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Norepinephrine metabolite DOPEGAL activates AEP and pathological Tau aggregation in locus coeruleus

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Aberrant Tau inclusions in the locus coeruleus (LC) are the earliest detectable Alzheimer’s disease–like (AD-like) neuropathology in the human brain. However, why LC neurons are selectively vulnerable to developing early Tau pathology and degenerating later in disease and whether the LC might seed the stereotypical spread of Tau pathology to the rest of the brain remain unclear. Here, we show that 3,4-dihydroxyphenylglycolaldehyde, which is produced exclusively in noradrenergic neurons by monoamine oxidase A metabolism of norepinephrine, activated asparagine endopeptidase that cleaved Tau at residue N368 into aggregation- and propagation-prone forms, thus leading to LC degeneration and the spread of Tau pathology. Activation of asparagine endopeptidase–cleaved Tau aggregation in vitro and in intact cells was triggered by 3,4-dihydroxyphenylglycolaldehyde, resulting in LC neurotoxicity and propagation of pathology to the forebrain. Thus, our findings reveal that norepinephrine metabolism and Tau cleavage represent the specific molecular mechanism underlying the selective vulnerability of LC neurons in AD.

Introduction
Alzheimer’s disease (AD) is an insidious, debilitating illness characterized by amyloid-β (A-β) plaques and Tau neurofibrillary tangles (NFTs), as well as neuronal cell death and neuroinflammation. Despite intense research into the pathologic hallmarks of AD, no disease-modifying therapies have emerged (1), and recent research efforts are focused on early AD detection and treatment. One area of interest in early AD involves the locus coeruleus (LC), the major brainstem noradrenergic nucleus that innervates and supplies norepinephrine (NE) to the forebrain and regulates attention, arousal, cognition, and stress responses (2). Hyperphosphorylated Tau, a “pretangle” form of the protein prone to aggregation, appears in the LC before any other area of the brain and is now considered the earliest detectable AD-like neuropathology (3–7). Moreover, degeneration of the LC is a ubiquitous feature of mid-to late-stage disease, contributing to the progression of forebrain pathology and cognitive impairment (6, 8–12), whereas restoration of noradrenergic signaling can ameliorate AD-like symptoms in animal models (13, 14). The progression of AD pathology follows a remarkably systematic pattern across individuals, and aberrant forms of Tau are capable of neuron-to-neuron propagation via a prion-like process of corruptive templating (15). Hyperphosphorylated Tau appears exclusively in the LC in Braak stage 0, spreads to the entorhinal cortex (Cx) in Braak stages I and II, then propagates to the hippocampus and frontal Cx in stages III–VI (3). Because hyperphosphorylated Tau can first be detected in the LC, and the LC sends dense projections to other vulnerable brain regions that display early Tau pathology (e.g., the transentorhinal Cx and hippocampus), some suggest that the LC might be one of the origins of Tau neuropathology in AD (4, 5). Although several theories have emerged to explain the selective vulnerability of LC neurons in AD, such as their unique anatomical, electrophysiological, morphological, and biochemical characteristics (7, 16), the underlying molecular mechanisms have yet to be identified.

Asparagine endopeptidase (AEP; gene name LGMN) is a lysosomal endopeptidase that specifically cleaves its substrates after asparagine under conditions of acidosis (17). Recently, we reported that AEP acts as a delta-secretase that cleaves both amyloid precursor protein (APP) and Tau in an age-dependent manner in mouse and human AD brains (18, 19). AEP cuts APP at the N373 and N585 residues in the ectodomain and facilitates A-β secretion, resulting in hyperphosphorylation, aggregation, and NFT formation (18). Of note, the AEP-generated Tau 1-368 fragment is highly neurotoxic. Genetic deletion or pharmacological inhibition of AEP in transgenic mice that overexpress mutant human APP or aggregation-prone mutant human Tau prevents APP N585 and Tau N368 cleavage, ameliorates A-β and Tau pathology, and reverses synaptic plasticity and cognitive deficits (18, 20). Thus, AEP plays a critical role in regulating A-β and Tau pathogenesis. Most recently,
we reported that AEP is activated by 3,4-dihydroxyphenylacetaldehyde (DOPAL), a highly toxic and oxidative dopamine (DA) metabolite, in the substantia nigra (21). Active AEP subsequently cleaves α-synuclein at N103 and promotes its aggregation and dopaminergic neuronal loss, leading to motor dysfunction in animal models of Parkinson’s disease (PD) (22).

Analogous to toxic DA metabolites killing substantia nigra pars compacta neurons in patients with PD, the noradrenergic phenotype of LC neurons itself may contribute to the vulnerability of these cells in AD. NE is converted into 3,4-dihydroxyphenylglycolaldehyde (DOPEGAL) by monoamine oxidase A (MAO-A) during the normal life cycle of catecholamine production and transmission, but is increased in the degenerating LC neurons of patients with AD (23–25). Injection of DOPEGAL into rodent brains elicits adrenergic neuronal loss (26), and DOPEGAL toxicity is likely due to the generation of free radicals and activation of mitochondrial permeability transition (27). The activation of AEP by DOPAL suggested to us that it might also be activated by DOPEGAL, thus triggering a cascade of events leading to Tau cleavage, hyperphosphorylation, aggregation, and neurotoxicity in LC neurons.

**Results**

**DOPEGAL triggers Tau aggregation in vitro and in cells.** To examine whether DOPEGAL directly modifies Tau, we incubated recombinant WT Tau with DOPEGAL for 24 hours. We found DOPEGAL induced Tau aggregation, indicated by the accumulation of high-molecular-weight bands, in a concentration-dependent manner. As expected, AEP-truncated Tau N368 recombinant proteins were more prone to aggregation than full-length Tau (Figure 1A). To explore the specificity of DOPEGAL, we tested various catecholamines and their oxidative metabolites (500 μM). Compared with vehicle control, DOPEGAL strongly provoked Tau aggregation, while NE, DA, and DOPAL instead caused Tau protein degradation (Figure 1B). Time course and dose-response assays demonstrated that both DA and NE triggered dose- and time-dependent degradation of Tau, whereas DOPAL elicited Tau aggregation at lower doses (2.5–25 μM) but promoted degradation at higher doses (Supplemental Figure 1, A–C; supplemental material available online with this article; https://doi.org/10.1172/JCI130513DS1). To investigate whether DOPEGAL elicits Tau fibrillization in vitro, we monitored the fluorescent intensity of recombinant Tau in the presence of Thioflavin T (ThioT) (30 μM). Compared with vehicle control, DOPEGAL strongly augmented Tau fibrillation over time, as revealed by the high-intensity ThioT fluorescence of Tau preformed fibrils (PFFs), while NE and DOPAL inhibited Tau fibrillation (Figure 1C). Transmission electronic microscopy analysis confirmed that Tau PFFs formed tight fibrillar structures following DOPEGAL exposure (Supplemental Figure 1D).

To assess whether DOPEGAL provokes Tau aggregation in intact cells, we transfected noradrenergic-like SH-SY5Y cells with WT human Tau, followed by treatment with different catecholamines or their oxidative metabolites for 24 hours. Immunoblotting revealed that DOPEGAL (but not DA or NE) produced demonstrable Tau aggregation, which was in alignment with its hyperphosphorylation (AT8-positive) status (Figure 1D, top 2 panels). The lack of an effect for NE is consistent with the limited conversion capacity for exogenous NE in SH-SY5Y cells due to the low expression of MAO-A. We next determined whether DOPEGAL, like DOPAL, activates AEP. We found DOPEGAL upregulated total AEP levels and its proteolytic activation, as well as the abundance of the AEP cleavage product Tau N368 (Figure 1D, bottom 2 panels, Figure 1E) and induced SH-SY5Y cell death (Figure 1F–H). DOPAL was less effective in these measures, while NE and DA failed to activate AEP or cause toxicity. To ascertain whether Tau is required for DOPEGAL-elicited cell death, we prepared primary cortical neurons from neonatal WT and Tau knockout (Tau−/−) mice and treated them with DOPEGAL. As expected, DOPEGAL triggered demonstrable Tau hyperphosphorylation, aggregation, and N368 cleavage in WT, but not Tau−/−, neurons (Figure 1I). Importantly, although AEP was similarly activated by DOPEGAL in both WT and Tau−/− neurons (Figure 1J), its toxicity was significantly attenuated in Tau−/− neurons (Figure 1K–M), suggesting that Tau is necessary for the full expression of DOPEGAL-induced cell death.

**NE oxidation by MAO-A facilitates AEP-cleaved Tau N368 cytotoxicity.** Oxidative deamination of NE by mitochondrial MAO-A generates DOPEGAL and H2O2, leading to oxidative stress (28, 29). To test whether MAO-mediated metabolism of NE results in oxidative stress and AEP activation, we transfected SH-SY5Y cells (30) with MAO-A or MAO-B. We found that MAO overexpression increased DOPEGAL levels, AEP enzymatic activity, and Tau N368 cleavage (Supplemental Figure 2, A–D), which was mimicked by H2O2 and prevented by the MAO-A inhibitor clorgyline, indicating the importance of oxidative stress (Supplemental Figure 2, E–G). Interestingly, H2O2 also increased MAO-A expression, which was blunted by clorgyline. To evaluate the contribution of NE metabolism, we transfected SH-SY5Y cells with siRNA against dopamine β-hydroxylase (DBH), which is required for NE synthesis (31). DBH depletion reduced AEP activity, diminished Tau N368 cleavage, and partially ameliorated the deleterious effects of H2O2 (Supplemental Figure 2, H and I). Combined, these results suggest that oxidative metabolism of NE to DOPEGAL by MAO activates AEP and Tau N368 cleavage in catecholaminergic SH-SY5Y cells.

To explore the neurotoxicity of AEP-cleaved Tau, we infected SH-SY5Y cells with AAV encoding full-length WT Tau, P301S Tau, the N368 Tau fragment, or uncleavable P301S/N255/368A Tau, and determined cell death using TUNEL staining and lactate dehydrogenase (LDH) assay (Supplemental Figure 2, J–M). WT Tau promoted cell death, which was escalated by the AEP-cleaved Tau N368 fragment. Notably, N368 Tau displayed neurotoxicity comparable with Tau P301S, whereas prevention of AEP cleavage blunted Tau P301S toxicity. These results indicate that AEP cleavage contributes to Tau neurotoxicity.

To ascertain whether DOPEGAL regulates Tau neurotoxicity, we overexpressed MAO-A or MAO-B in SH-SY5Y cells in combination with AAV-Tau. TUNEL and LDH analysis revealed that both MAO-A and MAO-B overexpression significantly activated AEP, increased Tau N368 cleavage, and magnified Tau neurotoxicity as measured by TUNEL, LDH, and loss of the catecholaminergic marker tyrosine hydroxylase (TH) (Supplemental Figure 3). By contrast, preventing DOPEGAL production by siRNA knockdown of DBH attenuated AEP activation and Tau-triggered cell death.
Figure 1. The NE metabolite DOPEGAL triggers Tau aggregation in vitro. (A) Recombinant Tau (rTau) or rTau N368 (1 μg) was incubated with DOPEGAL (0, 5, 50, or 500 μM) for 24 hours, and aggregation was detected by Western blot using Tau5 antibody. (B) rTau (1 μg) was incubated with vehicle, DA, NE, DOPAL, or DOPEGAL (500 μM) for 24 hours, and aggregation was detected by Western blot using Tau5 antibody. (C) rTau (1 μg) was incubated with vehicle or DOPEGAL, and ThioT assay was used to evaluate the kinetics of aggregation of Tau over 120 hours. SH-SY5Y cells were infected with hTau virus, then exposed to DA, NE, DOPAL, or DOPEGAL (60 μM) for 24 hours. (D) Western blot analysis was conducted with cell lysates using antibodies against AEP and different forms of Tau. (E) AEP activity was analyzed by the enzymatic assay. Data are shown as mean ± SEM (n = 3 per group). *P < 0.05 and **P < 0.01, by 1-way ANOVA. Cell death was analyzed by LDH assay (F) and propidium iodide staining (G–H). Scale bar: 50 μm. Data are shown as the mean ± SEM (n = 3 per group). *P < 0.05 and **P < 0.01, by 1-way ANOVA. Primary cortical neurons were cultured from Tau+/+ and Tau−/− mice and exposed to vehicle or DOPEGAL (60 μM) for 24 hours. (I) Western blot analysis was conducted with cell lysates using antibodies against AEP and different forms of Tau. (J) AEP activity was analyzed by the enzymatic assay. Data are shown as the mean ± SEM (n = 3 per group). *P < 0.05 and **P < 0.01, by 2-tailed t test. Cell death was analyzed by LDH assay (K) and propidium iodide staining (L and M). Scale bar: 50 μm. Data are shown as the mean ± SEM (n = 3 per group). *P < 0.05 and **P < 0.01, by 2-way ANOVA.
cleavage, and AT8 immunoreactivity in the presence of elevated MAO-A (Supplemental Figure 4, F and G). Interestingly, both L-DOPA and L-DOPS, but not ascorbic acid, exhibited cytotoxicity on their own, which was facilitated by MAO-A overexpression (Supplemental Figure 4H). The limited cytotoxicity of ascorbic acid may be attributed to its antioxidant activity.

DOPEGAL is metabolized in vivo via reduction by aldehyde reductase (AR) and further oxidation by aldehyde dehydrogenase (ALDH). To determine whether accumulation of DOPEGAL via blockade of its metabolism augments its cytotoxicity, we treated SH-SY5Y cells with the AR inhibitor imirestat, the ALDH inhibitor daidzein, or both drugs in the presence or absence of MAO-A overexpression. Although no treatment was effective on its own, imirestat and the mixture of both inhibitors augmented AEP activation, Tau N368 cleavage, and AT8 abundance in combination with MAO-A (Supplemental Figure 4, I and J). LDH assay indicated that the inhibitor cocktail modestly stimulated cell death without MAO-A overexpression, while either imirestat or imirestat plus daidzein escalated MAO-A cytotoxicity (Supplemental Figure 4H).

Figure 2. Tau is cleaved at N368 and hyperphosphorylated in the LC region of transgenic AD mouse models and human postmortem AD brains. WT, 3xTg, and P301S mouse brains from various ages, as well as postmortem human control and AD brains, were examined for Tau pathology in the LC by immunohistochemistry. (A) Representative images of staining for TH (green), AT8 or Tau N368 (red), and DAPI (blue). Scale bar: 100 μm (×60 panel is 20 μm). (B–D) Quantification of AT8 and Tau N368+ cells in 3XTG (B), Tau P301S mice (C), and human brains of controls and patients with AD (D). Data are shown as mean ± SEM (n = 5–6 per group). *P < 0.05 and **P < 0.01, by 2-way ANOVA.

death. These findings reveal that a proportion of Tau neurotoxicity is NE-dependent.

Our data support a model whereby MAO-A oxidizes NE into DOPEGAL and elicits oxidative stress, triggering AEP activation and Tau N368 cleavage and cytotoxicity. To further explore this idea, we overexpressed MAO-A and MAO-B into SH-SYSY cells and interrogated their roles in AEP activation in the presence of specific inhibitors. We confirmed that overexpression of either MAO-A or MAO-B substantially provoked AEP activation, and blockade of these enzymes by small molecular inhibitors attenuated AEP enzymatic activity, Tau hyperphosphorylation, and N368 cleavage (Supplemental Figure 4, A–E). To test whether elevation of NE in SH-SYSY cells facilitates MAO-A cytotoxicity, we transfected cells with MAO-A, followed by treatment with ascorbic acid, which stimulates NE biosynthesis (32), the DA and NE precursor L-3,4-dihydroxyphenylalanine (L-DOPA), or the synthetic NE precursor L-3,4-dihydroxyphenylserine (L-DOPS). Although these compounds alone barely activated AEP, both L-DOPS and ascorbic acid significantly increased AEP activation, Tau N368 cleavage, and AT8 immunoreactivity in the presence of elevated MAO-A (Supplemental Figure 4, F and G). Interestingly, both L-DOPA and L-DOPS, but not ascorbic acid, exhibited cytotoxicity on their own, which was facilitated by MAO-A overexpression (Supplemental Figure 4H). The limited cytotoxicity of ascorbic acid may be attributed to its antioxidant activity.

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AD brains (Figure 3B). These findings demonstrate that activated AEP is abundant in the diseased LC, where it is associated with Tau N368, hyperphosphorylated Tau, and aggregation.

NE is required for AEP activation and Tau N368 cleavage in the LC of Tau P301S mice. To investigate whether DOPEGAL is implicated in Tau pathology and LC neuronal degeneration, we crossed Tau P301S transgenic mice with DBH knockout (DBH–/–) mice that lack NE completely (and thus cannot produce DOPEGAL) and found Tau hyperphosphorylation and N368 cleavage at different ages. DBH+/– littermates with normal NE levels were used as controls (33). At 3 months, AT8 and Tau N368 immunoreactivity in the LC was minimal in both Tau P301S/DBH+/– and Tau P301S/DBH–/– mice. Although the abundance of pathological Tau escalated in both genotypes at 6 to 9 months, AT8, Tau N368, and Gallyas-Braak silver staining were significantly reduced in Tau P301S/DBH–/– mice (Figure 4, A–C). Consistent with our finding that suppression of NE production protected against Tau toxicity in vitro (Supplemental Figure 3), TH staining revealed a loss of LC neurons in Tau P301S/DBH+/– mice at 9 months that was partially abrogated by DBH–/– (Figure 5, A and B). To explore whether manipulation of NE synthesis affects cognitive impairment in Tau P301S mice, we tested spatial learning and memory using the Morris water maze (MWM). Comparable age-dependent deficits were observed at 3 and 6 months in NE-deficient and NE-competent Tau P301S mice. However, at 9 months, the performance of Tau P301S/DBH–/– mice was significantly better than Tau P301S/DBH+/– mice on several measures, including distance to platform during training and percentage of time spent in the target quadrant.
during a probe trial (Figure 5, C–E). Similar results were observed with cued and contextual fear conditioning, which test hippocampal-independent and -dependent associative learning and memory, respectively (Figure 5, F and G). AT8 and ThioS staining in the hippocampus also demonstrated that Tau pathology was increased age-dependently in Tau P301S/DBH+/– mice compared with Tau P301S/DBH−/– mice (Supplemental Figure 5D). Thus, the full expression of Tau pathology and cognitive impairment in Tau P301S mice depends on NE.

We next examined the impact of NE depletion on Tau pathology in the LC triggered by Tau cleavage. We reasoned that if DOPEGAL promotes the activation of AEP, then DBH−/− mice that cannot produce DOPEGAL would be protected from Tau pathology. We injected AAV encoding human Tau P301S or AEP-resistant Tau P301S N255A/N368A into the LC of DBH+/− and DBH−/− mice (Supplemental Figure 5D). Thus, the full expression of Tau pathology and cognitive impairment in Tau P301S mice depends on NE.

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**DOPEGAL is toxic to LC neurons in vivo.** MAO-A and DOPEGAL are upregulated in the LC of postmortem AD brains (24) and DOPEGAL (but not NE or its other oxidative or O-methylated metabolites) is toxic to differentiated catecholaminergic PC12 cells (23). To explore whether DOPEGAL is toxic to LC neurons in vivo and the potential involvement of AEP and Tau, we infused DOPEGAL (0.25 μg) into the LC of 2-month-old WT, Tau knockout (Tau−/−), MAPT transgenic mice that overexpress WT human Tau, and MAPT/AEP−/− mice. Four days following DOPEGAL administration, IF costaining with TH and TUNEL revealed diminished TH and greater apoptosis in MAPT mice compared with WT, and
these effects were abrogated by Tau or AEP depletion (Figure 6, A–C). These results indicate that DOPEGAL-induced degeneration of LC neurons in vivo is partially mediated by a Tau- and AEP-dependent mechanism.

To investigate whether acceleration of endogenous NE metabolism and DOPEGAL production would have similar effects, we injected AAV-MAO-A or control virus into the LC of WT and MAPT mice and assessed pathology 3 months later. MAO-A overexpression failed to trigger neuronal loss in WT mice but significantly enhanced apoptosis and loss of TH-positive cells in MAPT mice (Figure 7, A–C), which was accompanied by AT8 and Tau N368 staining in LC neurons (Figure 7E). We examined the levels of catecholamines and their metabolites in the LC by HPLC and verified that MAO-A overexpression significantly increased DOPEGAL and tended to decrease NE, as expected, without affecting any other analyte (Figure 7D). The staining of AT8 and ThioS in the hippocampus also demonstrated that Tau pathology was increased by MAO-A overexpression in MAPT mice compared with WT mice (Supplemental Figure 5E). MWM and fear conditioning tests demonstrated that MAO-A triggered significant cognitive dysfunction in MAPT mice compared with control virus, whereas it had no effect on WT mice (Figure 7, F–J). These data suggest that Tau exacerbates DOPEGAL-mediated LC neuron toxicity and cognitive impairment.

LC-derived human Tau can spread to interconnected brain regions in MAPT and 3xTg mice. Because the LC is the first brain structure to develop Tau lesions and has widespread connections to other areas of the brain, and Tau is capable of trans-synaptic propagation, LC neurons have been proposed to be the critical initiators of the stereotypical spread of Tau pathology in AD (4, 5). To explore this idea, we injected 3-month-old MAPT mice with either AAV-mCherry alone or AAV-mCherry + AAV-Tau under control of the noradrenergic-specific PRSx8 promoter into the LC (34), and then monitored Tau pathology throughout the brain 3 months later (Figure 8A). The staining of AT8 and ThioS in the hippocampus also demonstrated that Tau pathology was increased by MAO-A overexpression in MAPT mice compared with WT mice (Supplemental Figure 5E). MWM and fear conditioning tests demonstrated that MAO-A triggered significant cognitive dysfunction in MAPT mice compared with control virus, whereas it had no effect on WT mice (Figure 7, F–J). These data suggest that Tau exacerbates DOPEGAL-mediated LC neuron toxicity and cognitive impairment.
To determine whether AEP is necessary for Tau aggregation and spread in vivo, we injected 3-month-old WT and AEP–/– mice with AAV-PRSx8-Tau + AAV-PRSx8-mCherry in the LC and monitored Tau pathology in the LC and forebrain 3 months later. AT8, N368 Tau, and Gallyas-Braak staining indicated that hyperphosphorylated and AEP-cleaved Tau accumulation in the LC and subsequent spread to the HC, Cx, and EC were largely retarded in AEP–/– mice (Figure 10, A and B, and Figure 11A). A schematic map of Tau pathology spread from the LC in MAPT, 3xTg, WT, and AEP–/– mice as viewed by AT8 staining is shown in Supplemental Figure 7B.

Moreover, deletion of AEP prevented cognitive impairment induced by Tau overexpression in the LC (Figure 11, B–F). These data indicate that AEP contributes to the spread of LC-derived Tau pathology to the forebrain and associated cognitive deficits.

Discussion

Pathogenic forms of Tau first appear in noradrenergic LC neurons (3–6), which degenerate later in AD, and LC dysfunction contributes to AD-like neuropathology and cognitive deficits (6, 8, 9, 11–13, 16). However, the mechanism underlying the selective vulnerability of LC neurons has remained elusive. In the current study, we report that DOPEGAL, the NE metabolite produced by MAO-A oxidation, triggers AEP activation via oxidative stress, resulting in Tau N368 cleavage that is prone to hyperphosphorylation, aggregation, and propagation, and triggers noradrenergic neuronal death. In addition, DOPEGAL, but not NE, facilitates the fibrilization of purified recombinant Tau. Tau hyperphosphorylation and N368 cleavage were age-dependently increased in the LC of both MAPT and 3xTg mice, consistent with our previous finding that AEP is upregulated and activated in the brain during aging. Importantly, we made the same observations in the LC neurons of human AD brains, indicating that active AEP contributes to Tau pathology in the LC.

DOPEGAL is an appealing candidate to explain the selective vulnerability of LC neurons in AD for several reasons. First, it is only produced in noradrenergic cells. Second, it is increased in the LC of human AD brains and is toxic to noradrenergic neurons. AT8 IHC revealed that aberrant Tau had spread from the LC to the cerebellum, midbrain, hippocampus, entorhinal Cx, and Cx (Figure 8A). Moreover, IF and Gallyas-Braak staining indicated that hyperphosphorylated Tau, AEP-cleaved Tau, and aggregated Tau were present in the entorhinal Cx (EC), hippocampus (HC), and Cx of AAV-Tau injected mice (Figure 8C and Figure 9A). By contrast, neither aberrant Tau nor mCherry were detected in the forebrain of mice injected with AAV-mCherry alone (Figure 8C), indicating that Tau pathology, rather than the virus itself, spread from the LC to these distal regions. Similar results were obtained when we performed the identical experiment in 3xTg mice (Supplemental Figure 6A–C).

To test whether pathogenic Tau originating in the LC can induce cognitive deficits, we conducted MWM and fear conditioning 3 months after intra-LC Tau virus administration in MAPT and 3xTg mice. Tau-injected mice spent significantly less time in the target quadrant during the MWM probe trial (Figure 9, B–D) and were impaired in both cued and contextual freezing following fear conditioning (Figure 9, E and F). Together, these behavioral tests indicate that LC-derived Tau pathology can spread to the forebrain and produce cognitive impairment.

AEP cleavage of Tau increases its aggregation and neurotoxicity, and depletion of AEP diminishes Tau pathology in Tau P301S mice (18). Tau cleavage by AEP is necessary and sufficient for its neurotoxicity in LC neurons (Supplemental Figure 2), and the appearance of AEP-cleaved Tau N368 is tightly associated with AT8 immunoreactivity, suggesting that AEP cleavage of Tau may facilitate its spread. We prepared primary LC neurons from neonatal WT and AEP–/– mice and infected them with AAV-PRSx8-Tau or control virus. IF costaining with TH and Tau5 or the human-specific Tau HT7 antibody showed that human Tau was specifically overexpressed in the LC neurons (Supplemental Figure 7A, upper panels). TH and N368 or AT8 costaining demonstrated that human Tau was truncated at N368 and hyperphosphorylated in WT LC neurons, whereas both N368 and AT8 immunoreactivity were attenuated in AEP–/– LC neurons (Supplemental Figure 7A, lower panels).
Figure 7. Increasing DOPEGAL in the LC by MAO-A overexpression induces Tau pathology, cell death, and cognitive impairment in MAPT transgenic mice. AAV-MAO-A was injected into LC regions of WT or MAPT mice. (A) Representative images of TH (red) and TUNEL (green) staining in the LC 3 months following AAV-MAO-A injection. Scale bar: 100 μm. (B) Quantification of TUNEL. (C) Quantification of TH intensity. Data are shown as the mean ± SEM. n = 6 per group. *P < 0.05, by 2-way ANOVA. (D) Measurement of DOPEGAL in the LC of AAV-MAO-A–injected WT mice via HPLC analysis. Data were analyzed using 2-way ANOVA and are shown as the mean ± SEM (n = 3 per group). **P < 0.01, by 2-way ANOVA. (E) Representative images of staining for TH (green), AT-8 or Tau N368 (red), and DAPI (blue). Right panels are representative images of Gallyas-Braak staining. Scale bars: 20 μm and 100 μm, respectively. (F–J) Distance traveled to the platform (F), AUC for total distance traveled (G), and percentage of time spent in the quadrant (H) previously containing the platform during the probe trial in the MWM, and percentage of time spent freezing during the cued fear (I) and contextual fear (J) tests following fear conditioning. All data were analyzed using 1-way ANOVA and are shown as the mean ± SEM. n = 8 per group. *P < 0.055, by 1-way ANOVA.
Figure 8. Viral-mediated Tau expression in the LC drives propagation of pathology to the forebrain in MAPT transgenic mice. LC-specific AAV-PRSx8-Tau + AAV-PRSx8-mCherry, or AAV-PRSx8-mCherry alone were injected into the LC of MAPT mice, and mice were assessed for Tau pathology throughout the brain by immunohistochemistry 3 months later. (A) Representative images of AT8 immunostaining throughout the brain of MAPT mice. (B) Representative images of immunofluorescent staining for TH (blue), AT8 or Tau N368 (green), and mCherry (red) in the LC of MAPT mice. Scale bar: 100 μm (top panels) and 20 μm (bottom panels). (C) Representative images of immunofluorescent staining for AT8 (green) or Tau N368 (red), and DAPI (blue) in the EC, HC, and frontal Cx of MAPT mice. Scale bar: 20 μm.
Finally, the close neurochemical relationship between NE and DA suggests that the toxicity of the DA MAO metabolite DOPAL may have a correlate in the noradrenergic system. Previous studies show that physiological concentrations of DOPAL trigger the formation of α-Syn oligomers and aggregates in both a cell-free system and in cell culture, and produce cytotoxicity in vitro and in vivo. We recently reported that DOPAL activates AEP in dopaminergic neurons and leads to α-synuclein N103 cleavage by AEP, resulting in its aggregation and dopaminergic neuronal degeneration in the SN (21). Thus, DOPAL-α-Syn interactions may underlie the selective vulnerability of DA neurons in PD (35–39), and we suspected that a similar interaction might occur between DOPEGAL and Tau in AD. Here, we present multiple converging lines of evidence implicating DOPEGAL and AEP in the vulnerability of LC neurons to Tau pathology and toxicity. First, DOPEGAL, but not NE itself, increased AEP activity and cell death. Second, overexpression of MAO-A, which oxidizes NE into DOPEGAL, facilitated AEP activity, Tau N368 cleavage, and cell death, whereas decreasing MAO activity or blocking Tau cleavage had the opposite effect. Third, preventing DOPEGAL production via DBH depletion or Tau cleavage by AEP−/− mitigated the deleterious effects of Tau on LC neurons, whereas preventing the breakdown of DOPEGAL by AR or ALDH inhibition escalated AEP activity and cell death. Finally, full DOPEGAL toxicity was dependent on the presence of Tau. Together, these data suggest that DOPEGAL stimulates AEP activation via oxidative stress, resulting in Tau N368 cleavage and aggregation, LC neuronal degeneration, and the spread of Tau pathology to the forebrain and associated cognitive impairment (Figure 12). Given that DOPAL can directly modify α-Syn, it is possible that DOPEGAL may covalently modify the lysine residues and oxidize Met groups on Tau, thus facilitating its conformational change and fibrillization. We did find that DOPEGAL accelerated the aggregation of recombinant Tau in vitro, but further experiments will be required to pinpoint the covalent and/or noncovalent mechanisms.

MAO-A generates H2O2 and DOPEGAL, when it metabolizes NE. Both H2O2 and DOPEGAL induce AEP activation and trigger cell death, and we have shown that the transcription factor, CEBP-β, regulates the expression of AEP (40). Conceivably, there is positive feedback between NE metabolism and CEBP-β expression and activity, with increased levels of MAO-A and H2O2 activating CEBP-β, which, in turn, induces further MAO-A and AEP expression. MAO-A preferentially oxidizes serotonin and NE, whereas MAO-B preferentially oxidizes phenylethylamine, and both forms can oxidize DA (41). MAO-A oxidizes NE into DOPEGAL within less than 24 hours in SH-SY5Y cells (Supplemental Figure 2D). However, MAO-A overexpression by AAV infection is an indirect experiment to activate DOPEGAL in vivo, whereas DOPEGAL treatment is a direct approach to investigate DOPEGAL’s acute toxicity to LC neurons. AAV takes several weeks to reach steady-state gene expression, and the increased oxidation of NE to DOPEGAL is presumably gradual, because other enzymes are present in the LC that also metabolize DOPEGAL as well (i.e., aldehyde dehydrogenase and aldehyde reductase). The acute infusion of DOPEGAL likely overwhelms the normal machinery, leading to rapid toxicity, whereas the gradual production and buildup of DOPEGAL via AAV-MAO-A is likely to take more time. MAO-A is predominantly found in catecholaminergic neurons, with the highest concentration in the LC (42). Moreover, DOPEGAL and MAO-A levels are elevated 2.8- and 3.6-fold in AD LC neuronal cell bodies compared with controls (24), whereas MAO-B expression and activity are higher in multiple brain regions and cell types in AD (43, 44). The preference of MAO-A for NE and the enriched LC expression suggest that MAO-A is the main driver of DOPEGAL production in vivo, although both MAO-A and MAO-B were capable of increasing AEP activity, Tau N368 cleavage, and cell death in our experiments. We have previously demonstrated that MAO-B plays a greater role in...
generating the neurotoxic DA metabolite DOPAL in PD (21, 45), and here we show that NE metabolism into DOPEGAL by MAO-A in the progression of AD-like pathology. Combined, these results support our model that DA metabolism into DOPAL and MAO-A-mediated conversion of NE into DOPEGAL induce α-synuclein toxicity in SN and Tau toxicity in LC, respectively, leading to the selective vulnerability of catecholamine neurons in PD and AD (Figure 12).

Because DOPEGAL, but not NE itself, generates free radicals and activates mitochondrial permeability transition (27), DOPEGAL may contribute to the loss of noradrenergic LC neurons in AD (46). Employing Tau+/− neurons and mice, we showed that DOPEGAL-induced neurotoxicity is partially dependent on Tau. Knockout of AEP significantly ameliorated the LC neuronal degeneration and Tau pathology elicited by DOPEGAL exposure in MAPT mice, indicating that AEP cleavage of Tau is an important factor in LC vulnerability. This finding is further supported by our experiments with the AEP cleavage-resistant Tau P301S/N255/368A mutant. In SH-SY5Y cells, we found that the ability of AEP-cleaved Tau N368 to trigger cell death was more robust than WT Tau and comparable with P301S Tau. By contrast, introducing the N255/368A mutations onto a P301S background substantially reduced the toxicity of P301S Tau. Moreover, P301S Tau strongly induced Tau pathology in LC neurons in vivo that was associated with Tau N368 cleavage, while AEP-resistant P301S/N255/368A Tau was much less active. P301S-induced Tau pathology was also attenuated in the LC of DBH−/− mice, indicating that NE is required for Tau to exert its full neurotoxicity. This interpretation is further supported by our findings that siRNA-mediated knockdown of DBH (and thus NE) production blunted Tau-induced cell loss in catecholaminergic SH-SY5Y cells. Combined, these data support our model that DA metabolism into DOPAL and MAO-A–mediated conversion of NE into DOPEGAL induce α-synuclein toxicity in SN and Tau toxicity in LC, respectively, leading to the selective vulnerability of catecholamine neurons in PD and AD (Figure 12).

Previous research has shown that Tau pathology is detectable in 3xTg mice at 9 to 12 months (47) and in MAPT mice at 12 months (48), so in our experiments, any appearance of aberrant Tau at 6 months must be derived from the LC. We observed the appearance of hyperphosphorylated Tau in vivo that was associated with Tau N368 cleavage, while AEP-resistant P301S/N255/368A Tau was much less active. P301S-induced Tau pathology was also attenuated in the LC of DBH−/− mice, indicating that NE is required for Tau to exert its full neurotoxicity. This interpretation is further supported by our findings that siRNA-mediated knockdown of DBH (and thus NE) production blunted Tau-induced cell loss in catecholaminergic SH-SY5Y cells. Combined, these data support our model that DA metabolism into DOPAL and MAO-A–mediated conversion of NE into DOPEGAL induce α-synuclein toxicity in SN and Tau toxicity in LC, respectively, leading to the selective vulnerability of catecholamine neurons in PD and AD (Figure 12).
mCherry did not spread to distal brain regions, and there was no forebrain Tau pathology detected in mice that received mCherry alone. These results clearly show that, at least in these models, Tau pathology originating in the LC is capable of propagating to interconnected brain regions, including those that also show early vulnerability in AD. To determine whether this spread of pathogenic Tau had any functional effects, we tested mice for signs of cognitive impairment and found that the performances of 3xTg and MAPT mice in the MWM and fear conditioning assays worsened following Tau expression in the LC and subsequent spread to the forebrain. Remarkably, both the spread of Tau pathology and the accelerated cognitive deficits following Tau expression in the LC were alleviated in AEP–/– mice. Here, we provide a combination of in vitro, cellular, in vivo, genetic, and pharmacological evidence to describe a specific neurochemical/molecular pathway involving NE metabolism and AEP that may explain the selective vulnerability of LC neurons to developing Tau pathology, the later demise of these cells, and the potential spread of pathology from the LC to the forebrain. Identification of the DOPEGAL-AEP-Tau N368 pathway lays the foundation for development of LC/AEP-based therapies that could slow the progression of AD.

Methods

Animals

Tau P301S mice on a C57BL/6J background (line PS19), MAPT mice, 3xTg mice, and WT C57BL/6J mice were obtained from The Jackson Laboratory (stock nos. 008169, 005491, 34830, and 000664, respectively). The AEP–/– mice were maintained on a mixed 129/Ola and C57BL/6 background. DBH–/– mice were maintained on a mixed 129/SvEv and C57BL/6 background. Animal care and handling was performed according to the Declaration of Helsinki and Emory Medical School guidelines. Tau P301S mice were crossed with DBH–/– mice to generate Tau P301S/DBH–/– mice. Adrenergic agonists and/or L-DOPS were used to rescue the DBH–/– embryonic lethality and male mating deficit, as described (49). MAPT mice were crossed with AEP–/– mice to generate MAPT/AEP–/– mice. Investigators were blinded to the group allocation during the animal experiments.

Human tissue samples

Paraffin sections containing the LC from 5 postmortem AD cases of Braak stages IV–VI (age 74.5 ± 11.2 years, mean ± SD) and 5 cognitively normal controls (age 73.9 ± 12.7 years) were obtained from the Emory Alzheimer’s Disease Research Center. AD was diagnosed according to the criteria of the Consortium to Establish a Registry for AD and the National Institute on Aging. Diagnoses were confirmed by the presence of amyloid plaques and NFTs in formalin-fixed tissue.

Transfection and viral infection of cells

Primary cortical and noradrenergic neurons were cultured as previously described (50). All rats were purchased from The Jackson Laboratory. HEK293 cells and SH-SY5Y cells were transfected with plasmids encoding Tau, SiDBH, MAO-A, or MAO-B using polyethyleneimine or Lipofectamine 3000 (Thermo Fisher Scientific). AAV-Tau, AAV-Tau...
HPLC analysis

LC tissue was resuspended in 150 μL ice-cold 0.1 M perchloric acid (PCA) containing 0.1 mM EDTA and sonicated on ice by probe sonication on setting 3 with a 30% duty cycle. Homogenates were centrifuged at 10,000 × g for 10 minutes at 4°C, and supernatants were transferred into 0.22-μM PVDF microcentrifuge filter tubes to remove any remaining particulate matters at 5000 × rpm for 5 minutes at 4°C. Monoamine concentrations were determined by reverse phase HPLC with electrochemical detection. Tissue protein pellets were dissolved in 500-μL 2% SDS. Protein quantification was carried out in triplicate in 96-well microplates with SpectraMax M5e spectrophotometer (Molecular Devices) using the BCA method (Pierce BCA Protein Assay Kit, Thermofisher Scientific). For HPLC, an autosampler (ESA) 5600A CoulArray detection system, equipped with an ESA Model 584 pump and an ESA 542 refrigerated autosampler was used. Separations were performed at 23°C using an Xterra RP 4.6 × 250 mm, 3.5-μm particle size, C18 column (Waters). The mobile phase consisted of 72 mM sodium hydrogen phosphate, 28 mM citric acid, 0.6 mM 1-octanesulfonic acid sodium, and 0.2 mM EDTA at pH 5.35. A 20-μL sample was injected. The samples were eluted isocratically at 0.5 ml/min and detected using a 6210 electrochemical cell. Analytical cell potentials were –175, 100, and 350 mV. The analytes were identified by the matching criteria of retention time to known standards. Compounds were quantified by comparing peak areas with those of standards on the dominant sensor.

ThioT assay

ThioT stock solution (3 mM) was filtered through a 0.2-μm syringe filter (Sigma-Aldrich, catalog T3516). The stock solution was diluted into the aggregation buffer (20 mM Tris, pH 7.4, 100 mM NaCl, 1 mM EDTA) to generate the working solution (30 μM). ThioT working solution with heparin (30 μM), recombinant Tau (7 μM), and DOPEGAL (500 μM) was incubated on an orbital shaker at 37°C and excited at 440 nm and emitted at 482 nm to measure the fluorescence intensity on the plater reader (BioTek, 251639). Fluorescence values (excitation wavelength 440 nm and emission wavelength 482 nm) were plotted against the concentration of the target protein. The IC50 value was determined by fitting the data to a sigmoidal dose–response curve using Prism software (GraphPad).

Western blot analysis

Cultured cells were lysed in lysis buffer (50 mM Tris, pH 7.4, 40 mM NaCl, 1 mM EDTA, 0.5% Triton X-100, 1.5 mM Na3VO4, 50 mM NaF, 10 mM sodium pyrophosphate, 1.5 mM Na3VO4, 0.5% Triton X-100, 10 mM sodium β-glycerophosphate, supplemented with a cocktail of protease inhibitors) and sonicated for 15 minutes at 16,000 g. The supernatant was boiled in SDS loading buffer. After SDS-PAGE, the samples were transferred to a nitrocellulose membrane. Primary antibodies to the following targets were used: Tau5 (Santa Cruz Biotechnology, SC-8436), AT-8 (Thermo Fisher Scientific, MN1020), HT7 (Thermo Fisher Scientific, MN1000), β-actin (Sigma-Aldrich, A5316), MAO-B (GeneTex, GTX113771), MAO-A (Sigma-Aldrich, SC-58860), AT-8 (Thermo Fisher Scientific, MN1020), HT7 (Thermo Fisher Scientific, MN1000), β-actin (Sigma-Aldrich, A5316), MAO-B (GeneTex, GTX113771), MAO-A (Sigma-Aldrich, SC-58860), TH (Abcam, AB112), AEP (Cell Signaling Technology, 93627S), Iba1 (VWR, 019-19741), and Tau N368 (custom antibody, 18).

AEP activity assay

Assay buffer (20 mM citric acid, 60 mM NaHPO4, 1 mM EDTA, 0.1% 3-(3-cholamidopropyl)dimethylammonio]-1-propanesulfonate [CHAPS], and 1 mM DTT, pH 6.0) of 100 μL including 100 μM AEP substrate Z-Ala-Ala-Asn-AMC (ABchem) was added to cell or tissue lysates. AMC released by substrate cleavage was quantified using a fluorescence plate reader at 460 nm for 1 hour in kinetic mode.

MAO-A activity assay

Cell or tissue lysates (20 μg) were incubated with the working solution of 100 μL containing 400 μM AmplexÒ Red reagent, 2 U/mL HRP, and 2-mM p-tyramine substrate with MAO-B inhibitor, pargyline (Molecular Probes). The fluorescence of MAO-A activity was measured in a fluorescence plate reader using excitation in the range of 530–560 nm and emission at 590 ± 10 nm at 37°C for 2 hours in kinetic mode.

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tation at 440 nm and emission 482 nm) onto the plate reader (BioTek, 251639) were recorded for 5 days.

**Immunostaining**

Paraffin-embedded human brain sections or free-floating mouse brain sections sliced by cryotome were treated with 0.3% H2O2 for 10 minutes. Sections were washed 3 times in PBS and blocked in 1% BSA, 0.3% Triton X-100 for 30 minutes, followed by overnight incubation with TH antibody (1:1000), anti-Tau5, anti-Tau N368, or AT-8 antibody (1:500) at 4°C. The signal was developed using a Histostain-SP kit (Invitrogen, Thermo Fisher Scientific). For IF, sections were incubated overnight with various primary antibodies at 4°C. Then, the sections were incubated with the matched fluorocoujugated secondary antibody for 2 hours at room temperature, followed by 3 washes in PBS. The slides were washed 3 times in PBS and covered with a glass using mounting solution, after DAPI staining for 5 minutes.

**Stereotaxic injection**

AAV-PRSx8-mCherry, AAV-PRSx8-Tau, AAV-Tau P301S, AAV-Tau N255/368A, or AAV-MAO-A were injected into the LC region of mice. Three-month-old mice of each group were anesthetized with isofluorane (Piramal Healthcare). Meloxicam (2 mg/kg) was injected s.c. for analgesia (Loxicom, Norbrook). Unilateral intracerebral injection of virus was performed stereotaxically at the following coordinates: anteroposterior -5.4 mm and mediolateral -1.2 mm relative to bregma, and dorsoventral -3.7 mm from the dural surface. Two microliters of viral suspension was injected into each site using a 10-μl Hamilton syringe with a fixed needle at a rate of 0.25 μL/min. The needle remained in place for 5 minutes after the viral suspension was completely injected and then was slowly removed over 2 minutes. The mice were placed on a heating pad until recovery from the anesthesia.

**Behavioral testing**

MWM. Mice were trained in a round, water-filled tub (52-in. diameter) in an environment with extra maze cues. Each subject was given 4 trials/day for 5 consecutive days, with a 15-minute intertrial interval. The maximum trial length was 60 seconds, and if subjects did not reach the platform in the allotted time frame, they were manually guided to it. Following the 5 days of task acquisition, a probe trial was presented, during which time the platform was removed and the percentage of time spent in the quadrant that previously contained the escape platform during task acquisition was measured over 60 seconds. All trials were analyzed for latency and swim speed by MazeScan (CleverSys Inc.). We monitored body weight, visual ability, and swim speed during behavioral test to verify that the deficits did not result from motor dysfunction.

**Fear conditioning**

The ability to form and retain an association between an aversive experience and environmental cues was tested with a standard fear-conditioning paradigm occurring over a period of 3 days. Mice were placed in the fear-conditioning apparatus (7” W, 7” D × 12” H, Coulbourn Instruments) composed of Plexiglas with a metal shock grid floor and allowed to explore the enclosure for 3 minutes. Following this habituation period, 3 conditioned stimulus (CS)-unconditioned stimulus (US) pairings were presented with a 1-minute intertrial interval. The CS consisted of a 20-second 85-db tone, and the US consisted of 2 seconds of a 0.5-mA footshock, which was coterminal with each CS presentation. One minute following the last CS-US presentation, mice were returned to their home cage. On day 2, the mice were presented with a context test, during which subjects were placed in the same chamber used during conditioning on day 1, and the amount of freezing was recorded via camera (software provided by Colbourn). No shocks were given during the context test. On day 3, a tone test was presented, during which time subjects were exposed to the CS in a novel compartment. Initially, animals were allowed to explore the novel context for 2 minutes. Then the 85-db tone was presented for 6 minutes, and the amount of freezing behavior was recorded.

**Quantification and statistical analysis**

All cell culture data are expressed as mean ± SEM from 3 or more independent experiments, and the level of significance between 2 groups was assessed with Student’s t test. For experiments consisting of more than 2 groups, 1-way or 2-way ANOVA followed by Tukey’s post hoc test was applied. A P value of less than 0.05 was considered statistically significant. Sample sizes were determined by Power and Precision (Biostat).

**Study approval**

The protocol for animal experiments was reviewed and approved by the Emory Institutional Animal Care and Use Committee. The study using human tissue samples was approved by the Biospecimen Committee, and informed consent was obtained from the subjects prior to their deaths.

**Author contributions**

KY conceived the project, designed the experiments, analyzed the data, and wrote the manuscript. SSK designed and performed most of the experiments and analyzed the data. XJ, prepared primary neurons and assessed animal experiments. CJ performed the crosses, pharmacological embryonic lethality rescue, and genotyping for experiments with the DBH-/- mice. EHA and JX performed Tau purification and some of the in vitro experiments. FPM prepared the uncleavable Tau virus. HRL and XY assisted with data analysis and interpretation and critically read the manuscript. DW helped design the experiments and edit the manuscript.

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