



Development of a Reporter System for *In Vivo* Monitoring of γ -Secretase Activity in *Drosophila*

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<http://dx.doi.org/10.14348/molcells.2017.2294>

www.molcells.org

The γ -secretase complex represents an evolutionarily conserved family of transmembrane aspartyl proteases that cleave numerous type-I membrane proteins, including the β -amyloid precursor protein (APP) and the receptor Notch. All known rare mutations in APP and the γ -secretase catalytic component, presenilin, which lead to increased amyloid β -peptide production, are responsible for early-onset familial Alzheimer's disease. β -amyloid protein precursor-like (APPL) is the *Drosophila* ortholog of human APP. Here, we created Notch- and APPL-based *Drosophila* reporter systems for *in vivo* monitoring of γ -secretase activity. Ectopic expression of the Notch- and APPL-based chimeric reporters in wings results in vein truncation phenotypes. Reporter-mediated vein truncation phenotypes are enhanced by the *Notch* gain-of-function allele and suppressed by RNAi-mediated knockdown of *presenilin*. Furthermore, we find that apoptosis partly contributes to the vein truncation phenotypes of the APPL-based reporter, but not to the vein truncation phenotypes of the Notch-based reporter. Taken together, these results suggest that both *in vivo* reporter systems provide a powerful genetic tool to identify genes that modulate γ -secretase activity and/or APPL metabolism.

Keywords: γ -secretase, Alzheimer's disease, APPL, Notch, presenilin

INTRODUCTION

The γ -secretase complex, which consists of at least four proteins—Presenilin (PS), Nicastrin, Aph-1, and Pen-2, serves as a transmembrane aspartyl protease that plays a critical role in Alzheimer's disease (AD) and Notch signaling pathway (Bai et al., 2015; De Strooper, 2003; Fortini, 2009). Amyloid β -peptides (A β s), which are the main constituents of senile plaques present in the brain affected with AD, are generated by sequential cleavages of the β -amyloid precursor protein (APP) by β -secretase (BACE) and γ -secretase (Esler and Wolfe, 2001; Goedert, 2015). Another well-known substrate for γ -secretase is Notch, whose signaling controls a large number of cell fate decisions during development (Fortini, 2009; Louvi and Artavanis-Tsakonas, 2006). The cleavage of both APP and Notch within their single-pass transmembrane domains by γ -secretase releases cytosolic fragments and allows them to enter the nucleus, thereby regulating gene transcription (Brown et al., 2000; Cao and Südhof, 2001; Louvi and Artavanis-Tsakonas, 2006; Wang et al., 2014; Figs. 1A and 1B). This γ -secretase-dependent regulated intramembrane proteolysis (RIP) is tightly controlled (Brown et al., 2000), and thus alterations in the RIP of APP and Notch result in developmental defects and diseases (Esler and Wolfe, 2001; Louvi and Artavanis-Tsakonas, 2006).

Received 2 December, 2016; revised 25 December, 2016; accepted 28 December, 2016; published online 26 January, 2017

eISSN: 0219-1032

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AD is the most common form of senile dementia characterized by the deposition of aggregation-prone A β s and neurofibrillary tangles in the brain (Selkoe, 1998). Interestingly, early-onset familial Alzheimer's disease (EOAD) is caused by rare mutations in *APP*, *PS1*, and *PS2* (Tanzi, 2012). This is consistent with the fact that PS is the catalytic subunit of the γ -secretase complex whereas APP is a substrate for γ -secretase (De Strooper, 2003). In addition, γ -secretase activity and A β production were not detected in PS1/PS2 double-knockout cells (Herreman et al., 2000; Zhang et al., 2000). Therefore, γ -secretase serves as a therapeutic target to reduce A β production and development of γ -secretase inhibitors can provide an effective therapy for AD. However, there is still one major concern that γ -secretase inhibitors might cause deleterious side effects because they also affect the RIP of numerous other substrates including Notch (Esler and Wolfe, 2001; McCarthy et al., 2009). One possible approach to overcome this problem is to identify new therapeutic target genes that specifically affect the γ -secretase-dependent processing of APP rather than other substrates. To this end, we built a *Drosophila* reporter system for *in vivo* detection of γ -secretase activity by taking advantage of the Notch signaling pathway in the wing.

Canonical Notch signaling, which is highly conserved in the animal kingdom, is initiated by the binding of ligands including Delta, Serrate, and LAG-2 (Fortini, 2009; Louvi and Artavanis-Tsakonas, 2006). Ligand binding induces sequential cleavages of the Notch receptor by ADAM family metalloproteases and γ -secretase, releasing the Notch intracellular domain (NICD) (Fortini, 2009; Louvi and Artavanis-Tsakonas, 2006). NICD then enters the nucleus and forms an active transcriptional complex with the DNA-binding protein Suppressor of Hairless (Su(H)) and Mastermind (MAM), triggering expression of Notch target genes (Fortini, 2009; Louvi and Artavanis-Tsakonas, 2006; Fig. 1B). In the absence of Notch signaling, Su(H) proteins associate with various corepressors to actively repress the transcription of Notch target genes (Fortini, 2009; Louvi and Artavanis-Tsakonas, 2006; Fig. 1B). These findings demonstrate that Su(H) is the key effector of Notch signaling.

The *Drosophila* homolog of human APP is known as β -amyloid protein precursor-like (APPL) (Martin-Morris and White, 1990). Flies homozygous for *App1* deletion allele showed a fast phototaxis defect that can be restored by human APP as well as fly APPL, revealing their functional conservation (Luo et al., 1992). Furthermore, APPL is proteolytically cleaved by a BACE-like secretase (dBACE) and γ -secretase, producing a neurotoxic A β -like fragment in *Drosophila* (Carmine-Simmen et al., 2009). However, dBACE does not appear to recognize and cleave human APP at the β -secretase (BACE) cleavage site, suggesting the lack of a clear homolog of human BACE in *Drosophila* (Carmine-Simmen et al., 2009; Greeve et al., 2004). In contrast, each component and the activity of the γ -secretase complex appear to be evolutionarily conserved across invertebrates and vertebrates (De Strooper, 2003; Selkoe and Wolfe, 2007). In support of this claim, *Drosophila* γ -secretase can cleave human APP and thus produce A β peptides in combination with human BACE (Greeve et al., 2004).

Here, we have created a *Drosophila* reporter system for *in vivo* monitoring of γ -secretase activity and it provides a genetic tool to discover novel genes that modulate γ -secretase-mediated RIP of Notch and APP-like (APPL), and APPL metabolism.

MATERIALS AND METHODS

Drosophila strains

The *w¹¹¹⁸* *Drosophila* strain was used as a wild-type control. The following flies were obtained from the Bloomington Stock Center: *UAS-psn RNAi* (#27681), *UAS-p35* (#5073), *UAS-psn* (#8310), *e16E-GAL4* (#30557), and *N^{A^wE2}* (#51660). The *e16E-GAL4* enhancer trap line was shown to drive expression of a reporter gene in the striped pattern of *engrailed* during embryonic development (Weiss et al., 2001) and also in the posterior compartment of the wing imaginal disc during third instar larval stage (Johnson et al., 1995).

Expression constructs

The UAS-NTM-SV and UAS-APPL-SV constructs were generated by PCR and/or restriction enzyme-based strategies and subcloned into the pUAST vector (Brand and Perrimon, 1993). For the Litmus 28-Su(H)-VP16 (SV), a NcoI-PstI fragment from pGEX-Su(H) (Bailey and Posakony, 1995), which contains the DNA-binding domain (amino acids 109-457; GenBank: AAF53434.1) of Su(H), and a PstI-XbaI fragment from pAct-GAL4-VP16 (Han and Manley, 1993), which contains the transcriptional activation domain (amino acids 2-79; GenBank: AIZ65950.1) of VP16, were subcloned into the Litmus 28 vector (New England Biolabs, Inc.). For the Litmus 28-NTM-SV, a Sall-NcoI fragment from a *Notch* minigene clone (Wharton et al., 1985), which contains the NTM (amino acids 1710-1891; GenBank: AAF45848.2), and a BgIII-Sall fragment from Litmus 28-N^{AECN} (Ju et al., 2000), which contains the signal peptide region (amino acids 1-60) of Notch, were subcloned into the Litmus 28-SV. For the UAS-NTM-SV, a BgIII-XbaI fragment from the Litmus 28-NTM-SV was subcloned into the pUAST vector. For the UAS-APPL-SV construct, a EcoRI-NcoI fragment from GH04413, which encodes the full-length APPL (amino acids 1-887; GenBank: AAF45520), and a NcoI-XbaI fragment from the Limus 28-SV, were subcloned into the pUAST vector. These construct sequences were confirmed by DNA sequencing. Transgenic flies carrying these constructs were created according to standard *D. melanogaster* transformation procedures (KAIST transgenic fly service). Multiple independent lines for the same transgenic construct are indicated by superscripts. For example, *NTM-SV^{61f}* and *NTM-SV^{A2m}* are independent insertion lines for the construct UAS-NTM-SV.

Immunohistochemistry

Expression patterns of transgenes were visualized using anti-VP16 monoclonal antibody (14-5, Santa Cruz Biotech.) and 3,3'-diaminobenzidine reaction, as described previously (Jeong et al., 2012; Kim et al., 1995).

Preparation of *Drosophila* wings and phenotypic characterization

Using two pairs of forceps, the wings of female and male

adult flies (younger than 3 days) were carefully cut off and subsequently were arranged in the same orientation on a glass slide. A coverslip was applied and each corner of the coverslip was sealed with a regular nail polish. In wild-type wing, the average L4 to L3 vein length ratio was 0.97 and the average L5 to L3 vein length ratio was 0.58 (see Supplementary Fig. S1A). The average percentages of L4 and L5 vein truncations were calculated by the following equations (see Supplementary Fig. S1B):

$$\text{L4 vein truncation} = [(0.97x-y)/(0.97x)] \times 100 (\%)$$

$$\text{L5 vein truncation} = [(0.58x-z)/(0.58x)] \times 100 (\%)$$

$$x = \text{L3 vein length}$$

$$y = \text{L4 vein length}$$

$$z = \text{L5 vein length}$$

Statistical analysis

All quantitative data are presented as mean \pm standard error of the mean (S.E.M.). Statistical significance was determined using unpaired *t*-test ($*p < 0.05$, $**p < 0.01$, and $***p < 0.001$).

RESULTS

Generation of transgenic reporters for *in vivo* detection of the processing of Notch and APPL

We decided to take advantage of the Notch signaling pathway to generate transgenic reporters for *in vivo* detection of the activity of γ -secretase, which acts as a key player in Alzheimer's disease (AD) (Esler and Wolfe, 2001; Selkoe, 1998). Notch is cleaved within its single-pass transmembrane domain by γ -secretase (Fortini, 2009; Louvi and Artavanis-Tsakonas, 2006). The cleavage of the Notch receptor releases its intracellular domain (NICD) and translocates it into the nucleus, thereby driving the expression of Notch target genes through the assembly of an active transcriptional complex with Su(H) and Mastermind (Fortini, 2009; Louvi and Artavanis-Tsakonas, 2006; Figs. 1A and 1B). Vein pattern formation in the *Drosophila* wing is controlled by Notch signaling (de Cellis, 1998). Reduction in Notch signaling activity leads to the formation of thicker veins, whereas increased Notch activity results in loss of veins (de Cellis and Garcia-Bellido, 1994a). These observations prompted us to examine whether Notch activity is correlated with vein phenotypes in a dose-dependent manner. Normal vein patterning was observed in wild-type females (Fig. 2A), whereas in females heterozygous for N^{Ax-E2} , a dominant gain-of-function (GOF) *Abruptex* (*Ax*) allele (de Cellis and Garcia-Bellido, 1994b), vein L5 consistently failed to reach the wing margin (5.4% L5 vein truncation; Figs. 2B and 2I). This *Ax* phenotype of vein L5 was significantly increased up to 26.1% vein truncation in females homozygous for N^{Ax-E2} ($p = 0.002$, *t*-test; Figs. 2C and 2I). In addition, these homozygous mutants displayed a moderately truncated L4 vein (4.8% vein truncation; Figs. 2C and 2I). These findings suggest that the severity of *Ax* vein phenotype is proportional to Notch signaling activity. To further address whether γ -secretase activity is required for patterning of the *Drosophila* wing veins, we took advantage of the RNA interference

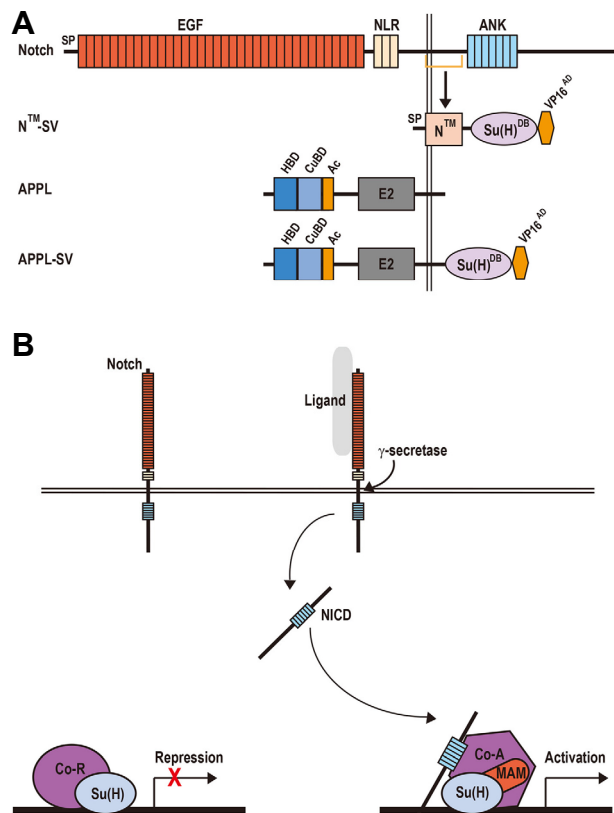


Fig. 1. Generation of the Notch- and APPL-based Fusion Reporters. (A) Schematic diagram of the Notch- and APPL-based reporter transgenes. SP, signal peptide; EGF, epidermal growth factor-like; LNR, Lin12-Notch repeats; ANK, seven ankyrin repeat domains; NICD, Notch intracellular domain; Su(H)^{DB}, the DNA-binding domain of Su(H); VP16^{AD}, the transcriptional activation domain of the virus VP16; HBD, heparin-binding domain; CuBD, copper-binding domain; Ac, acidic domain; E2, APP extracellular carbohydrate domain. (B) Transcriptional switch in the *Drosophila* canonical Notch signaling pathway. In the absence of Notch signaling, Su(H) protein functions as a transcriptional repressor through recruiting co-repressors (Co-R) including Hairless, Groucho and CtBP. Notch activation is triggered by ligand binding and sequential cleavages of ADAM family metalloproteases and γ -secretase. The second cleavage liberates the NICD, which then enters the nucleus and associates with Su(H) and Mastermind (MAM) to drive the transcription of Notch target genes. Co-A, co-activators.

(RNAi) technique. When the *Drosophila presenilin* (*psn*) gene was knocked down by overexpression of *psn RNAi* transgene using *e16E-GAL4*, which is expressed in the posterior compartment of the wing (Johnson et al., 1995), wing patterns showed robust vein thickening but no vein truncation phenotypes (Figs. 2D and 2I). This vein thickening phenotype, which is reminiscent of the phenotype found in *Notch* loss-of-function alleles (de Cellis and Garcia-Bellido, 1994a), was completely restored by coexpression of wild-

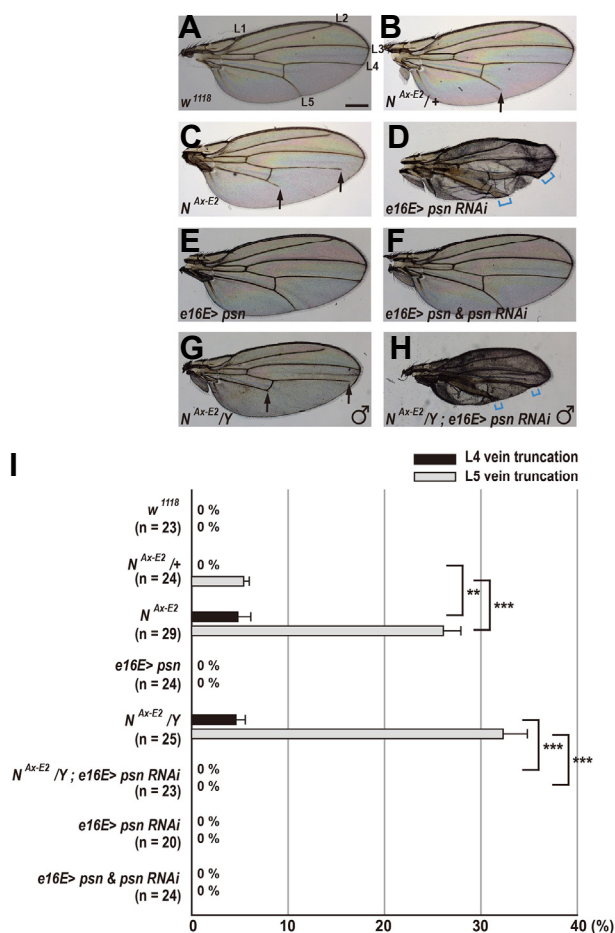


Fig. 2. Wing vein phenotypes of Notch gain-of-function and *presenilin* loss-of-function alleles. (A) Wild-type wing shows a normal pattern of five longitudinal veins (L1-L5). Scale bar indicates 50 μ m. (B) In females heterozygous for *N^{Ax-E2}*, the wing displays mild truncation only in L5 vein (arrow). (C) In females homozygous for *N^{Ax-E2}*, the wing displays mild truncation in L4 vein and moderate truncation in L5 vein (arrows). (D) Overexpression of *psn RNAi* in the posterior compartment results in vein thickening (square brackets in blue). (E) Female wing overexpressing wild-type Psn shows a normal vein pattern. (F) Overexpression of wild-type Psn completely rescues the *psn RNAi*-induced vein thickening phenotype. (G) In males hemizygous for *N^{Ax-E2}*, the wing exhibits mild truncation in L4 vein and moderate truncation in L5 vein (arrows). (H) Vein truncation phenotype observed in *N^{Ax-E2}* hemizygotes is suppressed by RNAi-mediated knockdown of *psn*. Square brackets indicate vein thickening phenotype. (I) Percentages of wing vein truncations in females and males with indicated genotypes (** $p < 0.01$ and *** $p < 0.001$, *t*-test). Error bars indicate S.E.M. by *t*-test. n = number of wings scored for each genotype.

type Psn (Fig. 2F). Overexpression of wild-type Psn alone hardly affected vein patterning (Figs. 2E and 2I). These results indicate target gene specificity of the *psn RNAi*

transgene. Furthermore, knockdown of *psn* in the posterior compartment using *e16E-GAL4* completely suppressed the vein truncation phenotype observed in males hemizygous for *N^{Ax-E2}* (Figs. 2G and 2H). Taken together, these findings strongly suggest that γ -secretase activity contributes to vein pattern formation in the *Drosophila* wing through the Notch signaling pathway.

To monitor the proteolytic cleavage of Notch and APPL by γ -secretase within their transmembrane domains, we explored a sensitive reporter system, in which the chimeric transcriptional activator Su(H)-VP16 (SV) is fused to either a subfragment of Notch (*NTM*) or full-length APPL (*UAS-NTM-SV* and *UAS-APPL-SV* in Fig. 1A). The *NTM* subfragment possesses a transmembrane domain and the N-terminal region of NICD (Fig. 1A). The SV was created by fusion of the DNA-binding domain of Su(H) transcription factor (Bailey and Posakony, 1995) and the transcriptional activation domain of the virus VP16 (Sadowski et al., 1988) (Fig. 1A). Therefore, in case of nuclear translocation of the SV proteins, they are supposed to activate some of the Notch target genes, mimicking increased Notch pathway activity. Increased Notch activity in the wing can be visualized by the severity of the resulting vein phenotypes. Based on the fact that the *NTM* subfragment and APPL contain a γ -secretase-mediated cleavage site, we reasoned that the expression of both reporter transgenes under the control of *e16E-GAL4* provides a sensitive assay for γ -secretase-dependent processing of Notch and APPL.

Expression of the notch- and APPL-based chimeric reporters in wings caused loss of wing veins

To determine whether both the *NTM-SV* and APPL-SV chimeric reporter proteins transduce Notch signaling activity, we ectopically overexpressed these transgenes using *e16E-GAL4* (Figs. 3E and 3H). Ectopic expression of *NTM-SV^{61f}* in the posterior compartment of the wing resulted in moderate truncations of L4 and L5 veins in females (Fig. 3D). These vein truncation phenotypes were not observed in wild-type, *e16E-GAL4/+*, and *NTM-SV^{61f}/+* female wings (Figs. 3A-3C). Interestingly, in females expressing APPL-SV^{85m}, we observed severe truncations of L4 and L5 veins (Fig. 3G). However, vein pattern formation in females heterozygous for the APPL-SV^{85m} transgene was normal (Fig. 3F). These findings might suggest that both the *NTM-SV* and APPL-SV reporter proteins were processed at least by γ -secretase, and they further demonstrate that the γ -secretase-cleaved C-terminal fragments of both reporters including the SV can transduce Notch activity in the wing.

To further address the difference in phenotypic severity between *NTM-SV* and APPL-SV (Figs. 3D and 3G), we obtained a total of 5 independent transgenic lines for *NTM-SV* and a total of 4 lines for APPL-SV. On average, ectopic expression in 5 different lines for *NTM-SV* caused 43.1% L4 vein truncations and 43.8% L5 vein truncations (n = 84), while expression in 4 different lines for APPL-SV resulted in 90.4% L4 vein truncations and 96.8% L5 vein truncations (n = 62) (Fig. 3I). An almost 2-fold difference in the percentage of both vein truncations between *NTM-SV* and APPL-SV ($p < 0.001$, *t*-test; Fig. 3I) strongly suggests that the *NTM-SV* and APPL-SV

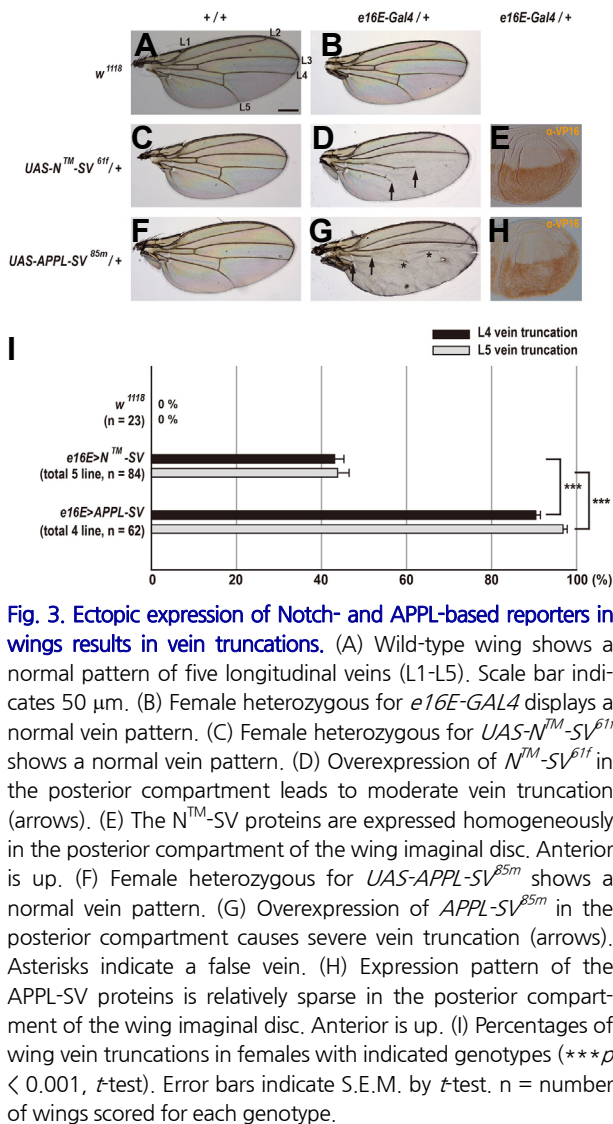


Fig. 3. Ectopic expression of Notch- and APPL-based reporters in wings results in vein truncations. (A) Wild-type wing shows a normal pattern of five longitudinal veins (L1-L5). Scale bar indicates 50 μ m. (B) Female heterozygous for *e16E-GAL4* displays a normal vein pattern. (C) Female heterozygous for *UAS-NTM-SV^{61f}* shows a normal vein pattern. (D) Overexpression of *NTM-SV^{61f}* in the posterior compartment leads to moderate vein truncation (arrows). (E) The *NTM-SV* proteins are expressed homogeneously in the posterior compartment of the wing imaginal disc. Anterior is up. (F) Female heterozygous for *UAS-APPL-SV^{85m}* shows a normal vein pattern. (G) Overexpression of *APPL-SV^{85m}* in the posterior compartment causes severe vein truncation (arrows). Asterisks indicate a false vein. (H) Expression pattern of the APPL-SV proteins is relatively sparse in the posterior compartment of the wing imaginal disc. Anterior is up. (I) Percentages of wing vein truncations in females with indicated genotypes ($***p < 0.001$, *t*-test). Error bars indicate S.E.M. by *t*-test. n = number of wings scored for each genotype.

proteins are differentially processed by γ -secretase and/or the γ -secretase-cleaved C-terminal portions of both reporter proteins generate different levels of Notch transducing activity. Another possibility is that the APPL-SV proteins are more abundantly expressed than the *NTM-SV* proteins.

Reporter-mediated wing vein phenotypes are enhanced by Notch and suppressed by presenilin

Based on wing vein phenotypes, we deduced that the *NTM-SV* and APPL-SV reporter proteins transduce Notch activity (Fig. 3). To further address this issue, we investigated genetic interactions among *NTM-SV*, *Notch*, and *psn*. Ectopic expression of *NTM-SV^{61f}* in the posterior compartment resulted in 38.3% L4 vein truncations and 33.6% L5 vein truncations (Figs. 4B and 4G). These *NTM-SV^{61f}* vein phenotypes were significantly enhanced by one copy of *N^{Ax-E2}* allele: from 38.3% to 69.2% for L4 vein truncations ($p < 0.001$, *t*-test)

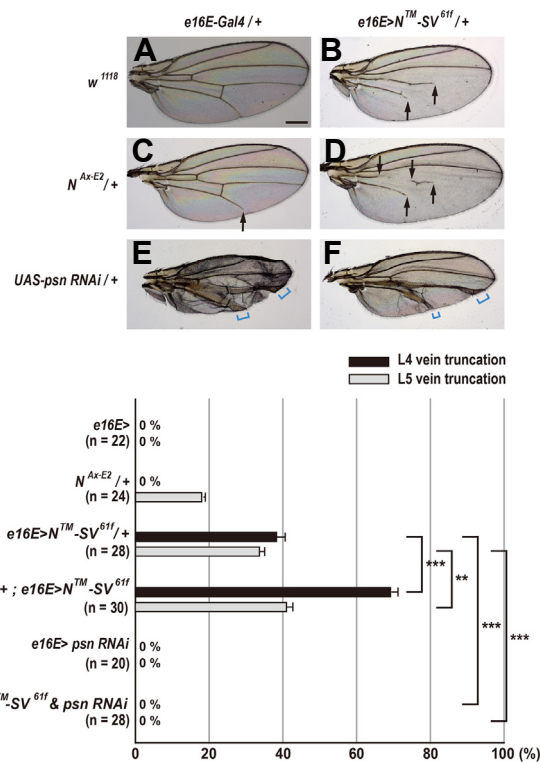


Fig. 4. Genetic interactions among *NTM-SV^{61f}*, Notch, and presenilin alleles. (A) Females heterozygous for *e16E-GAL4* show a normal pattern of five longitudinal veins. Scale bar indicates 50 μ m. (B) Overexpression of *NTM-SV^{61f}* in the posterior compartment results in moderate vein truncation (arrows). (C) In females heterozygous for *N^{Ax-E2}*, the wing shows mild truncation only in L5 vein (arrow). (D) One copy of *N^{Ax-E2}* mutation slightly enhances the gain-of-function (GOF) phenotype of *NTM-SV^{61f}* (arrows). (E) Overexpression of *psn RNAi* in the posterior compartment results in vein thickening (square brackets). (F) Overexpression of *psn RNAi* completely suppresses the GOF phenotype of *NTM-SV^{61f}*. Square brackets indicate vein thickening phenotype. (G) Percentages of wing vein truncations in females with indicated genotypes ($**p < 0.01$ and $***p < 0.001$, *t*-test). Error bars indicate S.E.M. by *t*-test. n = number of wings scored for each genotype.

and from 33.6% to 41.0% for L5 vein truncations ($p = 0.001$, *t*-test) (Figs. 4C, 4D, and 4G). Similar patterns of genetic interactions were also observed with a different insertion line, *NTM-SV^{A2m}* ($p < 0.001$, *t*-test for both veins; Supplementary Figs. S2C, S2D, and S2G). Overexpression of *NTM-SV^{A2m}* in the wing led to 57.0% L4 vein truncations and 57.7% L5 vein truncations, which are greater than those obtained after ectopic expression of *NTM-SV^{61f}* (Supplementary Figs. S2B and S2G). These findings support the claim that the *NTM-SV* proteins transduce Notch activity in the wing. In contrast, the *NTM-SV^{61f}* and *NTM-SV^{A2m}* vein truncation phenotypes were completely suppressed by RNAi-mediated knockdown of *psn* (Figs. 4E-4G and Supplementary Figs.

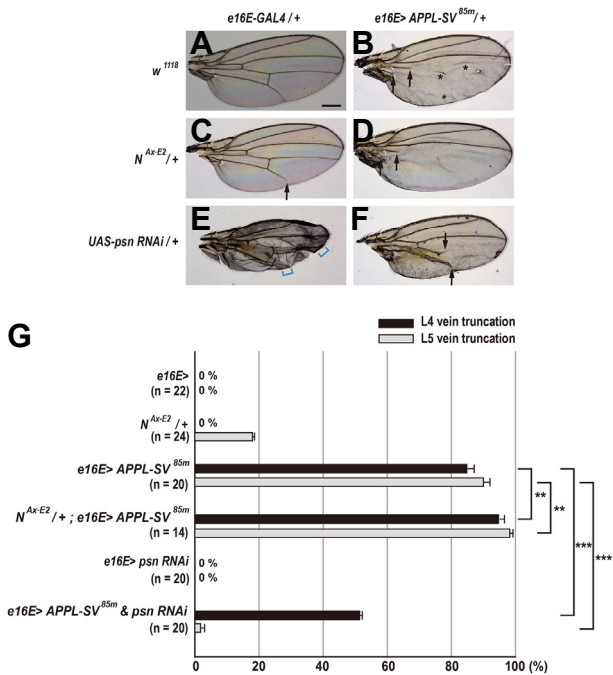


Fig. 5. Genetic interactions among *APPL-SV^{85m}*, Notch, and *pre-senilin* alleles. (A) Females heterozygous for e16E-Gal4 show a normal pattern of five longitudinal veins. Scale bar indicates 50 μ m. (B) Overexpression of *APPL-SV^{85m}* in the posterior compartment results in severe vein truncation (arrows). Asterisks indicate a false vein. (C) In females heterozygous for *N^{Ax-E2}*, the wing shows mild truncation only in L5 vein (arrow). (D) One copy of *N^{Ax-E2}* mutation slightly enhances the GOF phenotype of *APPL-SV^{85m}* (arrows). (E) Overexpression of *psn RNAi* in the posterior compartment results in vein thickening (square brackets). (F) Overexpression of *psn RNAi* partly suppresses the GOF vein phenotype of *NTM-SV^{61f}*. (G) Percentages of wing vein truncations in females with indicated genotypes (** $p < 0.01$ and *** $p < 0.001$, *t*-test). Error bars indicate S.E.M. by *t*-test. n = number of wings scored for each genotype.

S2E-S2G), indicating that Psn is required for the gain of NTM-SV function in the wing. Interestingly, the *psn* RNAi-mediated vein thickening phenotype was also suppressed, but it was not completely suppressed by overexpression in either *NTM-SV* transgenic line (Figs. 4E, 4F, Supplementary Figs. S2E and S2F). Therefore, knockdown of *psn* is epistatic to NTM-SV overexpression with respect to vein pattern formation in the wing.

We next analyzed genetic interactions among *APPL-SV*, *Notch*, and *psn*. Females overexpressing *APPL-SV^{85m}* in the posterior compartment showed highly penetrant vein truncation defects: 84.8% L4 vein truncations and 90.0% L5 vein truncations (Figs. 5B and 5G). These *APPL-SV^{85m}* vein phenotypes were enhanced by one copy of *N^{Ax-E2}* allele: from 84.8% to 94.7% for L4 vein truncations ($p = 0.001$, *t*-test) and from 90.0% to 98.2% for L5 vein truncations ($p = 0.003$, *t*-test), suggesting that the *APPL-SV* proteins can act as a Notch signal transducer in the wing (Figs. 5C, 5D, and

5G). Similar patterns of genetic interactions for both L4 and L5 vein phenotypes were not observed with the *APPL-SV^{82m}* transgene, a different insertion line ($p > 0.3$, *t*-test; Supplementary Figs. S3C, S3D, and S3G), probably due to high penetrance of vein truncation defects in females overexpressing *APPL-SV^{82m}* alone (92.8% for L4 veins and 100% for L5 veins; Supplementary Fig. S3G). In addition, we found that the *APPL-SV^{85m}* vein truncation phenotypes were robustly suppressed by RNAi-mediated knockdown of *psn*: from 84.8% to 51.2% for L4 vein truncations ($p < 0.001$, *t*-test) and from 90.0% to 1.6% for L5 vein truncations ($p < 0.001$, *t*-test) (Figs. 5E-5G). Partial suppression especially in L4 veins, which is different from that observed in females coexpressing *psn RNAi* and *NTM-SV* (Figs. 4F and 4G), might indicate that *APPL-SV*-induced Notch activity antagonizes the knockdown effects of *psn* to a greater extent than *NTM-SV*. Furthermore, similar patterns of suppressive genetic interactions were also observed with the *APPL-SV^{82m}* transgene ($p < 0.001$, *t*-test for both L4 and L5 vein truncations; Supplementary Figs. S3E-S3G), demonstrating the requirement of Psn for the gain of *APPL-SV* function in the wing. Females coexpressing *psn RNAi* and *APPL-SV* exhibited large reduction in vein thickening phenotype compared to females overexpressing *psn RNAi* alone (Figs. 5E and 5F, Supplementary S3E and S3F), which is indicative of the antagonistic relationships between knockdown of *psn* and *APPL-SV* overexpression in patterning of the wing veins.

The role of apoptosis in reporter-induced wing vein truncations

Since Notch plays a role in apoptosis during *Drosophila* visual system development (Bertet et al., 2014; Brachmann and Cagan, 2003), we investigated whether Notch-mediated regulation of apoptosis is involved in patterning of the wing veins. Interestingly, females overexpressing the baculovirus pan-caspase inhibitor p35, which has been shown to prevent apoptotic death in *Drosophila* (Hay et al., 1994), exhibited a mild phenotype only in L5 vein, indicating a positive, but a minor role of apoptosis in vein pattern formation (3.5% vein truncation; Figs. 6D and 6G). In females expressing both *p35* and *NTM-SV^{61f}* in the posterior compartment, we observed a small but significant increase in vein phenotypes compared to females overexpressing *NTM-SV^{61f}*: from 38.3% to 51.1% for L4 vein truncations ($p < 0.001$, *t*-test) and from 33.6% to 46.1% for L5 vein truncations ($p < 0.001$, *t*-test) (Figs. 6B, 6E and 6G). Similar enhancement in vein phenotype was also found in females coexpressing *p35* and *NTM-SV^{42m}* in the posterior compartment: from 57.0% to 60.5% for L4 vein truncations ($p = 0.03$, *t*-test) and from 57.7% to 76.6% for L5 vein truncations ($p < 0.001$, *t*-test) (Supplementary Figs. S4B, S4E, and S4G). Taken together, this seemingly additive genetic interaction between *p35* and *NTM-SV* suggests that NTM-SV-induced vein truncation is not due to apoptosis.

To examine whether *APPL-SV*-induced vein phenotypes result from apoptosis, we coexpressed *p35* and *APPL-SV^{85m}* in the posterior compartment of the wing. Unexpectedly, we observed that coexpression of *p35* and *APPL-SV^{85m}* robustly suppressed L4 vein truncation phenotypes of *APPL-SV^{85m}*

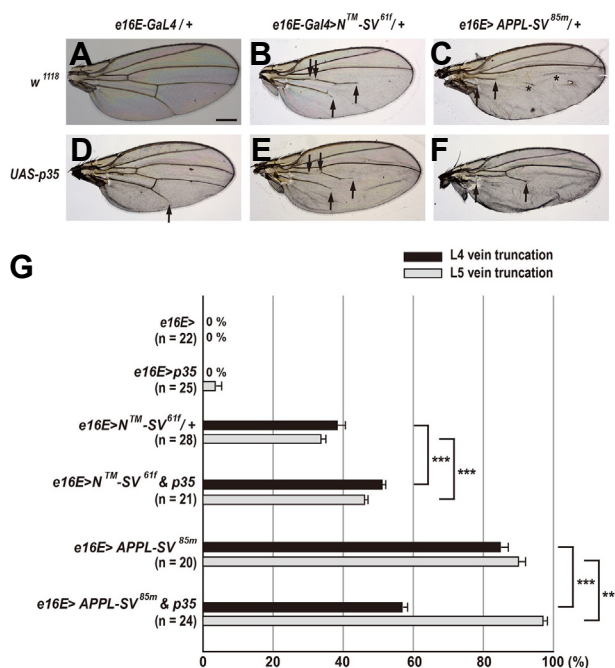


Fig. 6. Apoptosis partly contributes to vein truncations induced by the APPL-based reporter. (A) Females heterozygous for e16E-Gal4 show a normal pattern of five longitudinal veins. Scale bar indicates 50 μ m. (B) Overexpression of NTM-SV^{61f} in the posterior compartment results in moderate vein truncation (arrows). (C) Overexpression of APPL-SV^{85m} in the posterior compartment leads to severe vein truncation (arrows). Asterisks indicate a false vein. (D) Overexpression of p35 in the posterior compartment results in slight vein truncation (arrow). (E) Coexpression of p35 in the posterior compartment slightly increases the GOF vein phenotype of NTM-SV^{61f} (arrows). (F) Coexpression of p35 in the posterior compartment differentially affects the GOF vein phenotype of APPL-SV^{85m} (arrows). (G) Percentages of wing vein truncations in females with indicated genotypes (** $p < 0.01$ and *** $p < 0.001$, *t*-test). Error bars indicate S.E.M. by *t*-test. n = number of wings scored for each genotype.

from 84.8% to 56.8% ($p < 0.001$, *t*-test), whereas coexpression of p35 and APPL-SV^{85m} enhanced L5 vein truncation phenotypes from 90.0% to 97.0% ($p = 0.003$, *t*-test) (Figs. 6C, 6F, and 6G). Suppressive genetic interactions were also observed in L4 veins, but not in L5 veins with the APPL-SV^{85m} transgene: from 92.8% to 54.8% for L4 vein truncations ($p < 0.001$, *t*-test) and no change was observed in L5 vein truncations (Supplementary Figs. S4C, S4F, and S4G). Strong suppressive interactions between p35 and APPL-SV, especially in L4 veins suggest that overexpression of APPL-SV leads to cell death, and they further demonstrate that APPL-SV-induced cell death partly contributes to L4 vein truncation phenotypes. However, phenotypic enhancement observed in L5 veins, which is similar to that observed in females coexpressing p35 and NTM-SV (Fig. 6G), suggests that apoptosis is not attributable to L5 vein truncation phenotypes of APPL-SV.

DISCUSSION

Drosophila melanogaster provides a powerful genetic tool to analyze gene function associated with human diseases including Alzheimer's disease (AD) and evolutionarily conserved signaling pathways (Bier, 2005; Lu and Vogel, 2009). Indeed, many different *Drosophila* models of AD, which showed progressive degeneration through elevated production of amyloid β -peptides (A β s), were developed (Fernandez-Funez et al., 2015). Some of these AD models are based on overexpression of wild-type human APP, EOAD-associated mutant APP, or a secreted form of human A β 42 peptide (Fernandez-Funez et al., 2015). These findings demonstrate highly conserved mechanisms underlying A β -mediated neurotoxicity and neurodegeneration between *Drosophila* and humans. This is further supported by the fact that the fly genome contains functional homologs for most of the EOAD-associated genes including APP, PS1, and PS2 (Martin-Morris and White, 1990; Stempfle et al., 2010). In combination with human BACE, *Drosophila* γ -secretase is able to cleave human APP to generate A β peptides, indicating its conserved intramembrane proteolytic activity (Greeve et al., 2004). Although the A β sequence is not well conserved among APP homologs from different species, a growing body of evidence suggests that all APP homologs play an important and conserved role in neural development including neurite growth, axon guidance, and synaptogenesis (Nicolas and Hassan, 2014). However, little is known about how γ -secretase-mediated proteolytic processing of APP is regulated at the molecular and genetic levels.

One of the hallmarks of AD is senile plaques largely consisting of A β s, which are produced by sequential cleavages of APP by β -secretase (BACE) and γ -secretase (Esler and Wolfe, 2001; Goedert, 2015). Given that all known mutations responsible for early-onset familial Alzheimer's disease (EOAD) are localized in APP, PS1, and PS2 (Tanzi, 2012), γ -secretase represents a good therapeutic target for developing A β -lowering drugs. Therefore, brain-penetrant inhibitors of γ -secretase are emerging as one of the most effective therapies for AD (De Strooper et al., 2010). However, toxicity of γ -secretase inhibitors was found to be a major hurdle in preclinical studies since γ -secretase is also required for the RIP of numerous other substrates including Notch (De Strooper et al., 2010; Esler and Wolfe, 2001; McCarthy et al., 2009). Here, as an alternative approach to overcome this target-based toxicity, we report the development of a *Drosophila* reporter system for *in vivo* detection of γ -secretase activity. Based on several observations, we expect that this *in vivo* reporter system will provide a genetic tool to discover novel genes, which specifically modulate γ -secretase-mediated processing of Notch and APPL. First, overexpression of the NTM-SV reporter proteins in the wing induced vein truncation phenotypes that are totally dependent on γ -secretase activity. Second, not only γ -secretase activity, but also apoptosis contributes to the vein truncation phenotypes of the APPL-SV reporter transgene. This appears to be consistent with previous observations suggesting that human APP induces APP intracellular domain (AICD)-dependent cell death in *Drosophila* (Gunawardena and Goldstein, 2001; Wang et al.,

2014). Since the γ -secretase-cleaved C-terminal fragment of APPL-SV contains the APPL intracellular domain (dAICD) in addition to the SV, dAICD might be responsible for the contribution of apoptosis. Third, both NTM-SV and APPL-SV transduce Notch signaling activity in a dose-dependent manner. In support of this claim, reporter-induced vein phenotypes are enhanced by one copy of *N^{Δx-ε2}*, which is a *Notch* GOF allele. Fourth, APPL-SV-induced phenotype is almost 2-fold greater than the GOF phenotype of NTM-SV, probably due to differences in their RIP and the contribution of dAICD-dependent apoptosis.

Several γ -secretase associated proteins (GSAPs), in addition to the four main components of the γ -secretase complex, have been discovered (Chen et al., 2006; He et al., 2010; Teranishi et al., 2015; Wakabayashi et al., 2009; Zhou et al., 2005). These GSAP proteins, including TMP21, pigeon homologue protein, and proton myo-inositol cotransporter, regulate substrate selectivity and A β production. Therefore, these GSAPs can serve as therapeutic targets for the treatment of AD. These observations might indicate the existence of unknown GSAP proteins regulating A β production without affecting proteolytic cleavage of other γ -secretase substrates including Notch. Future work using the *in vivo* reporter system developed in this study will determine whether known and additional unknown GSAP proteins are required for modulation of γ -secretase activity, and how substrate specificity of γ -secretase is regulated. Our current reporter system can also be applied to screening of chemical compound libraries.

Note: Supplementary information is available on the Molecules and Cells website (www.molcells.org).

ACKNOWLEDGMENTS

We thank Jaeseob Kim, Eunkyung Bae, and Jungmo Kim for their comments on the manuscript; James Posakony, Kyuhyung Han, Spyros Artavanis-Tsakonas, and the *Drosophila* Genomics Resource Center for clones; and the Bloomington *Drosophila* Stock Center for the fly strains. This study was supported by NRF-2013R1A1A4A01011329 (S.J.).

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