorescence measurements. (a) Far-UV CD spectra of PS2 WT in the presence and absence of Ca2+. (b) Far-UV CD spectra of PS2 D263A/D366A in the presence and absence of Ca2+. (c) Deconvolution of the far-UV CD spectra. (d) Tryptophan fluorescence emission spectrum of PS2 WT in the presence and absence of Ca2+. (e) Tryptophan fluorescence emission spectrum of PS2 D263A/D366A in the presence and absence of Ca2+.
eukaryotic PSs [48]. The structures of cleaved PS1 were unable to explain the auto-inhibitory and activation mechanism of the PS zymogens, as most of the cytoplasmic loops containing the hydrophobic inhibitory segment and the auto-cleavage sites were not observed [52,71]. Furthermore, it was proposed that the inhibitory loop at the cleavage site plays an important role in the regulation of the suggested calcium leakage channel activity of PSs [72], a function of full-length PSs which is independent of their proteolytic activity [28]. Finally, it was also proposed that the loss of essential functions of PSs alone may be a more potent cause of AD pathogenesis than the accumulation of toxic Aβ peptides [25,26]. Therefore, we aimed to express, purify, and characterize full-length human PS2 zymogen to a sufficient quantity and quality for structural studies.

Upon extensive screening of expression, detergent solubilization, and purification, we overcame the aggregation problem of PS2, and milligram quantities of homogeneous hisPS2 were obtained by a combination of Ni-NTA and SEC. The removal of the periplasmic fraction helped to reduce protein degradation during purification. The ratio of the solubilization buffer to cell pellet amount was found to be crucial for the purity and homogeneity of the purified hisPS2. Less than 5 ml/g cell caused an increase in impurities and poor separation of the two major peaks on SEC. This is likely due to an insufficient number of micelles which caused the co-solubilization of aggregated material. A higher ratio (33 ml/g) only increased the separation slightly, but did not change the ratio of the two peaks significantly (data not shown). These results indicated that hisPS2 existed in FC14 micelles in different oligomerization states. However, it was reported that PS oligomerization is not required for γ-secretase activity [48], although the role of oligomerization for the proposed calcium leakage channel function is unclear. In the absence of other γ-secretase proteins, hisPS2 formed a trimer.

Biophysical characterization revealed that the purified hisPS2 was well-folded. Far-UV CD detected the spectrum for a typical helical protein. Deconvolution indicated that the purified hisPS2 contained the expected amount of the secondary structure (Table 1). Near-UV CD proved that the secondary structure of the purified hisPS2 was folded into a well-defined tertiary structure.

Protein tryptophan fluorescence had emission peaks ranging from 300 nm to 350 nm depending on the polarity of the local environment. The peak maximum at ~333 nm for purified hisPS2 revealed the buried hydrophobic microenvironment of tryptophan side chains [73]. In comparison with PS1, seven of the eight tryptophan residues of PS2 are conserved. They are either located within the transmembrane region (2 in the center, 4 at the ends of TMs) or located within the extra-membrane segment in the big loop between TM6 and TM7 (Fig. S1). The microenvironment of the majority of tryptophan residues were able to explain the observed emission maxima of the tryptophan fluorescence spectra. Protein tyrosine fluorescence is typically undetectable, mainly due to tryptophan residues having much higher absorption coefficients and quantum yields and also due to Förster energy transfer from tyrosine to nearby tryptophan [74]. The distances between tyrosine and nearby tryptophan in the cryo-EM structure of PS1 varied from 5 Å to 23 Å, indicating different energy transfer efficiencies [52]. PS2 contains 22 tyrosine residues, 16 of which are conserved (Fig. S1). For PS2, it is also likely that different energy transfer efficiencies exist between tyrosine and tryptophan. This explains the observation that when exiled at 275 nm, hisPS2 exhibited an emission maximum at ~326 nm instead of ~333 nm. The maximum of the fluorescence spectrum shifted to a lower wavelength because the tryptophan fluorescence was overlaid by unquenched tyrosine fluorescence [75]. The intrinsic protein fluorescence spectroscopy thus revealed the expected micro-environment of tryptophan side chains, which is another indicator of proper folding.

It was recently proposed that AD mutations cause AD pathology by impairing the overall protein stability of PS1 [76]. We therefore investigated the thermal stability of the secondary and tertiary structure of hisPS2. The thermal unfolding of hisPS2 in FC14 was an irreversible process, in which part of the helical structure transformed into beta-strand. Cross-beta aggregation prediction by Tango [77], reveals three regions in PS2 which are aggregation-prone: T142-160Y in TM2, W171-188L in TM3, and E286-K306 in TM8 (Fig. S1). Among these three regions, TM2 has the highest aggregation probability, which is in agreement with the observed structural flexibility of PS1 [52,71]. These regions account for 12.2% of the total sequence, which explained the transition from helical to beta-strand structure detected by CD. Such a transition likely causes the formation of the observed heat-stable aggregates by intermolecular beta-strand interaction during thermal unfolding [78]. This is in agreement with the irreversibility of structural transformations of hisPS2 as observed from the temperature reverse scan, the deconvoluted structure composition at different temperatures, and the detection of SDS-resistant aggregates. The presence of 22% helix at 98 °C was probably due to the extreme stability of the detergent-bound helix, which prevented it from unfolding – a phenomenon also observed for other membrane proteins [79,80]. Cholesterol significantly increased the thermal stability of hisPS2, which provides evidence that cholesterol acts as a natural ligand. It was observed that aberrant intracellular cholesterol transport alters PS localization and enhances Aβ production [81,82]. Therefore, cholesterol might regulate Aβ processing by altering the structural stability of PSs. The relatively low Tm value (34.2 °C) of the tertiary structure was in agreement with the reported loose packing of PS1 TMs in the cryo-EM structure [52] and the presence of the interconverting conformations of γ-secretase [83]. Such a Tm value also explained the difficulties (aggregation and impurities) encountered during purification.

For maturation, PSs undergo auto-cleavage within the hydrophobic sequence (E280 to E309 for PS1, E286 to K306 for PS2, Fig. S1) located in the large loop between TM6 and TM7. It was proposed that cleavage removes the inhibitory exon 9 loop which occupies the substrate binding site, thus resulting in an active enzyme [84]. The hydrophobic segment is not only involved in modulating enzyme activation, substrate binding, and processing [84,85], but might also be involved in controlling the potential calcium channel leakage function [72]. Experimental evidence has shown that the PS1-ΔE9 mutant not only exhibited proteolytic activity [86], but also functioned as a calcium leakage channel [28]. The corresponding sequence from the cryo-EM structure of PS1 exhibited a short beta-strand for residues 285–288, while the densities were missing for the rest of the residues [71]. It was proposed that the release of the inhibitory loop (S290 to S319 in PS1) by auto-cleavage after maturation controls both the proteolytic activity and the calcium leakage channel function. The ε-nitrogen of PS1-W294 is in close contact with the nearest δ-oxygen of the catalytic residue PS1-D385 (5 Å) in the immature PSs. This distance increases to > 10 Å in the mature PSs due to the movement of the inhibitory loop out of the proteolytic active site [54]. It is therefore reasonable to assume that after the release of the inhibitory loop, W294 becomes more exposed. However, experimental results showed that the Ca2+ channel leakage function of PSs was independent of their proteolytic activity, although residue D385 was indispensable. PS1-D275A abolished proteolytic activity (resulting in uncleaved PSs) but did not influence the Ca2+ leakage function while the PS1-D385A mutation lacked both the proteolytic activity and the Ca2+ leakage function [69,70,87,88]. In this study, it was observed that hisPS2-WT zymogen underwent conformational changes in the presence of calcium, which suggested that maturation might not be required for the calcium channel function. However, hisPS2-D263A/D366A, which abolishes both the proteolytic activity and the calcium channel function, adopted a different conformation and exhibited minor structural changes in the presence of calcium. These results are in agreement with the previous experimental observations [28]. W300 in PS2 (Fig. S1), which is located within the inhibiting sequence, might be (partially) responsible for the observed 11% quenching. For hisPS2-D263A/D366A, the quenching decreased to only 4.2%, probably because it adopted a different conformation. Indeed, the quantum yield of hisPS2-D263A/D366A was similar to the
hisPS2-WT in the presence of calcium, thus indicating that W300 might already be more exposed in hisPS2-D263A/D366A than in PS2-WT. The structural changes detected in the presence of calcium for hisPS2-WT probably reflected to some extent the movement of the inhibitory loop, which might open the calcium channel of the PSs. However, knowledge of the actual molecular mechanism of the maturation and calcium channel leakage function of PSs requires the atomic structure of the uncleaved yzomogen of PSs.

Conflict of interest

The authors declare that there are no conflicts of interest.

Author contribution

G.Y., K.Y., and J.L. conceived and designed the experiments. J.L. supervised the study. G.Y. performed the experiments. G.Y., K.Y., and J.L. analyzed the data and wrote the manuscript. All authors contributed to the discussion and revision of the manuscript.

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Appendix A. Supplementary data

Supplementary data associated with this article can be found, in the online version, at http://dx.doi.org/10.1016/j.pcbio.2017.09.012.

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