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γ-Secretase Composed of PS1/Pen2/Aph1a Can Cleave Notch and Amyloid Precursor Protein in the Absence of Nicastrin

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Introduction

γ-Secretase is a multiprotein, intramembrane-cleaving protease with a growing list of protein substrates, including the Notch receptors and the amyloid precursor protein. The four components of γ-secretase complex—presenilin (PS), nicastrin (NCT), Pen2, and Aph1—are all thought to be essential for activity. The catalytic domain resides within PS proteins, NCT has been suggested to be critical for substrate recognition, and the contributions of Pen2 and Aph1 remain unclear. The role of NCT has been challenged recently by the observation that a critical residue (E332) in NCT, which had been thought to be essential for γ-secretase activity, is instead involved in complex maturation. Here, we report that NCT is dispensable for γ-secretase activity. NCT-independent γ-secretase activity can be detected in two independent NCT-deficient mouse embryonic fibroblast lines and blocked by the γ-secretase inhibitors N-[N-(3,5-difluorophenacetyl)-l-alanyl]-S-phenylglycine t-butyler ester and L-685,458. This catalytic activity requires prior ectodomain shedding of the substrate and can cleave ligand-activated endogenous Notch receptors, indicating presence of this activity at the plasma membrane. Small interfering RNA knockdown experiments demonstrated that NCT-independent γ-secretase activity requires the presence of PS1, Pen2, and Aph1 but can tolerate knockdown of PS2 or Aph1b. We conclude that a PS1/Pen2/Aph1a trimeric complex is an active enzyme, displaying biochemical properties similar to those of γ-secretase and roughly 30% of its activity when normalized to PS1 N-terminal fragment levels. This PS1/Pen2/Aph1a complex, however, is highly unstable. Thus, NCT acts to stabilize γ-secretase but is not required for substrate recognition.

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We established the Institute for Biotechnology, Leuven, Belgium (Herreman et al., 1999). Bart De Strooper (Katholieke Universiteit Leuven, Leuven, and Flanders and wild-type (NCTPW) were generous gifts from Dr. Philip C. Wong (Johns Hopkins University, Baltimore, MD) (Li et al., 2003). PS1/PS2 double knock-out (PSDKO) cells were collected after 20 h and lysed with lysis buffer A containing 8M urea, 1% SDS, 50 mM Tris-HCl, pH 6.8, 2 mM EDTA, 2 mM DTT, and 1 mM disulfopiperazinyl L-alanyl) S-phenylglycine t-butyl ester (DAPT) or 1 mM L-685,458; Calbiochem).

Materials and Methods

Plasmids. pCS2+/N1ΔE-6MT, pCS2+/C99-6MT, and pCDNA3/PS1wt vectors have been described previously (Kopan et al., 1996; Schroeter et al., 2003).

Cell lines. NCT-deficient NcstnmutPw/NcstnmutPw (NCTPW−/−) mouse embryonic fibroblast (MEF) lines were generous gifts from Dr. Philip C. Wong (Johns Hopkins University, Baltimore, MD) (Li et al., 2003). PS1/PS2 double knock-out (PSDKO) and PS1/PS2 wild-type (PSWT+) MEF lines were generous gifts from Dr. Bart De Strooper (Katholieke Universiteit Leuven, Leuven, and Flanders Institute for Biotechnology, Leuven, Belgium) (Herremans et al., 1999). We established NcstnmutPw/NcstnmutPw MEF using NCTPW−/− breeders generously provided by Dr. Richard Rozmahel (University of Toronto, Toronto, Canada) (Nguyen et al., 2006) using a previously described method (Li et al., 2003) but with a slight modification. Briefly, embryonic day (E)9.5 embryos were minced and resuspended in DMEM containing 0.025% trypsin/EDTA and incubated at 37°C for 20 min. DMEM supplemented with 10% fetal bovine serum (FBS) was added to neutralize the trypsin/EDTA, and the cells were further dissociated by pipetting, plated into 24-well plates, and subsequently immobilized with polyoma large T antigen. Immortalized NCT knock-out clones were confirmed by both genotyping and Western blot analysis.

Cell culture and plasmid transfection. For most experiments, cells were maintained in DMEM supplemented with 10% FBS, 2 mM glutamine and incubated at 37°C and 5% CO₂. To examine the generation of Notch intracellular domain (NICD) and amyloid intracellular domain (AICD), cells were transfected with pCS2+/N1ΔE-6MT and pCS2+/C99-6MT vectors or empty vector using Lipofectamine 2000 (Invitrogen) or FuGENE HD (Roche) transfection reagents, according to the manufacturer’s instructions. The transfected cells were subsequently cultured in the presence or absence of proteasome inhibitors (4.5 μM lactacystin, 0.15 μM MG262; Calbiochem) and γ-secretase inhibitors [either 1 μM N-[3-(3,5-difluorophenacetyl-L-alanyl)] S-phenylglycine t-butyl ester (DAPT) or 1 μM L-685,458; Calbiochem].

Aβ analysis. pCS2+/C99-6MT-transfected cells were cultured in OPTI-MEM supplemented with 7% FBS, 2 mM glutamine, 10 μM nonessential amino acids, 10 μM phosphoramidon disodium (Sigma), with or without proteasome inhibitors and γ-secretase inhibitors. Both cells and conditioned media (CMS) were collected 20–22 h later. The CMSs were supplemented with 1 μM AEBSF [4-(2-aminoethyl)benzenesulfonyl fluoride hydrochloride; Roche], centrifuged, and analyzed by Aβ ELISA (Invitrogen) according to the manufacturer’s instructions, but with a slight modification. The incubation time for antibodies and stabilized chromogen and the time for each wash step were increased to make testing lower levels of Aβ more accurate. Each sample was run in duplicate. The CMSs from cells transfected with empty vector were used as blanks.

Coculture of NCTRR−/− cells with ligand-expressing cells. NCTRR−/− or PSDKO cells were coseeded with either CHO-DLL1-IRE-GFP (CHO: Chinese hamster ovary; IRES: internal ribosome entry site; GFP: green fluorescent protein) or CHO-GFP control cells (these two cell lines have been described previously) (Ong et al., 2008), cultured for 24 h, and then treated with or without lactacystin/MG262 and DAPT. The cells were collected after 20 h and lysed with lysis buffer A containing 8% urea, 1% SDS, 50 mM Tris·HCl, pH 6.8, 2 mM EDTA, 2 mM DTT, and 1% protease inhibitor cocktail (Roche) (Zhao et al., 2004).

siRNA transfection. ON-TARGETplus small interfering RNA (siRNA) smart pools targeting mouse PS1, PS2, Pen2, Aphi1a, and Aphi1b were purchased from Dharmacon (see supplemental Table 1, available at www.jneurosci.org as supplemental material for sequences). ON-TARGETplus nontargeting siRNA control (Dharmacon) was used as a negative control. siRNAs were transfected into NCTPW−/− cells using DharmaFect 1 (Dharmacon) according to the manufacturer’s instructions. The cells were cultured in complete medium for 48 h and then transfected with pCS2+/N1ΔE-6MT vector. Knockdown efficiency was determined by either Western blotting or quantitative reverse transcription (qRT)-PCR.

qRT-PCR. siRNA-transfected NCTPW−/− cells were cultured for 68 h. The total RNA was then isolated using RNeasy Micro kit (Qiagen) according to the manufacturer’s instructions. The reverse transcription was
performed using SuperScript II reverse transcriptase (Invitrogen) with oligo(dT) 

as the primer. Quantitative PCR was performed according to previously described methods (Lee et al., 2007) (see supplemental Table 2, available at www.jneurosci.org as supplemental material for primer sequences).

**EDTA treatment.** NCT 

/−/−, NCT 

+/−, PS 

DKO, and PS 

+/− cells were washed once with HBSS buffer (without Ca or Mg) and incubated in 1.5 mM EDTA/HBSS buffer for 40 min at 37°C. For γ-secretase inhibition, cells were treated with 1 μM DAPT for 4 h before and during EDTA incubation.

**Western blotting and antibodies.** Cells were lysed in lysis buffer A with brief sonication, and the protein concentration was determined using a BCA kit (Pierce). Twenty to 30 μg of total protein per sample was analyzed by SDS-PAGE/Western blot analysis. The following antibodies were used in this study: anti-V1744 (Cell Signaling Technology), anti-NCT (N1660, Sigma), anti-PS1-NTF (Santa Cruz Biotechnology), anti-PS2-CTF antibodies B24.2 and G24 [generous gifts from Dr. Bart De Strooper (Herreman et al., 1999) and Dr. Taikoe Tomita (Tomita et al., 1998, respectively), anti-Aph1a (Covance), anti-Notch1 ANK domain (mAN1) (Huppert et al., 2000), anti-β-actin (Sigma), and anti-myc (9E10).

### Results

**NICD is generated in NCT−/− cells but not in PSDKO cells**

All GxGD proteases prefer substrates that have undergone ectodomain shedding, and all, including PS, contain a substrate recognition domain (Kornilova et al., 2005). We hypothesized that if the role of NCT was to stabilize γ-secretase, but not to bind substrates, we should be able to detect γ-secretase activity in the absence of NCT. To test this hypothesis, we transiently transfected wild-type (NCT 

+/−) and NCT 

−/− cells (from Dr. P.C. Wong) with ectodomain-truncated, Notch1-expressing vector (pCS2+/N1ΔE-6MT). The truncated Notch1 ΔE-6myc protein is a ligand-independent and direct substrate of γ-secretase, similar to Notch extracellular truncation (NEXT) (Fig. 1A). To enhance detection, transfected cells were cultured in the presence of proteasome inhibitors that reduce the turnover of the NICD (Tagami et al., 2008) and increase the sensitivity of detection for γ-secretase-like activity.

To our knowledge, γ-secretase is the only enzyme that cleaves Notch1 at its S3 site, but the existence of other enzymes with γ-secretase-like activity has been proposed to compensate for PS loss (Huppert et al., 2005). To ask whether a Notch intramembrane protease exists in other γ-secretase-deficient cells, we examined NICD generation in PS 

DKO cells, which are deficient in N1ΔE-6MT, yet NICD was undetectable in samples from PS 

DKO cells under the same conditions that allow NICD accumulation in NCT 

+/− cells (Fig. 1B, lanes 9 and 13). The generation of NICD in PS 

DKO cells can be rescued by transfection of wild-type PS1 or PS2 but not an inactive aspartyl mutant PS1, confirming that the failure to generate NICD by PS 

DKO cells is not an indication of other defects (supplemental Fig. 2, available at www.jneurosci.org as supplemental material) (Schroeter et al., 2003). Overall, these results indicate that a γ-secretase-like activity in NCT 

+/− cells cleaves Notch to generate NICD and that this activity does not exist in PS 

DKO cells.

**Generation of NICD is observed in two independent NCT−/+ lines**

We noticed a faint cross-reacting band with mobility similar to that of immature NCT in NCT 

+/− cells (Fig. 1B), NCT 

+/− cells were derived from an NCT knock-out embryo in which the critical part of the DAP (DYIGS and peptidase homologous region) domain (exon 9, intron 9, and part of exon 10) was replaced
γ-secretase-like activity is independent of the specific NCT<sup>−/−</sup> line used.

**Endogenous Notch can be cleaved by the γ-secretase-like activity in NCT<sup>−/−</sup> cells but not in PS<sup>DKO</sup> cells**

The γ-secretase-like activity that survived removal of NCT may only cleave ectopically expressed Notch substrates lacking an extracellular domain. To ask whether this enzyme could cleave endogenous Notch, we examined the cleavage of endogenous Notch1 receptors under conditions that induce ectodomain shedding at the cell surface. In the absence of ligands, a calcium-stabilized negative regulatory region prevents metalloprotease access to the S2 cleavage site (Gordon et al., 2007). Ligand binding or treatment with EDTA (Rand et al., 2000) is thought to result in a conformational change that leads to the exposure of the S2 cleavage site and subsequent cleavage by ADAM (a disintegrin and metalloproteinase) metalloproteases (Kopan and Ilagan, 2009), generating NEXT (Fig. 1A), a direct substrate of γ-secretase. Western blot analyses showed that all the MEF lines (except for NCT<sub>DKO</sub><sup>−/−</sup> expressed endogenous Notch1. To test whether the generation of NICD from endogenous Notch1 can occur without NCT protein, PS<sup>DKO</sup> cells and NCT<sub>DKO</sub><sup>−/−</sup>, NCT<sub>RKO</sub><sup>−/−</sup>, and wild-type MEFs were treated with or without EDTA to induce S2 cleavage (see shift in mobility in Fig. 3A). Importantly, after treatment with EDTA, DAPT-sensitive NICD generation was detected in NCT<sub>RKO</sub><sup>−/−</sup> and wild-type cells (Fig. 3A, lanes 8–9 and 11–12). Moreover, after EDTA treatment, endogenous NICD was observed in NCT<sup>−/−</sup> cells even without proteasome inhibition (Fig. 3A, B). This result confirms that ectodomain shedding is required for the release of NICD in NCT<sub>RKO</sub><sup>−/−</sup> cells (Fig. 3A, compare lanes 7–8).

**Figure 3.** NICD is generated from endogenous Notch1 in NCT<sup>−/−</sup> cells but not in PS<sup>DKO</sup> cells. A, Representative Western blots of NICD levels in EDTA-treated cells. Metalloprotease-mediated shedding occurs after calcium chelation-induced dissociation of endogenous Notch1. EDTA treatment led to reduced amounts of furin-cleaved Notch (transmembrane/intracellular (TMIC)) and of NICD levels in EDTA-treated cells. Metalloprotease-mediated shedding occurs after calcium chelation-induced dissociation of endogenous Notch1 receptors after activation by ligands presented by neighboring cells. Note that NICD was only detected in coculture of NCTRR<sup>−/−</sup> endogenous Notch1 receptors after activation by ligands presented by neighboring cells. With EDTA, DAPT-sensitive NICD generation was detected in NCTRR<sup>−/−</sup> and wild-type cells (Fig. 3A, lanes 8–9 and 11–12). Moreover, after EDTA treatment, endogenous NICD was observed in NCT<sup>−/−</sup> cells even without proteasome inhibition (Fig. 3A, B). This result confirms that ectodomain shedding is required for the release of NICD in NCTRR<sup>−/−</sup> cells (Fig. 3A, compare lanes 7–8). In contrast, Notch1-expressing PS<sup>DKO</sup> cells accumulated the S2 cleavage product NEXT but did not produce NICD (Fig. 3A, lanes 1 and 2). Because removal of calcium with EDTA occurs at or near the cell surface and because both ADAM10 and Notch1 are mainly localized at the plasma membrane (van Tetering et al., 2009), this result suggests that the NCT-deficient γ-secretase-like protease performs its activity at or close to the plasma membrane.

To obtain physiologically relevant evidence that γ-secretase-like activity reached the cell surface, we asked whether the Notch1 receptors in NCT<sub>RKO</sub><sup>−/−</sup> cells could be activated by ligands presented on neighboring cells. We cocultured NCT<sub>RKO</sub><sup>−/−</sup> (or control PS<sup>DKO</sup>) cells with either ligand-expressing (CHO-DLL1) or control (CHO-GFP) cells in the presence or absence of proteasome inhibitors. Western blot analysis shows that coculture with CHO-DLL1 cells induced endogenous Notch1 S2 cleavage (Fig. 3B). As expected, if the enzyme reached the cell surface, endogenous NICD was detected in NCT<sub>RKO</sub><sup>−/−</sup> cells cocultured with

with the neomycin resistance gene (Li et al., 2003). To test whether γ-secretase-like activity in this specific cell line was due to residual NCT expression, we analyzed a second, independently generated NCT knock-out mouse line (from Dr. Richard Rozmahel) in which exon 3 and a part of exon 4 had been deleted, and therefore splicing downstream of the deletion would shift the reading frame (Nguyen et al., 2006). We isolated NCT<sub>RR</sub><sup>−/−</sup> MEF cells from E9.5 NCT<sub>RR</sub><sup>−/−</sup> embryos and examined NCT production and γ-secretase-like activity in these cells. No NCT immunoreactivity was observed in NCT<sub>RR</sub><sup>−/−</sup> cell extracts (Fig. 2, lanes 9–12). Importantly, as in NCT<sub>PW</sub><sup>−/−</sup> cells, endoproteolysis of PS1 in NCT<sub>RR</sub><sup>−/−</sup> was also observed, confirming that PS endoproteolysis does not require NCT. Accordingly, in the presence of proteasome inhibitors we observed NICD accumulation in NCT<sub>RR</sub><sup>−/−</sup> cells (Fig. 2, lane 11). In this cell line, NICD production was again blocked by either L-685,458 (Fig. 2, lane 12) or DAPT (Fig. 3A). Overall, these data clearly demonstrate that generation of NICD by residual

**Figure 3.** NICD is generated from endogenous Notch1 in NCT<sup>−/−</sup> cells but not in PS<sup>DKO</sup> cells. A, Representative Western blots of NICD levels in EDTA-treated cells. Metalloprotease-mediated shedding occurs after calcium chelation-induced dissociation of endogenous Notch1. EDTA treatment led to reduced amounts of furin-cleaved Notch (transmembrane/intracellular (TMIC)) and of NICD levels in EDTA-treated cells. Metalloprotease-mediated shedding occurs after calcium chelation-induced dissociation of endogenous Notch1 receptors after activation by ligands presented by neighboring cells. Note that NICD was only detected in coculture of NCTRR<sup>−/−</sup> endogenous Notch1 receptors after activation by ligands presented by neighboring cells. With EDTA, DAPT-sensitive NICD generation was detected in NCTRR<sup>−/−</sup> and wild-type cells (Fig. 3A, lanes 8–9 and 11–12). Moreover, after EDTA treatment, endogenous NICD was observed in NCT<sup>−/−</sup> cells even without proteasome inhibition (Fig. 3A, B). This result confirms that ectodomain shedding is required for the release of NICD in NCTRR<sup>−/−</sup> cells (Fig. 3A, compare lanes 7–8). In contrast, Notch1-expressing PS<sup>DKO</sup> cells accumulated the S2 cleavage product NEXT but did not produce NICD (Fig. 3A, lanes 1 and 2). Because removal of calcium with EDTA occurs at or near the cell surface and because both ADAM10 and Notch1 are mainly localized at the plasma membrane (van Tetering et al., 2009), this result suggests that the NCT-deficient γ-secretase-like protease performs its activity at or close to the plasma membrane.

To obtain physiologically relevant evidence that γ-secretase-like activity reached the cell surface, we asked whether the Notch1 receptors in NCT<sub>RR</sub><sup>−/−</sup> cells could be activated by ligands presented on neighboring cells. We cocultured NCT<sub>RR</sub><sup>−/−</sup> (or control PS<sup>DKO</sup>) cells with either ligand-expressing (CHO-DLL1) or control (CHO-GFP) cells in the presence or absence of proteasome inhibitors. Western blot analysis shows that coculture with CHO-DLL1 cells induced endogenous Notch1 S2 cleavage (Fig. 3B). As expected, if the enzyme reached the cell surface, endogenous NICD was detected in NCT<sub>RR</sub><sup>−/−</sup> cells cocultured with...
CHO-DLL1 (Fig. 3B, lanes 8 and 9) but not in NCTRR cells cocultured with CHO-GFP. DAPT blocked the generation of NICD in NCTRR cells (Fig. 3B, lane 10), and PS2KO cells cocultured with CHO-DLL1 cells did not release NICD, indicating that NICD was not generated by the CHO cells (Fig. 3B, lanes 4, 5, 11, and 12). Together, these results confirm that the γ-secretase-like activity in NCT−/− cells requires substrate ectodomain shedding and can reach the plasma membrane to mediate the cleavage of ligand-activated Notch.

C99 is converted into AICD and Aβ in NCT−/− cells but not in PS2KO cells

Another important substrate of γ-secretase is APP, which is involved in the pathogenesis of Alzheimer’s disease. Since mutations in NCT were shown previously to differentially affect APP and Notch processing (Chen et al., 2001), we asked whether a truncated APP substrate (C99) could be processed in NCT−/− cells by the γ-secretase-like activity. We transiently transfected PS2KO, NCTpw−/−, NCTRR−/−, and wild-type MEF cells with C99-6myc-expressing vector. Western blot analyses confirmed that a DAPT-sensitive fragment (AICD-6myc) present in wild-type cells but not in PS2KO cells was also seen in two NCT-deficient cell lines (Fig. 4A, lanes 3, 7, and 13), suggesting that NCT−/− cells are able to cleave C99 (Fig. 4A; note that proteasome inhibition was required to detect the cleavage products). A longer exposure detected low amounts of AICD-myc in NCTpw−/− cells not treated with proteasome inhibitors (supplemental Fig. 1, available at www.jneurosci.org as supplemental material). To address whether Aβ was also produced in NCT−/− cells, we performed ELISA analysis on conditioned media to determine the secreted Aβ levels. Aβ40 was detected in conditioned media from NCT−/− cells but not from PS2KO cells (Fig. 4B) and was abolished after DAPT addition (Fig. 4B). We were unable to detect Aβ42 levels with the Aβ42 ELISA kit, likely due to the low transfection efficiency of MEF cells. Collectively, these data indicate that not only can the residual γ-secretase activity cleave APP, it can proceed from the e-cleavage site to the γ-cleavage site, a hallmark of γ-secretase (Tanzi and Bertram, 2005; Haass and Selkoe, 2007), even in the absence of NCT.

γ-Secretase-like activity in NCT−/− cells is PS1 dependent

The fact that two unrelated γ-secretase inhibitors abolished the γ-secretase-like activity in NCT−/− cell lines, and that PS2KO cells did not exhibit this activity, strongly implies that PS is the active enzyme in NCT−/− cells. To test this, we used siRNAs to knock down PS1 and PS2 (alone or together) in NCTRR−/− cells transfected with N1ΔE-6MT. Western blot analysis confirmed that PS1 siRNA markedly decreased both full-length PS1 and PS1-NTF (Fig. 5A, lanes 3 and 4), whereas control siRNA and PS2 siRNA did not affect PS1 protein level (Fig. 5A, lanes 1, 2, 5, and 6). Interestingly, PS1 knockdown alone was sufficient to significantly diminish NICD production, whereas knockdown of PS2 did not show an obvious effect (Fig. 5A, lanes 5–6). Since we could not detect PS2 in these cells with two different established antibodies (Tomita et al., 1998; Herreman et al., 1999), the efficiency of PS2 siRNA knockdown was confirmed by qRT-PCR (Fig. 5B). Accordingly, cotransfection of PS1 siRNA together with PS2 siRNA did not further reduce NICD amounts (Fig. 5A, lanes 7 and 8). Similar results were obtained with NCTpw−/− cells (supplemental Fig. 3, available at www.jneurosci.org as supplemental material). These results suggest that PS1 or a PS1-containing enzyme in NCT−/− cells mediates NICD production.

To ask whether PS2 can contribute to γ-secretase activity in NCT−/− cells, we overexpressed PS2 in NCTRR−/− cells. We first performed PS1 mRNA knockdown and then asked whether cotransfection of the substrate (pCS2+/N1ΔE-6MT) with either human PS1 or PS2 expression vector into these mouse PS1-depleted NCT−/− cells would restore the γ-secretase activity. While both PS1 and PS2 restored γ-secretase activity equally well in PS2KO cells, only human PS1, but not human PS2, restored γ-secretase activity in PS1 siRNA-transfected NCT−/− cells (supplemental Fig. 2, available at www.jneurosci.org as supplemental material). These data confirmed that PS2 protein could not contribute to the γ-secretase activity in NCT−/− cells.

Pen2 and Aph1a are involved in the γ-secretase-like activity

To ask whether PS1 acted as a single-molecule protease like SPP (Goldé et al., 2009), we examined whether Pen2 and Aph1 were still required for the PS1 activity in the absence of NCT. We performed Pen2 or Aph1 knockdown in NCTR−/− cells using siRNA pools and confirmed the efficiencies of siRNA knockdown by qRT-PCR (Fig. 6B, D). Western blot analyses demonstrated that knockdown of Pen2 did not decrease the amount of full-length PS1 but dramatically reduced PS1-NTF levels and, concomitantly, the amounts of NICD (Fig. 6A, lanes 3–8). Of the three murine Aph1 genes (Ma et al., 2005; Serneels et al., 2005), only knockdown of Aph1a in NCTR−/− cells reduced PS1-NTF and NICD levels, whereas full-length PS1 levels remained unchanged (Fig. 6C, lanes 3–6). Aph1b siRNA did not impact PS1-
obtained with NCTPW PS2 antibodies (data not shown), knockdown of PS2 was confirmed by qRT-PCR. To accurately evaluate the remaining/H9253 complex lacking NCT is highly unstable in the detergent solution, although we could readily detect NCT composed of three proteins (PS1, Pen2, and Aph1a) provides frequently transfected with pCS2-transfected with PS1, PS2, PS1-plus-PS2, or nontargeting control siRNAs. The cells were subsequently transfected with pCS2+/NΔE-6MT and cultured in the presence of proteasome inhibitors. Cell lysates were then analyzed by Western blotting with antibodies against PS1NT, Myc, V1744, and β-actin. PS1 siRNA markedly decreased both full-length PS1 (FL-PS1) and PS1-NTF. Although the levels of NΔE-6Myc are similar in every extract, NICD accumulation was strikingly reduced in PS1 siRNA-transfected cells and in PS1-plus-PS2 siRNA-transfected cells but not in siRNA control (Ctr)- or PS2 siRNA-transfected cells. Since the PS2 proteins in these NCT−/− cells were undetectable by Western blot analysis using two well-characterized anti-PS2 antibodies (data not shown), knockdown of PS2 was confirmed by qRT-PCR.

Figure 5. γ-secretase-like activity in NCT−/− cells is PS1 dependent. A, Representative Western blots of NICD levels in PS siRNA-transfected NCTRR−/− cells. NCTRR−/− cells were transfected with PS1, PS2, PS1-plus-PS2, or nontargeting control siRNAs. The cells were subsequently transfected with pCS2+/NΔE-6MT and cultured in the presence of proteasome inhibitors. Cell lysates were then analyzed by Western blotting with antibodies against PS1NT, Myc, V1744, and β-actin. PS1 siRNA markedly decreased both full-length PS1 (FL-PS1) and PS1-NTF. Although the levels of NΔE-6Myc are similar in every extract, NICD accumulation was strikingly reduced in PS1 siRNA-transfected cells and in PS1-plus-PS2 siRNA-transfected cells but not in siRNA control (Ctr)- or PS2 siRNA-transfected cells. B, Since the PS2 proteins in these NCT−/− cells were undetectable by Western blot analysis using two well-characterized anti-PS2 antibodies (data not shown), knockdown of PS2 was confirmed by qRT-PCR.

NTF or NICD level (Fig. 6C, lanes 7–10). Similar results were obtained with NCTpw−/− cells (supplemental Fig. 3, available at www.jneurosci.org as supplemental material). These results suggest that in NCT−/− cells, an unstable γ-secretase isomorph composed of three proteins (PS1, Pen2, and Aph1a) provides γ-secretase activity that correlates strongly with the amount of endoproteolytically processed PS1-NTF but not full-length PS1. Thus, PS1 activity requires endoproteolysis and the functions of Pen2 and Aph1a.

Trimeric γ-secretase retains 50% of the enzyme activity To accurately evaluate the remaining γ-secretase activity in NCT−/− cells, we tried to establish a cell-free assay. However, although we could readily detect γ-secretase activity in solubilized membranes from wild-type cells, we failed to detect any γ-secretase-like activity in solubilized membranes from NCT−/− cells (data not shown). It is likely that the trimeric γ-secretase complex lacking NCT is highly unstable in the detergent solution, as it was unstable in blue native gels (data not shown). Therefore, we compared the relative activity of γ-secretase in NCT−/− cells and wild-type cells using a semiquantitative Western blot analysis of extracts isolated from cells cultured in the presence of proteasome inhibitors. As shown in supplemental Figure 4 (available at www.jneurosci.org as supplemental material), in cell lysates containing equivalent levels of PS1 NTF fragments, NCT−/− cells produced 50–55% of the NICD produced by wild-type lines. From these data, we can conclude that the trimeric enzyme is much more active than would be expected if NCT was providing the substrate recognition function in the complex. We cannot rule out an indirect contribution by NCT to overall activity due to increased enzyme stability.

Discussion

PS, NCT, Pen2, and Aph1 are the four proteins essential for reconstituting robust γ-secretase activity (De Strooper, 2003; Iwatsubo, 2004; Spasic and Annaert, 2008). With the exception of PS, which harbors the catalytic site of γ-secretase, the precise contributions of the other proteins to γ-secretase activity remain unclear, and PS remains the only GxGD protease that requires partners for its protease activity. Here, we have provided direct evidence that when PS1/Pen2/Aph1a are present, γ-secretase can assemble and acquire catalytic activity without NCT, cleaving both Notch and APP (C99) substrates. Furthermore, this three-protein enzyme remains biochemically similar to the four-protein enzyme: both are DAPT and L-685,458 sensitive (Fig. 1B), both generate NICD at the S3 site (Figs. 1–3), and both cleave APP at ε and γ to release AICD and Aβ40, respectively (Fig. 4). Importantly, ligand-mediated activation of endogenous Notch can also be observed in NCT−/− cells (Fig. 3B). However, this residual activity is not sufficient to permit survival of NCT−/− embryos (Li et al., 2003; Nguyen et al., 2006).

As this manuscript was being prepared, another group identified NCT-independent PS1 mutants (PS1 S438P or PS1 F411Y/S438P) (Futai et al., 2009), suggesting that perhaps this S438P mutation increased complex stability. We aligned PS proteins with their homologues and found that a Pro residue is located at a conserved position in TMD9 of SPP (P324; supplemental Fig. 5, available at www.jneurosci.org as supplemental material), suggesting that the α-helix-breaking Pro residue may increase stability of GxGD proteases. Interestingly, this analysis revealed that one of the two Caenorhabditis elegans PS proteins, Hop-1, also contains a Pro at the homologous position (P329) whereas the other (Sel-12) does not. Consistent with the hypothesis that γ-secretase complexes centered around Hop-1 may have reduced dependency on the Aph2/NCT gene (Goutte et al., 2000), Aph-2/Sel-12 double mutants are strongly affected but Aph-2/Sel-12 double mutants are not (Francis et al., 2002). This result indicates that Hop-1 is active without Aph-2, whereas Sel-12 is dependent on Aph-2 for full activity. NCT has been proposed to play a role in PS1 trafficking. In the absence of NCT, most PS1 proteins remain in the endoplasmic reticulum; PS1-NTFs were undetectable at the plasma membrane (Zhang et al., 2005). We have provided indirect evidence that at least some PS1 NTF-containing γ-secretase can reach the plasma membrane where it can activate a ligand-dependent Notch signal in the absence of NCT. Coupled with the genetic data (Francis et al., 2002) and with the analysis of NCT mutants (Chavez-Gutierrez et al., 2008), these observations favor a model in which NCT acts to stabilize γ-secretase but is not required for catalytic activity.

Although we cannot completely rule out the possibility that NCT contributes to γ-secretase activity and that this contribu-
tion requires E333 (Dries et al., 2009), our findings indicate that neither substrate recognition nor contribution to catalytic activity by NCT is necessary for \( \gamma \)-secretase activity. A distinct property of \( \gamma \)-secretase-mediated intramembrane cleavage is that it requires prior ectodomain shedding of the substrate; the large extracellular domain of NCT has been proposed to act as the gatekeeper that measures the size of the substrate ectodomain and the availability of a free N terminus (Hu et al., 2002; De Strooper, 2005; Shah et al., 2005). However, we found that NCT-deficient \( \gamma \)-secretase activity still requires prior ectodomain shedding of its substrates (Fig. 3) (Mumm et al., 2000). It is unlikely that an unknown protein may function as NCT in NCT-deficient cells, and no additional NCT homologues have been found in the mouse or \( C. \) elegans genome. Indeed, a substrate-binding site has also been identified at the interface of PS1 NTF/CTF (Kornilova et al., 2005); this site may also be sensitive to the presence of a large extracellular domain. Additionally, the GxGD motif in PS has been reported to contribute to substrate identification (Yamasaki et al., 2006). Furthermore, a PS relative, SPPLP2, requires substrate shedding yet does not require putative substrate binding partners (Martin et al., 2009).

How \( \gamma \)-secretase assembly occurs is a critical question in \( \gamma \)-secretase biology. One model posits that PS binds initially to Aph1 and NCT to form a subcomplex, and then this subcomplex binds to Pen2 to initiate PS endoproteolysis and NCT glycosylation (Takasugi et al., 2003). Alternatively, PS and Pen2 form a subcomplex in which PS endoproteolysis occurs while NCT and Aph1 form another subcomplex. The two subcomplexes then bind together (Spasic and Annaert, 2008). In both models, \( \gamma \)-secretase activity is conferred only after the complete assembly of the four-protein complex. The observations reported here suggest that a trimeric complex containing PS1, Pen2, and Aph1a is active but unstable and perhaps rapidly converted to the four-protein complex in wild-type cells. Whether such a subcomplex

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**Figure 6.** Pen2 and Aph1a are involved in the \( \gamma \)-secretase activity in the absence of NCT. Representative Western blots of NICD levels in Pen2 and Aph1 siRNA-transfected NCT-deficient cells are shown. A, Lysates from control (Ctr) siRNA- or Pen2 siRNA-transfected NCT-deficient cells were analyzed by Western blotting. Pen2 siRNA markedly decreased PS1-NTF level but not that of full-length PS1 (FL-PS1). Whereas Pen2 depletion slightly affected N1,\( \Delta \)E-6Myc expression, it significantly diminished NICD generation in a dose-dependent manner. Note that the amounts of a PS1 fragment (\( \approx 30 \) kDa) increased in Pen2 siRNA-treated samples. This PS1 fragment does not have \( \gamma \)-secretase activity. B, The efficiency of Pen2 knockdown (\( \approx 90 \% \)) was confirmed by qRT-PCR. C, Depletion of Aph1a in NCT-deficient cells markedly decreased PS1-NTF and NICD generation, whereas Aph1b siRNA treatment did not significantly affect NICD levels. D, Knockdown of Aph1b was confirmed by qRT-PCR.
occurs in the presence of NCT or only forms in its absence remains to be determined.

Why PS proteins require additional stabilizing components whereas SPP and SPLP do not and why PS2 is unable to act without NCT are interesting puzzles still to be resolved. The fact that γ-secretase activity could not be detected in PS-DRCO cells (Figs. 1, 3, and 4) and that PS1 siRNA greatly diminished the residual γ-secretase activity in NCT−/− cells (Fig. 5a; supplemental Figs. 2 and 3, available at www.jneurosci.org as supplemental material for sequences) confirms the centrality of PS for γ-secretase activity. Interestingly, although PS possesses both substrate-bonding and catalytic sites and thus has the potential to act as an enzyme on its own, knockdown of Pen2 and Aph1a eliminated γ-secretase activity in NCT−/− cells (Fig. 6). Despite establishing that PS requires these two proteins, the current data cannot distinguish whether Pen2 and Aph1 are required for stabilizing PS1 or for enhancing its catalytic activity. If Aph1 or Pen2 are required only to stabilize PS1, it is conceivable that other unstable trimericγ-secretase complexes can be formed in vitro. It is intriguing to suggest that isolation of such an active trimeric γ-secretase might facilitate the elucidation of its structure. It also appears that active Hop-1 trimeric γ-secretase complexes may exist in vivo and can be isolated from Aph-2 mutants.

In conclusion, the present data demonstrate that NCT is dispensable for PS1-containing γ-secretase activity and cell surface transport but critical for stabilizing γ-secretase. This study also provides a biochemical explanation for the differential requirement for Aph2 shown by the two C. elegans PS proteins, Hop-1 and Sel-12.

References


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