NGF controls APP cleavage by downregulating APP phosphorylation at Thr668: relevance for Alzheimer’s disease

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Summary

NGF has been implicated in forebrain neuroprotection from amyloidogenesis and Alzheimer’s disease (AD). However, the underlying molecular mechanisms are still poorly understood. Here, we investigated the role of NGF signalling in the metabolism of amyloid precursor protein (APP) in forebrain neurons using primary cultures of septal neurons and acute septo-hippocampal brain slices. In this study, we show that NGF controls the basal level of APP phosphorylation at Thr668 (T668) by downregulating the activity of the Ser/Thr kinase JNK(p54) through the Tyr kinase signalling adaptor SH2-containing sequence C (ShcC). We also found that the specific NGF receptor, Tyrosine kinase A (TrkA), which is known to bind to APP, fails to interact with the fraction of APP molecules phosphorylated at T668 (APP(T668)) directly. Accordingly, the amount of TrkA bound to APP is significantly reduced in the hippocampus of ShcC KO mice and of patients with AD in which elevated APP(T668) levels are detected. NGF promotes TrkA binding to APP and APP trafficking to the Golgi, where APP–BACE interaction is hindered, finally resulting in reduced generation of sAPPβ, CTFβ and amyloid-beta (1-42). These results demonstrate that NGF signalling directly controls basal APP phosphorylation, subcellular localization and BACE cleavage, and pave the way for novel approaches specifically targeting ShcC signalling and/or the APP–TrkA interaction in AD therapy.

Key words: AD; APP(T668); BACE; NGF; ShcC; TrkA–APP interaction.

Introduction

Accumulating evidence suggests that the NGF signalling system may be disrupted early in sporadic AD. In particular, alterations of the NGF signalling system in the basal forebrain correlate more robustly than the amyloid load with cognitive deficits in mild cognitive impairment (MCI) and with MCI progression towards AD (Mufson et al., 2012).

Consistent with these findings, antibody-mediated neutralization of NGF causes increased amyloid in AD111 mice (Ruberti et al., 2000). Moreover, in vitro models of NGF deprivation show that amyloid accumulation is induced and AD pathology is mimicked in PC12 and hippocampal neurons (Matrone et al., 2008a,b); this has been referred to as ‘Alzheimer’s-like molecular syndrome’ (Calissano et al., 2010).

Furthermore, NGF treatment can ameliorate cognition and amyloidogenesis in AD transgenic models (Yang et al., 2014), and AD clinical trials based on NGF gene therapy have produced promising results with respect to cognitive preservation (Rafi et al., 2014).

Despite these findings, the molecular mechanisms underlying the neuroprotective action of NGF in AD-like neurodegeneration remain unclear. In particular, elucidating how NGF modulates APP processing in neurons is critical for our basic understanding of the phenomenon and for the early diagnosis and treatment of AD.

Amyloid precursor protein (APP) can be processed via alternative physiological or amyloidogenic pathways depending on whether cleavage occurs via alpha secretase 10/17 (ADAM 10/17) or beta secretase 1 (BACE1) in neurons, which causes the generation of alpha (sAPPα, CTFα) or beta (sAPPβ, CTFβ) products, respectively (Bernstein et al., 2009; Vassar et al., 2014).

APP phosphorylation pattern has been reported to be crucial for APP binding to cytosolic interactors and for its subcellular trafficking and final processing (Suzuki & Nakaya, 2008; Tamayev et al., 2009). Particularly, phosphorylation of APP at T668 (APP695) increases the beta product levels by facilitating exposure to and cleavage by BACE in human patients with AD and transgenic models (Lee et al., 2003; Shin et al., 2007). The APP695 form is neuron-specific, highly expressed in AD dystrophic neurites and amyloid plaques, and induces neurodegeneration (Lee et al., 2003; Chang et al., 2006). Of interest, mice bearing a Thr to Ala substitution at 668 (T668A) prevent AD cognitive deficits when crossed to the FDD1 mouse model of dementia (Lombino et al., 2013). Accordingly, the inhibition/reduction of T668 phosphorylation may be a potential target for AD therapy (Chang et al., 2006).

Despite many studies on the anti-amyloidogenic action of NGF in the brain, to date the effect of NGF on APP695 phosphorylation has not yet been investigated. Thus, we investigated the NGF-mediated control of APP phosphorylation/processing in the forebrain and the underlying molecular mechanisms using in vitro (primary septal neurons), ex vivo (septo-hippocampal [SH] slices), and in vivo (KO mice and AD brain tissues) experimental paradigms.

Our results show that NGF rescues neurons from JNK(p54)-mediated phosphorylation of APP through its early-downstream signalling adaptor ShcC. The reduction in APP phosphorylation induced by NGF favours APP association with its receptor TrkA, while affecting APP interaction with BACE, and reduces the generation of beta products, such as CTFβ and amyloid (1-42). These findings suggest that NGF control of APP phosphorylation and the subsequent events are potentially disrupted in the human AD brain where the APP695 level is elevated and the APP–TrkA interaction is dramatically reduced.
Overall, our data support a role of APP (T668) phosphorylation in the NGF-driven control of amyloidogenesis in the basal forebrain and suggest early-downstream effectors of the NGF pathway as targets in the development of novel anti-amyloidogenic therapies for AD.

**Results**

**NGF controls basal APP phosphorylation at T668**

The phosphorylation of APP at T668 is known to facilitate APP amyloidogenic processing and AD pathology (Lee et al., 2003; Chang et al., 2006).

To assess the effect of NGF on APP phosphorylation at T668 in the forebrain, SH slices from control mice were incubated with NGF (100 ng mL\(^{-1}\); 1 h) and analysed by Western blot (WB) using specific antibodies against APP\(^{pT668}\) and total APP (Fig. 1A). The APP\(^{pT668}\) level was significantly reduced in the NGF-treated SH slices compared with the untreated slices (CTR, 44.4 ± 17.5%; *P < 0.05, Fig. 1B). Next, we restricted our analysis to the septal neurons (DIV10, E18), which are TrkA-expressing neurons and the primary cellular target of NGF in the SH system. We incubated mouse septal neurons with NGF (100 ng mL\(^{-1}\); 1 h) and confirmed that the APP\(^{pT668}\) level (Fig. 1C) was significantly decreased after NGF treatment (72.3 ± 6.1% of CTR, *P < 0.05; Fig. 1D), indicating that the effect of NGF on APP\(^{pT668}\) is specifically directed towards these neurons. The NGF-driven decrease in APP\(^{pT668}\) does not result from APP protein degradation because the total amount of APP protein was unaffected by NGF in both paradigms (Fig. 1A,C). The same experiment was performed using rat primary neurons with comparable results (Fig. 1A,B).

We next assessed whether in turn NGF withdrawal promotes APP phosphorylation by incubating SH slices with anti-NGF neutralizing antibody. We analysed APP\(^{pT668}\) and total APP by WB (Fig. 1E) and found almost doubled levels of APP phosphorylation following 6-h incubation with anti-NGF antibody (195.9 ± 5.3% CTR, **P < 0.01; Fig. 1F).

We also used the classic NGF deprivation paradigm in the pheochromocytoma cell line (PC12). PC12 cells were differentiated into neurons by culturing in NGF (100 ng mL\(^{-1}\)) under serum-free conditions for 10 days. Then, the PC12-derived neurons were switched to serum-free and NGF-free media for the indicated time periods (30’, 3 h) and analysed by culturing in NGF (100 ng mL\(^{-1}\)/C6 or 3 h of NGF deprivation.

Our data indicate that NGF controls basal APP phosphorylation at T668 in septal neurons in vitro and ex vivo.

**NGF treatment decreases activation of JNK (p54) kinase**

Given that JNK (Standen et al., 2001; Colombo et al., 2009; Sclip et al., 2011) and cyclin-dependent kinase 5 (CDK5) (Iijima et al., 2000; Liu et al., 2003) are Ser/Thr kinases and regulate APP phosphorylation under basal and pathological conditions, they were analysed as potential mediators of the NGF effect on APP\(^{pT668}\). We incubated SH slices with the CDK5 and JNK inhibitors roscovitine (R; 14 μM, 1 h) and SP600125 (SP; 2 μM, 1 h), respectively and found a dramatic reduction in the APP\(^{pT668}\) level (Fig. 2A). The inhibitor concentrations and incubation times were selected to produce the maximum inhibitory effect on the specific kinase without affecting cell survival (data not shown). We analysed APP phosphorylation levels also in mouse models of genetic ablation of the three JNK isoforms (JNK1-3) and found reduced APP\(^{pT668}\) levels in the hippocampal extracts of JNK1, JNK2 and JNK3 KO mice (n = 2 mice per genotype) (Fig. 2B).

We observed that NGF does not affect the level and/or activation of CDK5. Briefly, SH slices were incubated with NGF (100 ng mL\(^{-1}\), 1 h) or roscovitine (R; 14 μM, 1 h), and WB was performed using antibodies against CDK5 or p35, the CDK5 activator (Fig. 2C). NGF was not able to...
modulate CDK5 or p35 levels, while the p35 level was affected by roscovitine, as expected (Fig. 2C). SH slices were then incubated with NGF (100 ng mL\(^{-1}\), 1 h) or SP600125 (SP; 2 \(\mu\)M, 1 h), and JNK activation was analysed by WB. A decrease in the phosphorylation of the two JNK bands of 46 (p46) and 54 (p54) kDa was observed upon SP treatment, while NGF specifically affected the activation of only the p54 isoform (Fig. 2D).

Therefore, we repeated the analysis to quantify the phosphorylated p54 level in the SH slices treated with NGF (100 ng mL\(^{-1}\), 1 h) and observed a significant reduction in the phosphorylation level of the JNK (p54) isoform compared with the untreated slices (59.4 ± 4.4% of CTR, \(P < 0.05\); Fig. 2E,F). A similar correlation was also observed in mouse septal primary neurons, in which we observed a significant decrease in the levels of phosphorylated p54 in SH slices (F) or mouse septal neurons (G) (Fig. 2F). The levels of phosphorylated p54 in SH slices (F) or mouse septal neurons (G) were measured, and the ratio was calculated over the total p54 level, and expressed as percentage of the control (59.4 ± 4.4% of CTR, \(P < 0.05\) and 61.5 ± 2.1% of CTR, \(P < 0.01\), respectively). Beta-actin was used as a loading control. Triplicate data from 4 to 5 independent experiments were used for the analysis. SH = septo-hippocampal, HP = hippocampus.

**Deletion of the ShcC gene causes increased pJNK activation and APP\(^{T668}\)**

Isoform C of the SH2-containing sequence adaptor (ShcC) couples NGF/TrkA activation with neuronal inhibition of JNK (Molton et al., 2003). We thus hypothesized that ShcC signalling could act early downstream of NGF to downregulate pJNK and, consequently, APP\(^{T668}\). To investigate this potential mechanism, we examined pJNK (Fig. 3A) expression in hippocampal tissue from ShcC KO mice by WB and found an almost 50% increase in the pJNK level compared with the age-matched WT mice (144 ± 5% of WT, \(P < 0.05\); Fig. 3C). Consistent with our hypothesis, the basal APP\(^{T668}\) level determined by WB (Fig. 3B) was found to be almost 50% increased (141 ± 8% of WT, \(P < 0.05\), Fig. 3C). Moreover, elevated amounts of the APP C-terminal fragment (CTF), and in particular CTFβ (as indicated by the anti-APP\(_{6E10}\) antibody; Fig. 3D), in the presence of normal BACE levels (Fig. 3E) were observed in the ShcC KO mice. The ablation of the gene product was verified probing ShcC KO and WT tissues with specific antibodies against ShcC (Fig. 3F). The beta-amyloid (1-42) levels, measured by a highly sensitive rodent-specific EUSA, increased in the hippocampus of ShcC KO (16 ± 0.6 pg mg\(^{-1}\) proteins), as compared to age-matched control mice (10 ± 1.4 pg mg\(^{-1}\) proteins, \(P < 0.05\); Fig. 3G).

Furthermore, NGF stimulation (100 ng mL\(^{-1}\), 1 h, RT) of SH slices from ShcC KO mice (ShcCKO+NGF) failed to reduce JNK activation (102 ± 11% of ShcC KO, \(P = 0.096\)) and APP phosphorylation (104.5 ± 5.7% of ShcC KO, \(P = 0.37\)), as shown in Fig. 3H-I. Also, levels of CTFβ are comparable between NGF-treated and untreated SH slices from ShcC KO (Fig. 3H).

Our results indicate that ShcC deletion prevents NGF from its basal regulation of pJNK and APP\(^{T668}\) and suggest that releasing the NGF-TrkA-ShcC control of APP metabolism leads to increased CTFβ and amyloid (1-42) generation in vivo.

**APP does not interact nor colocalize with TrkA, upon phosphorylation at T668**

TrkA receptor has been shown to interact with APP in vitro and in vivo (Tarr et al., 2002; Matrone et al., 2011). Because APP undergoes a conformational change (Ramlot & Nicholson, 2001) upon T668 phosphorylation that detaches its cytoplasmic interactors (Ando et al., 2001), we investigated whether T668 phosphorylation affects TrkA binding to APP. We analysed the intracellular localization of Trk, total APP and APP\(^{T668}\) using confocal microscopy. Septal neurons were treated with NGF (NGF; 100 ng mL\(^{-1}\), 1 h) and assayed by double immunofluorescence staining for Trk and APP (Fig. 4A and at low magnification in Fig. 5A), or Trk and APP\(^{T668}\) (Fig. 4B and at low magnification in Fig. 5B).
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**Fig. 3** Deletion of the ShcC gene results in increased levels of pJNK, APP\(^{pT668}\), sAPP\(^b\), CTF\(^b\) and amyloid-beta (1-42). (A, B) Representative WB of hippocampal extracts from ShcC \(KO\) (\(n = 3\)) and WT (\(n = 3\)) mice using antibodies against pJNK and total JNK (A), or APP\(^{pT668}\) and C-terminal APP (B). (C) The graph illustrates the optical density analysis of pJNK (144 \(\pm\) 5\% of WT, \(\ast P < 0.05\)) and APP\(^{pT668}\) (141 \(\pm\) 8\% of WT, \(\ast P < 0.05\)) measured over total JNK and APP in ShcC \(KO\) mice, and the values are reported as a percentage of the values in WT mice. (D–F) A representative WB of hippocampal extracts from ShcC \(KO\) and age-matched WT mice probed with specific antibodies against APP (APP C-ter and 6E10; D), BACE (E) and ShcC (F). (G) Endogenous amyloid-beta (1-42) levels in hippocampal extracts of ShcC \(KO\) (16 \(\pm\) 0.6 pg mg\(^{-1}\) proteins) and age-matched wild-type mice (10 \(\pm\) 1.4 pg mg\(^{-1}\) proteins), \(\ast P < 0.05\). \(N = 3\) mice per genotype were used for this analysis. (H, I) SH slices from ShcC \(KO\) mice were incubated with NGF (100 ng ml\(^{-1}\), 1 h), extracted and probed for phosphorylated and total APP and JNK, and the CTF\(^b\). The data are reported as percentage of control in the graph in (I). Levels of pJNK (102 \(\pm\) 1.1\% of ShcC \(KO\); \(P = 0.096\)) and APP\(^{pT668}\) (104.5 \(\pm\) 5.7\% of ShcC \(KO\); \(P = 0.37\)) are unchanged following NGF incubation. Beta-actin was used as a loading control. Data from two separate experiments with brain slices from four animals per genotype were used for the analysis. HP = hippocampus; WT = wild-type; SH = septo-hippocampal.

**Fig. 4** (A) Synaptic terminals innervating the CA1 region of the hippocampus (HP) were subjected to IP with a specific anti-TrkA antibody, and then probed with anti-APP\(^{pT668}\) or total APP (22c11) and trk antibodies. We confirmed that APP co-immunoprecipitates with TrkA, as expected (Fig. 4C). In contrast, under the same experimental conditions, APP\(^{pT668}\) failed to co-immunoprecipitate with TrkA (Fig. 4C). The specific interaction of TrkA with APP and not APP\(^{pT668}\) was further confirmed by IP studies with SH slices. The lack of an APP\(^{pT668}\)–TrkA association was also observed in both the untreated and NGF-treated SH slices (Fig. 4D) even though the amount of total APP that immunoprecipitated with TrkA increased upon NGF treatment (Fig. 4D).

Our data indicate that NGF stimulation promotes APP binding to its receptor TrkA but that only the fraction of APP that is not phosphorylated at T688 is able to interact with TrkA. Therefore, it was not surprising to observe that TrkA IP by APP was almost completely lost in the ShcC \(KO\) mice (Fig. 4E), a mouse model with elevated APP\(^{pT668}\) (Fig. 3B) Furthermore, we investigated the Tyr 490-phosphorylated form of TrkA (pTrkA) because of the selective NGF-driven docking of Shc to pTrkA. The levels of APP (Fig. 3B,D; Fig. 4E) and pTrkA (Fig. 4E) in the ShcC \(KO\) mice were found to be unaffected. The weak residual APP–TrkA binding observed in the ShcC \(KO\) mice may result from the presence of other Shc isoforms in the adult brain, presumably ShcB (Liu & Meakin, 2002).

We also studied APP\(^{pT668}\) and Trk localization in the absence of ShcC by double IF of coronal brain sections from perfused ShcC \(KO\) (Fig. 4F) and WT age-matched mice (not shown). In particular, the cholinergic cell soma in the medial septum (MS) and diagonal band of Broca and their terminals innervating the CA1 region of the hippocampus (HP) were analysed in ShcC \(KO\). The analysis confirmed in vitro confocal imaging and showed that Trk and APP\(^{pT668}\) do not colocalize in vivo in ShcC \(KO\) mice, neither in the MS (\(r = 0.03 \pm 0.01\)) nor in the CA1 (\(r = 0.08 \pm 0.01\)).

These data indicate that TrkA will colocalize and interact with APP as long as none of it is phosphorylated at T688. Accordingly, the APP–TrkA interaction is significantly affected in the ShcC \(KO\) brain, where high levels of APP phosphorylation occur.

**The APP–TrkA interaction is significantly affected in the hippocampus of AD patients**

We hypothesized that the APP–TrkA interaction is affected in the AD brain, where an elevated APP\(^{pT668}\) level is often reported (Lee et al., 2003; Chang et al., 2006). To test this hypothesis, we analysed the
APP\textsubscript{pT668} level using WB (Fig. 5A–C) and the APP–TrkA interaction using IP (Fig. 5D, E) in brain samples from control subjects without neurological diseases (CTR; \( n = 3 \)), patients with AD (AD; \( n = 3 \)) and subjects affected by other neurological diseases (OND; \( n = 3 \)). We observed a dramatic increase in the APP\textsubscript{pT668} level in the hippocampus of patients with AD (Fig. 5A), as compared to both CTR (319.5 \pm 46.1\% of CTR),
**P < 0.01; Fig. 5B) and OND (202.4 ± 15.6% of OND, *P < 0.05; Fig. 5C), as expected. Then, we immunoprecipitated the APP from the CTR, OND and AD hippocampal tissue and probed the samples with a specific anti-TrkA antibody. We found that the amount of TrkA bound to APP was significantly lower in all three AD samples, as shown in Fig. 5D. Analysis of the TrkA-positive co-immunoprecipitated band relative to the input showed lower binding in AD (*P < 0.05; 56.4 ± 12, %CTR), as compared to both CTR and OND (108 + 15, %CTR), while no difference was found between CTR and OND (P = 0.6) hippocampal samples.

Next, we repeated the IP analysis using the hippocampal and cerebellar extracts from the same AD subject (AD2) and found that the APP-TrkA interaction was almost completely undetectable in the hippocampus of this AD patient but preserved in the AD cerebellum (Fig. 5E).

Our data reveal that a lack of APP-TrkA binding correlates with an elevated APP\textsubscript{T668} level and it is specific of neurodegenerating tissues in humans with AD.

**NGF promotes the APP binding with TrkA versus BACE1, by inducing APP trafficking to the Golgi, and reduces beta products generation**

Finally, we investigated the effect of NGF on the APP interaction with BACE and APP subcellular localization. Both of these events are known to be modulated by APP phosphorylation at T668 (Lee et al., 2003; Muresan & Muresan, 2005). For this investigation, we treated SH slices with NGF (100 ng mL\textsuperscript{-1}, 1 h), performed an IP analysis with anti-APP antibody, and probed the samples with specific anti-TrkA or anti-BACE1 antibodies. We observed that NGF increases APP binding to TrkA and concomitantly decreases APP binding to BACE1 (Fig. 6A).

To study APP cleavage following NGF treatment, we performed WB for APP, sAPP\textsubscript{\alpha}, sAPP\textsubscript{\beta}, CTF\textsubscript{\beta}, BACE1 and ADAM17 (Fig. 6B) on untreated and NGF-treated (100 ng mL\textsuperscript{-1}, 1 h) SH slices. We found increased sAPP\textsubscript{\alpha} (129.4 ± 9.3% of CTR, *P < 0.05) and reduced sAPP\textsubscript{\beta} (50.6 ± 12.2% of CTR, *P < 0.05) and CTF\textsubscript{\beta} (52.5 ± 7.3% of CTR, *P < 0.05) expression upon NGF treatment (Fig. 6C). The levels of full-length APP (86.2 ± 6.9% of CTR, P = 0.12), BACE1 (92.2 ± 18.7% of CTR, P = 0.69) and ADAM17 (113.6 ± 5.8% of CTR, P = 0.08) were unaffected (Fig. 6C). Moreover, we measured endogenous amyloid-beta (1-42) levels in the hippocampus of NGF-treated SH slices (10 ± 0.1 pg mg\textsuperscript{-1} proteins) and found a significant reduction compared to control (14.5 ± 0.5 pg mg\textsuperscript{-1} proteins, *P < 0.05, Fig. 6D).

To determine whether NGF reduces the interaction between APP and BACE by modulating subcellular APP trafficking, we performed a confocal imaging analysis on NGF-treated (100 ng mL\textsuperscript{-1}, 1 h) septal neurons. Neurons were double-immunolabelled with several markers for different intracellular compartments, such as giantin for the Golgi, Rab5 for early endosomes, and LAMP1 for late endosomes/lysosomes. Upon NGF stimulation, we found an increase in APP localization at the plasma membrane (Fig. 6E and inset of Fig. 6E, asterisks) and in cell soma, in particular in the Golgi, as quantified with the Imaris 7.6 software (130 ± 9.7% of CTR, *P < 0.01; Fig. 6F; lower magnification in Fig. 5B).

To address APP-BACE interaction in the Golgi, where accumulates upon NGF treatment, we performed a triple immunostaining for APP (red)/BACE\textsubscript{1} (green)/giantin (purple) in septal neurons (Fig. 6G). BACE\textsubscript{1} immunoreactivity was found to be diffuse and widely distributed throughout the cytoplasm of both unstimulated (CTR) and NGF-treated (NGF) septal neurons. Moreover, APP and BACE immunofluorescent signals were found to colocalize in the Golgi, although at a low extent (Fig. 6G). Pearson's correlation between APP and BACE within the Golgi

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has been measured by creating a 3D mask of the giantin-positive volume, with the help of Volocity 6.0 (PerkinElmer). The 3D rendering of APP (red)- and BACE (green)-positive voxels within the giantin (purple)-positive volume in CTR and NGF-treated neurons has been reported in the inset 3D (Fig. 6G). The results of this analysis indicate a moderate correlation of APP and BACE colocalization in the Golgi of CTR neurons.
(r = 0.579 ± 0.084), while poor correlation was found upon NGF treatment (r = 0.323 ± 0.049; P < 0.05). Of note, BACE localization in the Golgi was not affected by NGF treatment (P = 0.314; r = 0.363 ± 0.02), as compared to unstimulated neurons (r = 0.315 ± 0.05).

APP trafficking to endosomes and lysosomes was also analysed. The amounts of APP localized to the early (Rab5; Fig. 53A) endosomes were comparable (P = 0.7) between the CTR (r = 0.6 ± 0.2) and NGF-treated septal neurons (r = 0.7 ± 0.1). APP localization to lysosomes (LAMP1; Fig. 53C) was present at a similar extent (P = 0.99) in CTR (r = 0.5 ± 0.2) and NGF (r = 0.5 ± 0.1)-treated neurons, in our experimental conditions.

Taken together, our data show that NGF treatment favours the APP interaction with TrkA, promotes APP trafficking towards the Golgi, reduces APP exposure to BACE cleavage, and finally results in the reduction of sAPPβ, CTFβ and amyloid-beta (1-42) levels in the basal forebrain neurons.

**Discussion**

Our study aimed to investigate NGF control of APP phosphorylation and metabolism in the basal forebrain under physiological conditions, and following NGF system perturbation described in AD. NGF is a pleiotropic factor for cholinergic neurons of the basal forebrain, the main target of Alzheimer’s disease (AD) pathology. Septal cholinergic neurons have been shown to accumulate intracellular amyloid during aging and AD, thus affecting neuronal activity and cognition in rodents, monkeys and humans (Baker-Nigh et al., 2015; Norvin et al., 2015). The data presented in this study indicate that NGF downregulates the phosphorylation of APP at the neuron-specific residue T668, promotes binding with TrkA, and favours APP trafficking to the Golgi, at the expense of BACE interaction and cleavage.

In detail, our findings indicate that ectopic NGF stimulation (Fig. 1A–D) and NGF withdrawal (Fig. 1E–H), control basal APPpT668 levels in neurons. Moreover, our results correlated the effect of NGF on the APPpT668 level with a specific reduction in JNKs (p54) activation. In fact, APPpT668 is affected both by pharmacological inhibition of JNKs in SH slices and by genetic ablation of the three JNK isoforms in KO mice, and of JNK2 in particular. Of interest to this study, JNK3 activity has been found to increase by 35% in FAD models, which induces amyloidogenesis and neuronal apoptosis, affects cognition (Yoon et al., 2012), and controls APP localization (Muresan & Muresan, 2005).

As expected, also CDK5 inhibitors reduced the APPpT668 level in the SH slices (Fig. 2A). However, NGF did not affect the expression levels of CDK5 or its activator p35 (Fig. 2C). Conversely, NGF was able to significantly reduce JNK activation in both the SH slices (Fig 2D–F) and septal neurons (Fig 2G,H). In particular, NGF specifically affected phosphorylation of the JNK(p54) band (Fig. 2F and H). Thus, a crucial role for NGF modulation of JNK(p54) activation may be in APP phosphorylation, metabolism and the related cognition in mammals.

Further analyses focused on the involvement of the SH2-containing sequence C (ShcC) in these events. ShcC docking to TrkA is an early downstream event induced by NGF and causes the p35/akt-mediated inhibition of JNK (Pellicci et al., 2002). Thus, we analysed JNK activation and the APPpT668 level in hippocampal extracts from ShcC KO mice and found a dramatic rise of JNK phosphorylation (Fig. 3A,C) and a concomitant increase in the APPpT668 level compared with age-matched WT mice (Fig. 3B,C). Furthermore, we found a significant elevation in the total CTF level, and in particular CTFβ (Fig. 3D), while full-length APP (Fig. 3B) and BACE (Fig. 3E) were unaffected (Fig. 3B,D). Endogenous amyloid-beta (1-42) levels were also significantly augmented in ShcC null mice, as measured by a rodent-specific ELISA (Fig. 3D). As a further confirmation of the key role of ShcC in the anti-amyloidogenic NGF pathway, NGF failed to modulate JNK and APP phosphorylation in SH slices from ShcC KO mice (Fig. 3H,I), as well as to reduce CTFβ levels (Fig. 3H). Thus, ShcC is necessary for negative regulation of JNK mediated APP phosphorylation, and the control of the consequent CTFβ and amyloid-beta generation in vivo.

Our data also indicate that TrkA colocalizes with APP in unstimulated septal neurons (CTR; Fig. 4A) and interacts with APP in both hippocampal tissues (Fig. 4C) and SH slices (Fig. 4D). These results are consistent with previous studies demonstrating the APP–TrkA interaction in TrkA-transfected 293T cells and hippocampal neurons in vitro and in vivo (Tarr et al., 2002; Matrone et al., 2011). Moreover, using both confocal imaging (Fig. 4A) and IP analysis (Fig. 4D), we found that the APP association with TrkA increases after NGF treatment. However, TrkA does not colocalize with APPpT668 under basal conditions or after NGF treatment (Fig. 4B,C). Indeed, the lack of an APPpT668–TrkA interaction was observed in both the control and NGF-treated SH slices, despite the increase in the amount of total APP bound to TrkA after NGF stimulation (Fig. 4D).

Previous studies have shown that transgenic mice lacking the APP–TrkA interaction demonstrate forebrain damage and cognitive deficits (Matrone et al., 2012). Based on the effect of APP phosphorylation on the APP–TrkA binding, the APP–TrkA interaction was investigated in the AD human brain where elevated APPpT668 levels have been reported (Lee et al., 2003; Chang et al., 2006). An increase in the APPpT668 level in patients with AD was also observed in our analysis (Fig. 5A–C). Consistent with our hypothesis, the APP–TrkA interaction was found to be significantly affected (Fig. 5D), and in some cases lost (Fig. 5E), in the AD hippocampus. Interestingly, the AD cerebellum (Fig. 5E), a relatively unaffected tissue in AD (Andersen et al., 2012), and the HD hippocampus (Fig. 5D) maintained the APP–TrkA interaction. These data suggest that reduced APP–TrkA binding is a good correlate of AD pathology and not just a general marker of degeneration.

Next, we investigated neuronal APP trafficking and BACE binding/ cleavage upon NGF stimulation in primary septal neurons. We found that NGF treatment favoured binding of APP to its specific receptor TrkA at the expense of BACE; (Fig. 6A) and stimulated APP trafficking to the Golgi (arrows, Fig. 6E,F), where APP–BACE colocalization is quite weak in control neurons and further reduced upon NGF treatment of septal neurons (Fig. 6G; inset 3D). These results are in line with recent papers showing that although APP and BACE colocalization is detected in the Golgi, the main neuronal site for APP shedding are the endosomes (Das et al., 2013, 2016; Buggia-Prevot et al., 2014; Vassar et al., 2014).

Accordingly, NGF treatment promoted the anti-amyloidogenic processing of APP in the basal forebrain, as demonstrated by the increase in sAPPα and the concomitant decrease in sAPPβ (Fig. 6B,C), CTFβ (Fig. 6B, C) and amyloid-beta 42 (Fig. 6D) levels in SH slices. Further, these events occurred independently of both BACE and ADAM17 levels (Fig. 6B,C), which were unaffected (Fig. 6C).

Intriguingly, it has been reported that APP is transported using the activated NGF/TrkA-Shc signalling system in neurons (Heese et al., 2004). Therefore, the idea of a NGF-driven trafficking of APP in septal...
primary neurons is consistent with previous and recent studies on neuronal APP trafficking and deserves future investigation.

Conclusions

All together, these data suggest that NGF directly controls the pattern of APP phosphorylation, its trafficking, interaction with and cleavage by BACE, which ultimately reduces CTFβ and amyloid generation in basal forebrain neurons.

Interestingly, APP processing was found to be normal in APP(T668A) mice, bearing a Thr to Ala point mutation (Sano et al., 2006), suggesting that APPpT668B reduction is not sufficient per se to affect APP processing and that instead an active NGF/TrkA–ShcC signalling system is required for promoting APP-TrkA interaction and anti-amyloidogenic APP processing. Further, the differential binding of APP to its interactors, namely TrkA and BACE, impacting APP trafficking and processing in basal cholinergic neurons is of potential interest for AD pathology, where APP–TrkA interaction was found to be specifically affected.

Therefore, novel strategies aimed at stimulating ShcC signalling and/or promoting TrkA–APP binding should be considered as promising therapeutic tools for early neurodegeneration in AD.

Experimental procedures

Reagents and antibodies

Murine NGF was purified from submaxillary glands (Bocchini & Angeletti, 1969). NGF from Xiangen Bioway (Biotec Co., Ltd., Xiamen, Fujian, China) was also used in the study. Roscovitine was purchased from SIGMA (St. Louis, MO, USA; R7772), SP600125 from Calbiochem (San Diego, CA, USA; 420119). Rabbit anti-TrkA (WB 1:1000, IF 1:100) was a generous gift from Prof. L. Reichardt. The polyclonal anti-NGF antibody used for NGF neutralization was kindly provided by D. Mercanti (CNR-IBCN, Rome). The commercial antibodies used are listed in Table 1.

SH slices

Young adult C57/Bl6 male mice 3–5 months of age were used for the preparation of SH slices. The animals were sacrificed by cervical dislocation. The SH region was removed and cooled in ice-cold oxygenated artificial cerebrospinal fluid (aCSF; 126 mM NaCl, 2.5 mM KCl, 1.2 mM NaH2PO4, 24 mM NaHCO3, 2.4 mM CaCl2, 1.3 mM MgCl2 and 10 mM glucose). Next, 300-μm-thick slices were cut using a McIlwain tissue chopper (Ted Pella, Redding, CA, USA) and incubated in aCSF (1 h) under appropriate oxygenation and pH conditions (95% O2, 5% CO2). Slices were then exposed to the different treatments, snap frozen and stored at −80 °C until use.

NGF withdrawal experiments were performed by incubation at different time points (3, 6 h) of control SH slices with neutralizing titres of anti-NGF antibody, as previously described (Matrone et al., 2008b).

Septal neurons primary culture

Septal neurons were prepared from embryonic day 17/18 (E17/18) pregnant Wistar rats or embryonic day 16 (E16) pregnant C57/Bl6 mice, as previously described (Hartikka & Hefti, 1988). They were seeded as 104 cells on glass coverslips in 24-well plates (BD Falcon; 351147) for biochemistry analyses and 9 × 106 cells on poly-l-lysine (SIGMA)-coated plates (BD Falcon; 353001) for immunofluorescence analyses. The dissociated cells were plated in Neurobasal medium supplemented with 2% B27 (Invitrogen Inc., Carlsbad, CA, USA) for 10 days (37 °C, 5% CO2) and then used for the experiments. At day 10, more than 95% cells showed positive staining for ChAT and around 90% of ChAT-positive cells were also TrkA positive (data not shown).

PC12 cell culture

PC12 cells were cultured in RPMI medium containing 10% horse serum and 5% FBS. NGF deprivation was performed as previously described (Matrone et al., 2008b). PC12 cells were incubated with NGF (100 ng mL−1) for 10 days until fully differentiated and then used as a control (CTR) or kept in serum-free and NGF-free medium for 30 min or 3 h (NGF-dep 30°; NGF-dep 3 h). Plates were frozen at −20 °C until use.

Rodent strains

All animals were handled in compliance with the national (D.Lgs26/2014) and international guidelines for animal welfare (2010/63/EU). All efforts were made to minimize the number of animals used and suffering. C57/Bl6J mice and Wistar rats (Harlan Laboratories Ltd., Füllinsdorf, Switzerland) were housed at the IBCN-CNR (Italy) animal facility. Brain tissues from male ShcC KO mice (Sakai et al., 2000), maintained in a C57/Bl6 background, were collected from mice housed at the Animal Facility of the ‘University of Siena’ (Italy). The JNK1-3 KO mouse hippocampal tissues were provided by M.F. (Nisticò et al., 2015). Age-matched C57/Bl6J mice and/or tissues were used as controls in this study.

Human brain samples

Brain tissue samples from human subjects were provided by the Regional Neurogenetic Center (CRN, ASP CZ) of Lamezia Terme, Italy. All brain

| Antibody name | Company and code | Working dilution |
|---------------|------------------|------------------|
| APP (22c11)   | MILLIPORE (Temecula, CA, USA) | WB 1:10000, IF 1:200 |
| APP C-ter (Y188) | ABCAM (Cambridge, UK) | AB23126 | IF 1:100 |
| BACE1         | MILLIPORE (Temecula, CA, USA) | AB35308 | WB 1:500 |
| ChAT          | MILLIPORE (Temecula, CA, USA) | AB144P | WB 1:1000 |
| APP C-ter     | SIGMA (St. Louis, MO, USA) | A8177 | WB 1:1000 |
| APPB          | SIGMA (St. Louis, MO, USA) | A9103 | WB 1:1000, IF 1:100 |
| p35           | SIGMA (St. Louis, MO, USA) | C6118 | WB 1:1000 |
| β-actin-HRP   | IBL INTERNATIONAL (Hamburg, DE) | 11088 | WB 1:500 |
| sAPPx         | IBL INTERNATIONAL (Hamburg, DE) | 18957 | WB 1:2000 |
| APPB          | ABCAM (Cambridge, UK) | AB18211 | IF 1:100 |
| pTrkA         | SIGMA (St. Louis, MO, USA) | A3854 | WB 1:1000 |
| pJNK1/2/3     | SIGMA (St. Louis, MO, USA) | A3854 | WB 1:1000 |
| pJNK1/2/3      | SANTA CRUZ (Santa Cruz, CA, USA) | sc7268 | WB 1:1000 |
| Trk            | SANTA CRUZ (Santa Cruz, CA, USA) | sc411 | IF 1:100 |
| Rab9           | SIGMA (St. Louis, MO, USA) | A8717 | WB 1:1000 |
| Rab8           | SIGMA (St. Louis, MO, USA) | A8717 | WB 1:1000 |
| LAMP1          | STRESSGEN (Collegeville, PA, USA) | ADIVAMEN001 | IF 1:100 |
donors or their legal tutors gave written informed consent during their lifetime, and the protocol was approved by the local ethics committee. The study was conducted with brain samples from 3 unrelated patients affected by familial AD (FAD) and bearing the Met146Leu mutation of the Presenilin1 gene. The mean age of death of the patients was 48 ± 8.2 years, and they were at the terminal stage of dementia (stage 5, according to the Clinical Dementia Rating scale). For all patients, the Braak staging for AD pathology (AD-related neurofibrillary pathology) was VI. Brain tissues from three human subjects (CTR; 60 ± 7 years of age) without neurological diseases were used as control. Samples from three subjects affected by other neurological diseases (OND) were also analysed in this study. Specifically, the three OND subjects were affected by spinocerebellar ataxia 17 (SCA17), Huntington’s disease (HD), and the third subject died from accidental intracerebral bleeding in the left brain hemisphere (the sample was taken from the counter-lateral right brain hemisphere). Histopathological lesions suggestive of AD (neurofibrillary pathology, deposition of Aβ protein) were excluded. In details, tissues were sampled from hippocampal gyrus (CA1, CA2, CA3 and dentate gyrus, excluding parahippocampal gyrus and subiculum) and from cerebellar hemispheres. The brain samples were collected at 4–36 h postmortem and stored at −80 °C until use.

**WB analysis**

Brain slices, tissues samples and cultured neurons were digested in a RIPA buffer with phosphatase and protease inhibitors (Pierce) and centrifuged (10,000 rpm, 20°). The supernatants were collected, and the amount of total protein was determined (Quick Start Bradford Dye Reagent, Biorad, Hercules, CA, USA). Each sample (40 μg) was separated by SDS-PAGE in 4–12% Bis-Tris gels (Invitrogen), transferred to nitrocellulose membranes (0.45 μm, GE healthcare, Little Chalfont, UK) and incubated for 1 h at room temperature with 5% nonfat dry milk in TBS-T (10 mM Tris, pH 7.5, 100 mM NaCl and 0.1% Tween-20). The overnight incubation with primary antibody (4 °C) was followed by incubation with the appropriate HRP secondary antibody (1:3000, Pierce, 1 h, RT) and the ECL method (GE Healthcare). The films were digitalized using a professional scanner (HP) and quantified using IMAGE 1.63 software (NIH). Omission of the primary antibody was routinely performed as control for antibody specificity (not shown).

**Immunoprecipitation**

A total of 1 mg of protein extract was added to Dynabeads-Protein G and eluted with 0.1 m citrate buffer, pH 2.3, according to the manufacturer’s instructions (Invitrogen). IP of APP was performed using rabbit anti-APP C-terminal antibody (Sigma; 10 μg/50 μl beads). IP of TrkA was performed using specific anti-TrkA antibody (4 μL/50 beads). There were no specific bands detected by WB, when the same amount of normal rabbit IgG (Santa Cruz, sc-2027) was used as isotypic control for the IP (Fig. S1E).

**Immunofluorescence and confocal imaging**

Primary septal cell cultures were fixed for 20 min in PBS containing 4% paraformaldehyde (PFA), permeabilized and quenched. Statistical analysis was performed and representative images were chosen among 50 primary neurons from multiple assays.

ShCC KO (n = 3) and age-matched control mice (n = 3) were anesthetized with avertin (500 mg kg⁻¹, i.p.; Sigma-Aldrich) and perfused transcardially with PBS, pH 7.4, followed by 4% PFA in PBS. Then, the brains were collected, postfixed overnight (4% PFA, 4 °C) and cryoprotected in 30% (w/v) sucrose in PBS 0.1 M (4 °C) until they sank to the bottom, as previously described (Müller et al., 2012). Coronal brain sections (25 μm thick) were cut on a cryostat (Leica Microsystems) and kept floating in PBS at 4 °C until use. Sections spanning from bregma 0.74 mm to 1.34 mm (medial septum) and from bregma −1.34 to −2.3 mm (hippocampus) were used for this analysis (Paxinos & Franklin, 2001).

Coronal brain sections or fixed primary neurons were incubated with ammonium chloride (50 mM, RT) for 30 min, to quench autofluorescence and then blocked with 10% normal donkey serum (code: 01700121, Jackson Lab, Bar Harbor, ME, USA). The overnight incubation (4 °C) with primary antibody was followed by incubation with the appropriate combination of secondary antibodies. In particular, donkey anti-mouse-546 and donkey anti-rabbit-488, or donkey anti-mouse-488 and donkey anti-rabbit-546 were used for double immunofluorescence; donkey anti-mouse-647, donkey anti-goat-546 and donkey anti-rabbit-488 were used for triple immunostaining (Life Technologies, 1 h, RT, Carlsbad, CA, USA). Finally, DAPI (Life Technologies) was used for counterstaining and Prolong Gold (Life Technologies) for mounting on Superfrost Plus slides. Confocal microscopy was performed with the laser scanning confocal microscope TCS SP5 (Leica Microsystems, Mannheim, Germany) using a 40X (NA = 1.25) and a 63X (NA = 1.4) oil-immersion lens. An UV diode laser operating at 405 nm, an Argon laser at 488 nm and HeNe lasers at 543 or 633 nm were used as excitation sources. Confocal Z-stacks were collected at 5X zoom at 0.29 mm intervals in a 4 μm total optical depth. Images for direct comparison were collected using the same parameters and were analysed with the help of IMARIS 7.6 (Bitplane, Zurich, Switzerland) or VOLOCITY 6.0 (PerkinElmer, Waltham, MA, USA) softwares. Pearson's correlation coefficient (r) was used for correlation analysis, with an r value between +0.5 and 1.0 indicating a high strength of correlation. Omission of the primary antibody was routinely performed as control for antibody specificity (not shown). The specificity of the antibody against BACE1 was tested by pre-incubation of the primary antibody with the BACE peptide (PEP 0077, Thermofisher, Waltham, MA, USA), resulting in no staining (not shown).

**Amyloid-beta (1-42) ELISA**

To measure endogenous amyloid-beta (1-42) levels, a mouse/rat-specific and high-sensitive enzyme immunoassay kit was used, according to the manufacturer’s protocol (code: re45721; IBL Int.).

**Statistical analysis**

All experiments using SH slices and primary neurons were performed at least four times independently, each in triplicate. Tissues from four ShCC KO and four control mice were analysed. The graphs were generated using PRISM (GraphPad Software, Inc., San Diego, CA, USA), and the data are presented as the mean ± standard error. Student’s t-test or ANOVA followed by the Tukey–Kramer post hoc was used to analyse the data depending on the number of variables and groups (Statview-SAS, Cary, NC, USA). A P value ≤ 0.05 was considered statistically significant.

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Supporting Information

Additional Supporting Information may be found online in the supporting information tab for this article.

Fig. S1 NGF reduces pAPP (A) and pJNK (B) levels in rat primary septal neurons (E18, DIV10).

Fig. S2 Low magnification confocal images of APP, APP pT668 and TrkA localization in septal neurons.

Fig. S3 Low magnification confocal images of APP trafficking under NGF treatment showing that NGF (100 ng mL⁻¹, 1 h) affects Golgi accumulation of APP but not endosomal or lysosomal APP localization in primary septal neurons (E17, DIV10).