**Holo-APP and G-protein-mediated signaling are required for sAPPα-induced activation of the Akt survival pathway**

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Accumulating evidence indicates that loss of physiologic amyloid precursor protein (APP) function leads to reduced neuronal plasticity, diminished synaptic signaling and enhanced susceptibility of neurons to cellular stress during brain aging. Here we investigated the neuroprotective function of the soluble APP ectodomain sAPPα (soluble APPα), which is generated by cleavage of APP by α-secretase along the non-amyloidogenic pathway. Recombinant sAPPα protected primary hippocampal neurons and SH-SY5Y neuroblastoma cells from cell death induced by trophic factor deprivation. We show that this protective effect is abrogated in neurons from APP-knockout animals and APP-depleted SH-SY5Y cells, but not in APP-like protein 1- and 2- (APLP1 and APLP2) depleted cells, indicating that expression of membrane-bound holo-APP is required for sAPPα-dependent neuroprotection. Trophic factor deprivation diminished the activity of the Akt survival pathway. Strikingly, both recombinant sAPPα and the APP-E1 domain were able to stimulate Akt activity in wild-type (wt) fibroblasts, SH-SY5Y cells and neurons, but failed to rescue in APP-deficient neurons or fibroblasts. The ADAM10 (a disintegrin and metalloproteinase domain-containing protein 10) inhibitor GI254023X exacerbated neuron death in organotypic (hippocampal) slice cultures of wt mice subjected to trophic factor and glucose deprivation. This cell death-enhancing effect of GI254023X could be completely rescued by applying exogenous sAPPα. Interestingly, sAPPα-dependent Akt induction was unaffected in neurons of APP-ΔCT15 mice that lack the C-terminal YENPTY motif of the APP intracellular region. In contrast, sAPPα-dependent rescue of Akt activation was completely abolished in APP mutant cells lacking the G-protein interaction motif located in the APP C-terminus and by blocking G-protein-dependent signaling with pertussis toxin. Collectively, our data provide new mechanistic insights into the physiologic role of APP in antagonizing neurotoxic stress: they suggest that cell surface APP mediates sAPPα-induced neuroprotection via G-protein-coupled activation of the Akt pathway.

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(and the mammalian paralogs APP-like protein 1 and 2 (APLP1/APLP2)) are integral transmembrane proteins with proposed functions as membrane-bound signaling receptors and/or adhesion molecules.8–10 Despite the experimental evidence supporting the different functions of APP, it is currently unclear whether these functions are primarily mediated by transmembrane APP or by its soluble ectodomain shed from the cell surface. Based on existing evidence, it is also conceivable that membrane-anchored APP and secreted APP fragments share equal relevance in mediating the diverse physiologic roles of APP.

In vivo APP appears to be involved in damage responses and was found to be upregulated during brain injury in mammals and lower organisms.11–15 In line with this hypothesis, sAPPα was shown to exert protective effects following traumatic, ischemic and excitotoxic brain injury.16–20 There is also strong evidence supporting a neuroprotective function of APP and sAPPα in vitro21–31 but the cellular receptor and the intracellular downstream targets mediating these neuroprotective effects remain to be identified.32 Earlier work pointed to a role of sAPPα in the modulation of ion homeostasis and it was proposed that sAPPα-mediated neuroprotection may be associated both with rapid effects on ion channel function and with delayed transcription-dependent processes.33 sAPPα was proposed to activate potassium channels and suppress NMDA currents to limit Ca2+ overloading and excitotoxic damage in neurons.21,34,35 However, these studies also revealed that prolonged incubation with sAPPα over several hours was required to achieve significant effects on neuronal cell viability. Therefore, the neuroprotective function of APP/sAPPα is likely to be mediated by the prolonged activation of cell survival signaling pathways.32

We have previously demonstrated that APP/sAPPα can antagonize the activation of the c-Jun N-terminal kinase (JNK) pathway that represents a central stress signaling pathway and critical upstream modulator of the mitochondrial pathway of apoptosis.36 Furthermore, we and others have shown that the phosphatidylinoitride 3-kinase (PI3K)/Akt pathway is involved in mediating the protective function of APP.37–39 Of note, upstream JNK kinases such as mixed lineage kinase 3 and apoptosis signal-regulating kinase-1 (ASK1) can be inhibited via phosphorylation by Akt, suggesting crosstalk between stress and survival pathways modulated by APP.32 In addition to suppression of stress signaling, activation of survival pathways by sAPPα may also lead to the induction of prosurvival genes involved in stress responses and neuronal survival (manganese superoxide dismutase, peroxiredoxins, catalase).25,29 Despite these observations, the cellular receptor mediating the neuroprotective functions of sAPPα and the exact molecular mechanisms underlying sAPPα-dependent neuroprotective signaling have hitherto remained elusive.

Here we applied purified recombinant sAPPα in combination with a large set of knockdown (KD) and knockout (KO) models as well as APP mutants to further investigate the molecular mechanisms of sAPPα-induced neuroprotection. We found that transmembranous APP acts as an sAPPα receptor-mediating activation of the Akt survival pathway in a G-protein-dependent manner.

Results

Recombinant sAPPα and E1 promote cell survival only in the presence of holo-APP. As outlined above, it has been shown that sAPPα has neuroprotective properties under various stress conditions. To investigate the potential contribution of endogenous APP for sAPPα-induced neuroprotection, we performed a stable lentiviral KD of APP in human SH-SY5Y cells (Figure 1a). Cell viability/death was analyzed by quantification of ATP levels (Figures 1b and c) and microscopical evaluation of propidium iodide (PI) uptake (Figures 1d and e). Human insulin-like growth factor 1 (IGF1) served as a positive control for activation of cell survival. Recombinant, yeast-derived sAPPα and its subdomain E1 exerted dose-dependent antiapoptotic effects in wild-type (wt) human SH-SY5Y subjected to trophic factor withdrawal (Figures 1b and d). Both sAPPα and its subdomain E1 failed to antagonize stress-triggered cell death in cells lacking endogenous APP, suggesting that the expression of holo-APP may be necessary for the neuroprotective functions of exogenously applied sAPPα (Figures 1c and e). The requirement of endogenous APP for the protective effects of sAPPα was also confirmed in mouse embryonic fibroblasts (MEFs) derived from APP-KO versus littermate control mice that were subjected to trophic factor/glucose withdrawal using a calcein/ethidium homodimer-3-based assay (Figure 2a) and FACS analysis of PI uptake (Figure 2b).

The neuroprotective function of sAPPα was further substantiated in tissue models when analyzing organotypic (hippocampal) slice cultures prepared from wt or APP-KO mice (Figure 3). There was a pronounced increase of PI-positive cells in serum/glucose-deprived hippocampal slice cultures, which was significantly reduced in the presence of recombinant sAPPα. Importantly, this effect was visible only in wt slices (Figure 3b, upper panel) and not in slices from APP-KO animals (Figure 3b, lower panel). When applying the specific ADAM10 (β-secretase) inhibitor GI254023X (5 μM)40 to serum/glucose-deprived slices, PI counts were significantly increased in comparison with DMSO (carrier)-treated controls. This observation suggests that the absence of endogenous sAPPα exacerbates cell death under these conditions. This cell death-enhancing effect of GI254023X could be completely rescued by applying exogenous sAPPα, thereby supporting the notion that recombinant sAPPα can substitute endogenous sAPPα function (Figure 3c). Microscopic quantification of PI-stained cells in the cornu ammonis (region) 1 (CA1) region of the hippocampus confirmed sAPPα-dependent neuroprotection in wt versus APP-KO cultures and increased cell death in GI254023X-treated slices that was compensated by recombinant sAPPα (Figure 3d).

Recombinant sAPPα and APP-E1 domain activate the PI3K/Akt survival pathway, which requires the expression of holo-APP but not APLP1 or APLP2. Next, we performed in vitro Akt kinase assays with serum-deprived SH-SY5Y neuroblastoma cells to investigate directly the activation of the PI3K/Akt survival pathway by sAPPα.37,38 As seen in Figure 4, trophic factor withdrawal lead to a
pronounced decrease of Akt activity and pGSK3/β (glycogen synthase kinase 3/β) levels, which was prevented by increasing doses of yeast-derived sAPPα and the APP-E1 domain alone. Again, this was only observed in APP-expressing wt cells (Figure 4a, left panel), while SH-SY5Y APP-KD cells did not show any sAPPα-mediated Akt activation (Figure 4a, right panel). To rule out possible protective activities caused by components of the yeast
medium still present in purified sAPP2 and E1, we also tested heat-inactivated fractions, which did not show any rescuing effects (Supplementary Figure 1A). Quantification of the blots (n = 3) confirmed significant induction of Akt activity (Figure 4b, left panel) and enhanced pGSK3β levels (Figure 4b, right panel) for sAPP2 and recombinant E1 in wt cells. In APP-KD cells, only treatment with IGF1 induced phosphorylation of GSK3β to a significant degree. Retrotransfection of APP-KD cells with a holo-APP wt construct restored the sAPP2-dependent Akt activation (Figure 4c).

To further substantiate our findings, we subsequently used hippocampal neurons derived from APP-KO and wt mice. As seen in APP-depleted neuroblastoma cells before, APP-KO neurons failed to show sAPP2-dependent Akt activation that could, however, be readily detected in wt neurons (Figure 4d). Again, these results could be verified by quantification of western blot data (Figure 4e).

To test the possible redundancy of endogenous APP with the APP family members APLP1 and APLP2, we also performed experiments with stable SH-SY5Y APLP1 and APLP2 KD cells (Figure 5a). Interestingly, both Akt pathway activity (Figures 5b and c) and cell survival, as measured by PI staining (Figures 5d and e), were mediated by sAPP2 and the E1 domain independent of APLP1 or APLP2 expression.

The APP C-terminal domain but not the YENPTY motif is required for sAPP2-mediated Akt signaling. Next, we investigated the possible contribution of the APP C-terminal domain (Figures 5f and g). The APP C-terminal domain but not the YENPTY motif is required for sAPP2-mediated Akt signaling.

**Figure 2** sAPP2-mediated suppression of cell death depends on the presence of holo-APP. (a) MEFs prepared from wt or APP-KO mice were cultured in glucose- and serum-free medium for 24 h to induce cell death. In parallel, cells were treated with yeast-derived sAPP2 or IGF1 as a positive control. Cells were stained with green-fluorescent calcein-AM to indicate intracellular esterase activity. This dye exclusively stains live cells (green). Cultures were simultaneously stained with red fluorescent ethidium homodimer-3 to visualize loss of plasma membrane integrity (red, dead cells). Scale bar: 400 μm. (b) MEFs obtained from wt (left panel) or APP-KO mice (right panel) were cultured in glucose- and serum-free medium for 24 h and treated with the indicated amounts of sAPP2 or IGF1. PI-stained cells were analyzed with a FACS cytometer and the extent of cell death was normalized to control cells (+ FCS + Gluc). Data are means from four cultures ± S.E.M. Statistical significance: *P < 0.05 compared with controls (+ FCS + Gluc); *P < 0.05 compared with controls (+ FCS + Gluc); **P < 0.05 compared with controls (+ FCS + Gluc); ***P < 0.05 compared with controls (+ FCS + Gluc).

**Figure 3** sAPP2 induces neuroprotection in organotypic hippocampal slices from wt mice but not from APP-deficient mice. (a) Representative light microscopic image of a hippocampal slice culture. (b) Organotypic hippocampal slices dissected from wt or APP-deficient (APP-KO) mice were cultured on permeable membrane inserts in full medium containing 5% (v/v) horse serum and glucose. After 5 days in vitro (DIV), slices were transferred into serum- and glucose-free neurobasal A medium with 25/50 nM sAPP2 or 20 nM IGF1 for 24 h. Cell death was visualized microscopically by PI staining. Whole slices are shown in the upper panel of each genotype; magnification: × 4; scale bar: 400 μm. A representative area in the CA1 region of the hippocampus (as depicted in the white square) is shown in the lower panel of each genotype; magnification: × 20; scale bar: 100 μm. (c) Organotypic slices from wt mice subjected to serum/glucose deprivation in the presence of the ADAM10 (γ-secretase) inhibitor GI254023X show increased PI staining in comparison with serum/glucose-deprived controls, which is rescued by recombinant sAPP2 (50 nM). (d) Extent of cell death in wt versus APP-KO slices and slices treated with ADAM10 inhibitor was calculated by counting PI-stained (dead) cells in a specific area in the CA1 region and subsequent normalization to serum-glucose-treated controls. Data are means from four to six cultures ± S.E.M. Statistical significance: *P < 0.05 compared with controls (+ HSI + Gluc); **P < 0.05 compared with serum/glucose withdrawal + dimethyl sulfoxide (DMSO) or ADAM10 inhibitor in the absence of sAPP2.
Soluble and membranous APP cooperate to induce Akt

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domain for sAPPβ-induced Akt activation. To this end, we expressed a lentiviral construct encoding the extracellular part from the amino terminus to the β-secretase cleavage site (sAPPβ) of APP coupled to the platelet-derived growth factor receptor (PDGFR) transmembrane domain (TM) in MEF APP-KO cells (Figure 6a). This construct

**a**

![Image of Figure 6a](image1)

**b**

![Image of Figure 6b](image2)

**c**

![Image of Figure 6c](image3)

**d**

![Image of Figure 6d](image4)

**e**

![Image of Figure 6e](image5)
(sAPP/β-PDGFR-TM) lacks the complete APP C-terminal domain. Similar to SH-SY5Y cells and neurons, significant sAPPβ-dependent induction of Akt activity under serum deprivation was also observed in MEF wt cells (Supplementary Figure 1A). However, as presented in Figure 6b (left panel), sAPPα/E1 were not able to activate the Akt pathway in the absence of the APP C-terminal domain as determined by conventional western blots or in vitro kinase assays. To further delineate the domain within the APP C-terminus that mediates Akt activation, we tested primary hippocampal neurons from APP-ΔCT15 mice. These lack the last 15 C-terminal amino acids (aa) of APP, including the YENPTY motif41 that interacts with several adaptor proteins including Fe65 and is crucial for transcriptional activation by the (released) APP intracellular domain (AICD).42 We found that Akt signaling was potently activated in these cells, despite the absence of the YENPTY motif (Figure 6b, right panel) excluding

Figure 5  Akt pathway activation and neuroprotection by sAPPα does not require APLP1 or APLP2. (a) KD of APLP1 and APLP2 in SH-SY5Y neuroblastoma cells. Akt pathway activation by sAPPα in APLP1 (b) and APLP2 KD (c) cells under serum deprivation. (d and e) Cell death was assessed microscopically by counting PI-stained (dead) cells in three random visual fields (>150 cells) and calculated as a percentage of the total number of visualized cells (Hoechst staining). Data are means from four cultures ± S.E.M. Statistical significance: *P<0.05 compared with controls (+ FCS); #P<0.05 compared with serum withdrawal in the absence of sAPPα/E1.
AICD-mediated effects on transcription as the underlying cause of sAPPα-dependent Akt signaling. Quantification of protein levels confirmed significant Akt activity induction in APPΔCT15 mice as compared with MEF cells expressing sAPPβ-PDGFR-TM in an APP-deficient background (Figure 6b, graphs).

**Figure 6b**

- **AMEF sAPPβ-PDGFR-TM**
  - Akt activity
  - pGSK3β
  - GAPDH
  - FCS + - - - - B27 + - - -

- **APP ΔCT15 hippocampal neurons**
  - Akt activity
  - pGSK3β
  - GAPDH
  - FCS + - - - - B27 + - - -

**Figure 6c**

- **Cell surface expression of APP constructs**
  - wt ΔPEER ΔNPTY
  - FCS + - - - - APP KO ΔPEER ΔNPTY

**Figure 6d**

- **AMEF APP ΔPEER and APP ΔNPTY**
  - Akt activity
  - GAPDH
  - FCS + - - - - APP KO ΔPEER ΔNPTY

**Figure 6a**

- **sAPPβ-PDGFR-TM**
  - ectodomain
  - TM
  - AICD
  - G protein interaction motif
  - YENPTY motif

- **APP ΔCT15**
  - E1 E2 A5
  - KGYTSIHGVVEVDATPEERHLKS
  - TM

- **APP ΔNPTY**
  - E1 E2 A5
  - KGYTSIHGVVEVDATPEERHLKS

- **APP ΔPEER**
  - E1 E2 A5
  - KGYTSIHGVVEVDATPEERHLKS
Activation of the PI3K/Akt pathway by sAPPα/E1 is mediated by G-protein-dependent signaling. To further map the regions of the APP C-terminal domain required for sAPPα-induced Akt activation, we transfected two APP deletion mutants (illustrated in Figure 6a) lacking the YENPTY motif (ΔNPTY) or a central sequence motif (ΔPEER) of the APP C-terminal domain into an APP-deficient background. The ΔPEER mutant lacks the membrane-proximal G-protein binding site. Transfection efficiency and overall expression of the APP constructs was verified by western blot (Figure 6a). We also applied an on-cell-Western approach to quantify their cell surface expression after transfection in comparison with wt control cells. As shown in Figure 6c, cell surface expression levels of both constructs were comparable with those detected in wt cells. In line with the result obtained in ΔCT15 neurons, Akt signaling was strongly activated in ΔNPTY-transfected cells (Figure 6d). In contrast, expression of the ΔPEER mutant, despite being expressed at similar levels as APP ΔNPTY, did not induce Akt pathway activation (Figure 6d). This suggests that Akt pathway activation by sAPPα requires G-protein-mediated signaling. Indeed, applying pertussis toxin (PTX), a specific G-protein inhibitor, efficiently blocked sAPPα-mediated Akt signaling in wt MEFs (Figure 7a), which was readily confirmed in blot quantifications (Figure 7a, graph). We also performed FACS analysis with MEF wt cells simultaneously treated with sAPPα and PTX or the PI3K inhibitor LY294002 to quantify cell death of cultures under serum/glucose deprivation. As depicted in Figure 7b, treatment both with PTX and the PI3K inhibitor completely abolished sAPPα-mediated cell survival. This observation further substantiates our findings with APP deletion constructs that sAPPα initiates cell survival via G-protein-mediated activation of the PI3K/Akt pathway.

Discussion

Previous studies from our and other labs suggested that sAPPα exerts potent neuroprotective effects, whereas the amyloidogenic pathway of APP processing and AICD have largely been linked to neurodegeneration and apoptosis. Despite the fact that this apparent dichotomy of APP-dependent biologic effects was proposed several years ago, the overwhelming part of existing experimental work has been devoted to decipher the pathophysiologic, neurotoxic roles of APP. Therefore, the molecular mechanisms of sAPPα-mediated neuroprotection have not been clearly established so far.32 Our data from the present study unequivocally demonstrate that recombinant sAPPα is capable of providing a robust neuroprotection. In line with the observation that synthetic inhibitors of PI3K can abrogate sAPPα-dependent neuroprotection,17–20 they also provide first direct evidence that sAPPα indeed represents a key activator of the PI3K/Akt survival signaling pathway.

It is clearly established that holo-APP can homodimerize, but the cellular receptor coupling sAPPα to downstream survival signaling was hitherto unidentified. In another study, APP dimers were proposed to exert a proapoptotic function, which in turn is antagonized by sAPPα in B103 neuroblastoma cells.33 Despite the somewhat different conclusions drawn by the authors and the fact that sAPPα-dependent intracellular signaling events were not addressed in that study, its principal finding, that is, that APP is required for the antiapoptotic effect of sAPPα, is in line with our own observations. Recently, Jimenez et al.48 also demonstrated that sAPPα can limit induction of the GSK3β stress signaling pathway. Based on experimental evidence exclusively obtained with pharmacologic inhibitors (which often exhibit a limited target specificity and thereby make interpretation of results difficult), this study also suggests that sAPPα-dependent neuroprotection may require the IGF-1 receptor and the insulin receptor (IR) to mediate its effects. The authors proposed a model in which sAPPα acts through IGF-1 and/or IRs to induce the PI3K/Akt pathway and to phosphorylate and inhibit the activity of GSK3β.48 Despite the proposed contribution of coreceptors such as IR and insulin-like growth factor 1 receptor (IGF1-R), our own experiments in IGF1-R-deficient cells did not confirm a role of IGF1-R in sAPPα-mediated neuroprotection (unpublished data). However, our data obtained in this study clearly demonstrate that in the absence of endogenous holo-APP, Akt activation by sAPPα is completely abolished. Our data also suggest that the E1 domain alone can substitute for sAPPα in rescuing the activity of the Akt pathway under conditions of serum deprivation. This could not be observed in APP-depleted cells, emphasizing that the E1 domain, containing the growth factor-like domain, encompasses the neuroprotective active part of sAPPα. Surprisingly, we and others have previously demonstrated that sAPPβ, which is generated via β-secretase cleavage of APP, lacks the neuroprotective properties of sAPPα.37,49 Given the fact that sAPPβ also carries the E1 domain, this lack of neuroprotective properties of sAPPβ versus sAPPα is currently difficult to explain. Therefore, the molecular mechanisms of sAPPα-mediated neuroprotection have not been clearly established so far.32
understand, but may involve conformational differences between these two molecules. Future studies will have to address this important topic of APP biology in more detail.

In line with our findings on the GSK3α/β-regulating function of sAPPα, GSK3/β activity has been found to be upregulated in the AD brain.48,50 These investigations support the hypothesis that the aging-associated decline of sAPPα and its neuroprotective function5,7,32 may contribute to this upregulation, thereby sensitizing neurons to apoptosis. Based on this notion, we hypothesize that the shift toward amyloidogenic APP processing (promoting enhanced generation of Aβ) and loss of the sAPPα-dependent function in neuroprotection may synergize in rendering neurons more prone to neurodegeneration in the aging brain.

In light of the high structural similarity between APP and APLPs,2 we hypothesized that APLP1 and APLP2 may possess similar or overlapping physiologic functions as APP in neuroprotection. It is well established that APP and APLPs can form homo- and heterodimers, arguing for a functional connection between these molecules.10 APLP1 and APLP2 do not contain an Aβ domain, but their ectodomains are shed in an ADAM10-dependent manner similar to APP.51,52 In line with this hypothesis, neuroprotective IGF-1 signaling induces antiamyloidogenic processing of APP and ectodomain shedding of APLP1 and APLP2 in human SH-SY5Y neuroblastoma cells.53 To analyze the potential role of APLP1 and APLP2 in sAPPα-mediated neuroprotection, we also established stable lentiviral KDs in SH-SY5Y cells. Our data clearly demonstrate that APP, but not APLP1 and APLP2, specifically functions as a surface receptor for sAPPα-mediated neuroprotection.

Our observation that endogenous APP was required for sAPPα-mediated neuroprotection suggested a signaling role of the APP C-terminal domain and its interactors in this context. In an alternative scenario, we hypothesized that membrane tethering of APP alone may suffice for sAPPα-mediated activation of the Akt pathway. To further characterize the functional domains of holo-APP required for sAPPα-mediated signaling, we used a construct composed of the E1 and E2 domains of APP fused to the heterologous PDGFR TM. We then overexpressed this construct in an APP-KO background. However, it was not able to restore
sAPPα2-mediated Akt activation under serum deprivation, indicating that the APP C-terminal domain is indeed required for neuroprotection.

The APP C-terminal domain couples APP to diverse intracellular signaling pathways. It was proposed that the Src homology 2 domain of Abl or the phosphotyrosine-binding domain (PTBD) of Shc may interact with the YENPTY motif of APP. This would suggest a role for APP in tyrosine kinase-mediated signal transduction.54 Consequently, to further investigate the putative role of YENPTY interactors, we used primary neuron cultures of APP ΔCT mice (lacking the 15 C-terminal aa of APP) to investigate the possible contribution of PTBD-containing interactors to sAPPα2-mediated Akt activation. Intriguingly, our data show that the last C-terminal 15 aa of the APP C-terminal domain are dispensable for mediating the neuroprotective effect of sAPPα2. This domain comprises the YENPTY motif to which the vast majority of the APP C-terminal domain interactors bind.55

Our finding that the APP C-terminal domain, but not the YENPTY motif, is essential for mediating sAPPα2/APP-induced neuroprotection suggested that YENPTY-independent interactors are required for sAPPα2-induced neuroprotection. Indeed, the heterotrimeric G-protein subunit Gα(S), which binds in a YENPTY motif-independent manner to the APP C-terminal domain, was recently shown to be involved in APP-mediated modulation of GSK3β and neurite outgrowth.45 To further map the APP C-terminal domain region mediating sAPPα2-dependent neuroprotection, we reconstituted APP-KO cells with a deletion mutant (ΔAPEER) lacking the G-protein interaction motif.45,55 In contrast to the ΔNPTY mutant, expression of the ΔAPEER mutant43,44 did not rescue sAPPα2-induced Akt activation after serum starvation, indicating that the G-protein-mediated signaling is causally involved. Consistently with these data, the G-protein inhibitor PTX completely abolished sAPPα2-induced Akt activation and cell survival. Of note, G-protein-dependent activation of the PI3K/Akt pathway has been previously demonstrated in different cell models.56,57

In conclusion, we were able to identify key signaling components and mechanisms involved in sAPPα2-mediated neuroprotection: our data indicate that sAPPα2 signals either by direct binding or via an indirect mechanism through membrane-tethered APP. As APP harbors a G-protein interaction domain and sAPPα2-mediated neuroprotection is lost upon deletion of this motif, this strongly suggests that APP serves as a receptor to trigger G-protein-dependent activation of PI3K. This activation leads to the recruitment and downstream activation of the prosurvival kinase Akt and subsequent inhibition of its proapoptotic target GSK3β (Figure 8). These findings support the hypothesis that sAPPα2 and holo-APP share equal relevance in mediating the neuroprotective function of APP. They also provide novel mechanistic insights into the physiologic function of APP in limiting neuron damage and death in response to neurotoxic stress conditions, as well as the loss of this function during brain aging.

Materials and Methods
Materials. Unless stated otherwise, cell culture media and supplements were purchased from Invitrogen (Darmstadt, Germany), human IGF1 from Sigma-Aldrich (Seelze, Germany) and Millicell cell culture inserts (0.4 µm, 30 mm diameter) for hippocampal slice cultures from Merck Millipore (Darmstadt, Germany). PTX and LY294007 were purchased from Enzo Life Sciences (Lörrach, Germany). All other biochemicals and chemicals were provided in analytical grade purity from Roth (Karlsruhe, Germany) or Sigma-Aldrich. Plasticware was purchased from Corning Life Sciences (Wiesbaden, Germany) or Greiner Bio-One (Frickenhausen, Germany).

Cell and tissue culture. Immortalized MEFs were derived from wt or APP-KO mice.55 Hippocampal neurons and organotypic slices were dissected from the same mice. SH-SY5Y cells were cultured in Dulbecco’s modified Eagle’s medium (DMEM) with F-12 Nutrient Mixture (Ham) supplemented with 10% fetal calf serum (FCS), 2 mM l-glutamine, 100 U/ml penicillin and 100 µg/ml streptomycin (pen/strep). MEFs were grown in DMEM supplemented as above. Organotypic (hippocampal) slice cultures were prepared from P4 to 6 transgenic and non-transgenic littermate mouse pups as described in Stoppini et al.59 and cultured on cell culture inserts (interface method). Broken or otherwise lesioned slices were discarded. Slices were maintained in culture medium containing 45% minimum essential medium, 25% basal medium Eagle, 25% horse serum (HS), 3% glucose (20%), 1% l-glutamine, 0.5% pen/strep and 0.5% fungizone, adjusted to pH 7.2, and kept at 5% CO2 and 37°C. Culture medium was changed every 2–3 days. For cell death experiments, the culture medium was replaced with glucose-free neurobasal A medium without HS (full medium as control) after 5–6 days in vitro. Until then, slices had shrunk from 400 µm thickness to ~200 µm. PI stain (0.8 µg/ml) was directly added to the medium before microscopic evaluation. Primary hippocampal neurons were dissociated from hippocampal slices obtained from P1 to 3 pups and plated into full medium (neurobasal A medium supplemented with 2% B27, 2% Glutamax, 0.2% pen/strep, 1% gentamycin) on poly-D-lysine-coated plates. Neurons were maintained at 5% CO2 and 37°C for 7 days before experiments.

Animals. All mice were bred and maintained according to the FELASA and National Animal Experimental Ethics Committee Guidelines. We used wt, APP-KO (APP/−/−) and APP-ΔCT15 C57Bl/6 mice.41,58 APP-KO mice lack membrane-anchored full-length APP and all of its proteolytic fragments; APP-ΔCT15 mice lack the last 15 aa of the C-terminus of APP, which includes the YENPTY motif. APP-KO and APP-ΔCT15 mice had been backcrossed for at least six generations to C57Bl/6 mice.

Viral vectors and plasmids. Stable APP/APLP1/APLP2 KOs were achieved by transducing wt SH-SY5Y cells with lentiviral particles to stably express shRNAs directed against huAPP or huAPLPs. Mission shRNA Plasmid...
DNA containing the pLKO.1-puro vector was purchased from Sigma-Aldrich (clone IDs for APP: NM_000484.2-3202s1c1; APLP1: NM_005166.2-1711s1c1 and APLP2: NM_001642.1-1582s1c1). Packaging (psPAX2) and enveloping (pMD2.G) plasmids were from Addgene (Cambridge, MA, USA). Production of lentiviral particles was performed following Addgene’s pLKO.1 TRC Cloning Vector Protocol (2006). Lentiviral vectors encoding the fusion protein sAPP /pDGFR-TMD were generated by triplicate transfection of HEK-293T cells with the envelope plasmid pMD2.G, the packaging plasmid pCMVΔR8.9 and the transfer vector plasmid pHAIN as described.43 The plasmid pHAIN encodes human-gal4::VP16 under control of the CMV promoter fused at its N-terminus to the HA tag and the IgG signal peptide. At the C-terminus, sAPP was then fused to a myc tag followed by the TM of the PDGFR as provided by the pDisplay plasmid (Invitrogen). Thus, upon transfection with this vector, MEFs expressed the extracellular part of APP but not its TM. The APP mutant constructs lacking the YENPTY motif (Δaaa 680–691 of APP695) or the G-protein interaction motif (Δaaa 668–679) have been described before and were subconed in-frame from pUAS into pCDNA3.1 behind the N-terminal myc tag.43 Plasmids were purified with the Plasmid Max Kit from Qiagen (Hilden, Germany) and commercially sequenced to confirm the mutations. Transfection of MEF APP-KO cells with APP constructs was performed with FuGENE HD Transfection Reagent (Roche Diagnostics, Mannheim, Germany) according to the manufacturer’s instructions and verified in a western blot.

### Purification of recombinant sAPPβ/E1 domain.

The coding sequences of sAPPβ695 (aa 18–611) and APP-E1 (aa 18–188) were cloned in frame as 6 x His-tagged fusions in a pBLHIS-SX-derived expression vector and expressed in Pichia pastoris GS115 cells. The asparagine residue at position 467 of APP695 (glycosylation site) was replaced with serine by site-directed mutagenesis. The yeast was cultured in buffered glycerol complex medium with peptone. Addition of methanol induced the expression of 6-His-tagged APP (C1 from pUAS into pCDNA3.1 behind the N-terminal myc tag. 43 Plasmids were expressed in Pichia pastoris according to the manufacturer’s instructions and verified in a western blot.

### Cell death and viability assays.

Cell death was quantified based on plasma membrane permeabilization. To this aim, cells or tissue cultures were cultured under normal serum conditions (control) and serum/trophic factor deprivation (– FCS; – B27; – HS) or double deprivation with additional removal of glucose (– FCS – Gluc) for 24–48 h. Cultures were treated in parallel with either 25–50 nM sAPPβ; 50 nM APP-E1 domain or 20 nM IGF1, as indicated. PTX (Enzo Life Sciences, 100 ng/ml) or P38K inhibitor LY294002 (Enzo Life Sciences, 10 μM) were applied 30 min before sAPPβ was added to the medium. Following collection, cells were stained with 0.8 μg/ml PI, a DNA-intercalating and fluorescent agent, to visualize dead cells that emit red light at 617 nm when excited at 535 nm. Cells were counted manually under a fluorescence microscope in three random visual fields (> 150 cells) and counter-stained with Hoechst to calculate the percentage of dead cells versus the total number of visualized cells. Alternatively, cells were stained with PI and analyzed with a FACS cytometer (BD Biosciences, Heidelberg, Germany). Organotypic hippocampal slices were incubated with 0.8 μg/ml PI for 20 min at 37 °C. When applying the ADAM10 (α-secretase) inhibitor G254023X (5 μM), slices were cultured in serum-free medium for 48 h containing the inhibitor or its respective carrier (DMSO) as control. Round circles of identical size (Ø 500 μm) were positioned in equivalent locations within the CA1 region of each hippocampus image and all PI-stained cells were counted using ImageJ software (NIH, Bethesda, MD, USA). Cell viability assays were performed with a commercial kit (CellTiter-Glo Luminescence Assay, Promega, Mannheim, Germany) according to the manufacturer’s instructions. The assay quantitates the cell number, an indicator of cellular metabolic activity, photometrically with a fluorescence plate reader. Additionally, we applied the live-dead cell staining kit II from Promokine (Heidelberg, Germany) according to the manual. Cells were simultaneously stained with green fluorescent calcium-AM (4 mM; ex/em: 495/515 nm) to detect intracellular esterase activity (viable cells) and red fluorescent ethidium homodimer-3 (2 mM; ex/em: 530/635 nm) to indicate loss of plasma membrane integrity (dead cells).

### Akt kinase activity assays.

Cell cultures were pre-treated with 10–50 nM yeast-derived sAPPβ/E1 or 20 nM human IGF1 for 24 h before removal of glucose and/or serum. During starvation for 24–48 h, the same treatments were administered. IGF1 was added every 24 h owing to its short half-life. In the experiment using PTX, 100 ng/ml of the toxin was applied 30 min before sAPPβ was added to the medium. Akt kinase activity was measured in vitro with a commercial kit (Akt kinase assay kit; Cell Signaling, Frankfurt/Main, Germany) according to the manufacturer’s protocol. Briefly, endogenous levels of pAkt were immunoprecipitated from whole-cell extracts with immobilized pAkt (Ser473) mAb (bead conjugate) overnight. After extensive washing, the kinase assay was performed using 10 mM ATP and GSK3 β fusion protein (27 kDa) as a substrate. Subsequent inactivation of GSK3 β was measured by western blot detecting pGSK3 β/ Ser (Ser21/9, 27 kDa).

### Immunoblotting.

For western blotting, cells were washed with PBS, harvested and lysed with SDS lysis buffer (2% SDS, 68.5 mM Tris-HCl, 10% glycerin, 1 mM protease/phosphatase inhibitor cocktail) or lysis buffer from the Akt kinase assay kit supplemented with 1 mM PMSF followed by sonication. The protein amount was quantified using the Pierce BCA Protein Assay Kit (Thermo Fisher, Schwerte, Germany). Equal amounts were used for the Akt kinase assays or directly loaded onto 10–12% bi-acrylamide/SDS gels for conventional western blots and electrophoretically transferred to nitrocellulose membranes (Whatman Protran BA 83, 0.2 μm; GE Healthcare, Little Chalfont, UK). Unspecific binding was blocked for 1 h in 5% non-fat powdered milk in 0.05% Tween-20 (v/v in TBS) followed by overnight incubation at 4 °C with primary antibodies specific for GSK3 β/ (rabbit; Ser21/9; Cell Signaling), pGSK3 β/ (rabbit, D44A; Cell Signaling), GSK3 β/ (mouse, 3D10; Cell Signaling), APP (mouse, 22C11; Millipore, Darmstadt, Germany), Bim (rabbit; Cell Signaling) or APLP1/APLP2 (rabbit; Millipore). Equal loading was monitored by probing membranes with glyceraldehyde 3-phosphate dehydrogenase (GAPDH) (Millipore). The corresponding secondary antibodies coupled with infrared dyes in red (680 RD) or green (800 CW) against rabbit or mouse (IRDye goat anti-rabbit or anti-mouse from LI-COR Biosciences, Bad Homburg, Germany) were diluted in 5% bovine serum albumin, followed by detection with the LI-COR Odyssey Infrared Imager (LI-COR Biosciences). On-cell-Western assays were performed to evaluate cell surface expression of APP. The experiment was performed by adding primary antibody (22C11, 1:160) directly into the medium of live, unpermeabilized cells for 4 h. After three washing steps with PBS, secondary antibody (IRDye goat anti-mouse from LI-COR) was applied for 1:1000 incubated for 1 h. Following washing steps, fluorescence signals were measured using the LI-COR Odyssey Infrared Imaging and Image Studio 3.1 software (LI-COR Odyssey, Bad Homburg, Germany). Values are arbitrary fluorescence units calculated from trim signals (means of n = 3 experiments ± S.E.M.) normalized to background staining controls.

### Statistical analyses and microscopy.

Results are expressed as mean ± S.E.M. All experiments were repeated at least three times yielding similar results. Statistical analyses were performed using SPSS (IBM, Armonk, NY, USA) with one-way ANOVA followed by Tukey’s HSD post hoc comparisons. P-values < 0.05 were considered as statistically significant. Western blot quantifications were performed with LI-COR Odyssey Image Studio 3.1 software to guarantee analysis of fluorescence signals in a linear range or ImageJ software for ECL-developed blots. Values were normalized to serum/glucose-treated controls (dashed line in figures). Band intensities in blots with whole-cell lysates were normalized to their corresponding loading control (GAPDH) and plotted as pGSK3 β/GAPDH ratios. All graphs show means of n = 3 bi-quantifications ± S.E.M. For microscopy, we used the Nikon Eclipse TE2000-S fluorescence microscope with Plan Fluor × 4, × 10 or × 20 dry objectives, a 100 W mercury lamp and FITC (green; ex: 465–495 nm; dichroc mirror: 505 nm; em: 515–555 nm) or Texas Red (ex: 540–580 nm; dichroc mirror: 595 nm; em: 600–660 nm) excitation filters. Images were acquired with a D5-SMC cooled color digital camera (Nikon, Düsseldorf, Germany) and NIS Elements AR (version 3.22) software from Nikon. Image adjustments such as changes of contrast and brightness were applied equally across the entire image.

### Conflict of Interest

The authors declare no conflict of interest.

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