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SUMMARY
It has been suggested that aberrant activation of glycogen synthase kinase-3-beta (GSK-3β) can trigger abnormal tau hyperphosphorylation and aggregation, which ultimately leads to neuronal/synaptic damage and impaired cognition in Alzheimer disease (AD). We examined if isoform-selective partial reduction of GSK-3β can decrease pathological tau changes, including hyperphosphorylation, aggregation, and spreading, in mice with localized human wild-type tau (hTau) expression in the brain. We used adeno-associated viruses (AAVs) to express hTau locally in the entorhinal cortex of wild-type and GSK-3β hemi-knockout (GSK-3β-HK) mice. GSK-3β-HK mice had significantly less accumulation of hyperphosphorylated tau in synapses and showed a significant decrease of tau protein spread between neurons. In primary neuronal cultures from GSK-3β-HK mice, the aggregation of exogenous FTD-mutant tau was also significantly reduced. These results show that a partial decrease of GSK-3β significantly represses tau-initiated neurodegenerative changes in the brain, and therefore is a promising therapeutic target for AD and other tauopathies.

INTRODUCTION
Glycogen synthase kinase 3 (GSK-3) is an ubiquitously expressed, highly conserved serine/threonine kinase that regulates multiple major biological processes such as cell differentiation, metabolism, immunity, and cell survival (Maurer et al., 2014). GSK-3 is expressed as two highly homologous isoforms, α and β, that have partially overlapping functions. GSK-3β is the most abundant isoform in neurons (Leroy and Brion, 1999; Takahashi et al., 1994; Woodgett, 1990) and is involved in the pathogenesis of multiple neurological disorders including schizophrenia, bipolar disorder, fragile X syndrome, brain tumors, stroke, Parkinson disease, and Alzheimer disease (AD) (Chuang et al., 2011; Jope and Roh, 2006; Kozikowski et al., 2006; Mills et al., 2011; Mines and Jope, 2011; Takashima, 2006).

One of the hallmark pathological changes in the AD brain is the hyperphosphorylation and intraneuronal aggregation of the microtubule-associated protein tau in form of neurofibrillary tangles (NFTs) (reviewed in Gomez-Isla et al., 2008). Tau NFT pathology spreads through the brain with disease progression (Braak and Braak, 1991) and correlates with neuronal death and cognitive decline (Gomez-Isla et al., 1997). Furthermore, soluble phosphorylated forms of tau appear to be synaptotoxic (Tai et al., 2012) and can propagate between neurons across neuronal networks (de Calignon et al., 2012; Dujardin et al., 2014; Wegmann et al., 2019).

In AD, GSK-3β is a pivotal kinase responsible for tau phosphorylation, and multiple GSK-3β phosphate- sited tau sites are present in NFTs (Hanger et al., 1992; Lucas et al., 2001; Mandelkow et al., 1992). It has been demonstrated that GSK-3β is aberrantly activated in human AD brains (DaRocha-Souto et al., 2012; Leroy et al., 2007) and is a key player in the pathogenesis of AD (Takashima, 2006). Overexpression of GSK-3β leads to tau hyperphosphorylation, microtubule dissociation, and cognitive impairment (Engel, 2006;
Isoform-selective reduction of GSK-3β in GSK-3β-HK mice

In AD, hyperphosphorylation of tau by GSK-3β and other kinases is suggested to promote its aggregation into insoluble aggregates (Figure 1A) (Hanger et al., 1992) and is a strong correlate of synapse loss and neurodegeneration (Gomez-Isla et al., 1996; Gong and Iqbal, 2008). GSK-3β phosphorylates more than 30 sites in the human tau protein and is considered one of the main tau kinases (Figure 1B). Using immunohistochemistry (IHC), we confirmed in mice as well as in primary neurons that the constitutively active form of GSK-3β (pGSK-3β-Tyr216) co-localizes with tau in the neuronal cytosol and in some neurites (Figure 1C).

To study the effect of GSK-3β reduction on tau, we used GSK-3β-HK mice (Hoeflich et al., 2000). First, we confirmed that in these mice total GSK-3β levels were reduced by 45% (p < 0.0001), and the levels of pGSK-3β-Tyr216 (enzyme active form) and pGSK-3β-Ser9 (enzyme inhibited form) are reduced by about 40% compared with WT mice (Tyr216: p = 0.0004 and Ser9: p = 0.0045; Figures 2A and 2B). Importantly, no alteration in the protein levels of other relevant tau kinases—GSK-3α, pGSK-3α-Tyr279 (active form of GSK-3α), FYN, and CDK5—were detected (GSK-3α: p = 0.1035; pGSK-3α-Tyr279: p = 0.7100; FYN: p = 0.3213; CDK5: p = 0.9874; Figures 2A and 2B). These data confirmed that the reduction of GSK-3 levels in the brain of GSK-3β-HK mice is selective for the β-isoform of the enzyme, does not lead to compensatory upregulation of remaining GSK-3β activity, and does not induce significant upregulation of other kinases implicated in the pathologic phosphorylation of tau protein in AD.

We also tested if WT and GSK-3β-HK mice had comparable total tau levels at baseline (non-injected: p = 0.9905; Figures 3K and 3N) as well as after AAV injections (AAV-injected: p = 0.9052; Figures 2C, 2D, 3K, and 3N) to ensure that genetic manipulation of GSK-3β did not affect tau protein expression.

AAV-mediated expression of hTau in the EC of WT and GSK-3β-HK mice

To generate a model that allows us to study the effect of GSK-3β reduction on tau accumulation and spreading, we utilized targeted intracranial AAV injections to express hTau in the EC of WT and GSK-3β-HK mice. The AAV construct used was designed to express eGFP and hTau as individual proteins separated by a self-cleaving 2a-peptide under the CBA promoter (AAV CBA-eGFP-2a-hTau; Figure 3A) (Wegmann et al., 2015, 2017, 2019). AAV-transduced “donor neurons” express both eGFP and hTau, whereas “recipient neurons,” which acquired hTau protein from transduced donor neurons, contain only hTau (no eGFP) that is detectable by immunostaining (Figure 3B). AAV-injections into the superficial layers (II–III) of the EC (Figures 3C and 3D) successfully led to the expression of hTau around the injection site.
Twelve weeks after AAV injection, brain sections of WT and GSK-3β-HK mice were immunolabeled for hTau (HT7 or TauY9 antibody) and GFP to identify and distinguish tau-transduced donor from tau recipient neurons. Most neurons around the injection site were AAV-transduced donor neurons expressing both GFP (green) and hTau (red) (Figure 3E). Brain sections (and primary neurons) were immunolabeled for GFP and a neuronal marker (NeuN), to confirm neuronal expression of the AAV (Figure 3F).

Comparable AAV transduction and hTau expression in GSK-3β-HK and WT mice

Unbiased stereological counts of neurons (Nissl stained) in the medial and lateral EC showed no differences in the total number of neurons between GSK-3β-HK versus WT non-injected or injected mice (p = 0.9682 and p = 0.6685, respectively; Figures 3G and 3H). This excluded both the possibility of a baseline difference in neuronal numbers between the two genotypes, as well as a neurotoxic effect of the AAV.
injections themselves, which could have artificially impacted the rate of trans-cellular tau spreading in the brain.

Differences in AAV transduction and hTau expression could potentially result in differences in detected tau spreading. To rule out such artifacts, we compared the volume of the injection sites (area with GFP-positive neurons in consecutive brain sections; p = 0.6702, Figure 3I) and the total number of transfected neurons (number of GFP-positive neurons; p = 0.6414, Figure 3J) between WT and GSK-3b-HK mice. We found comparable AAV transduction and hTau expression in both groups of mice. This was further confirmed by western blot analysis of EC brain lysates, in which the amount of total tau (mouse and human tau (ms + h);
Figure 3. AAV-mediated human tau expression in the entorhinal cortex of GSK-3β-HK and WT mice
(A) AAV construct designed to express eGFP and hTau as individual proteins, separated by the self-cleaving 2a peptide under the CBA promoter (AAV CBA.eGFP-2a-hTau).

(B) Schematic representation of proteins expected to be identified in the EC of mice injected with AAV. AAV-transduced neurons express eGFP (green) and hTau (red) (“donor neurons”), whereas some neurons received hTau protein from other cells and do not have GFP (“recipient neurons” (red only)).

(C) Stereotactic coordinates for unilateral intracranial AAV injection into the left EC of adult WT and GSK-3β-HK mice.

(D) Representative image of a Nissl-stained horizontal section of a mouse brain with the approximate area corresponding to the EC shaded in green.

(E) Horizontal brain sections of a WT and a GSK-3β-HK-injected mouse with AAV expression in the EC and interconnected neighboring regions. Twelve weeks after AAV injection, brain sections were immunostained for hTau (antibodies HT7 or TauY9) and GFP.

(F) Primary neurons and brain sections immunostained for GFP and neuronal marker NeuN show that AAV is expressed in neurons (“AAV-transduced neurons”; white arrowheads).
Figure 3. Continued

(G) Stereologically based counts of neurons (Nissl stained) in the non-injected (right hemisphere) and (H) injected (left hemisphere) EC of WT and GSK-3β-HK mice showed no significant differences, indicating that there are no baseline differences in neuronal numbers between the two genotypes or neuronal cell death due to the AAV injections. Data are presented as mean ± SEM, N = 14 mice: 7WT, 7HK, 5 brain sections per mouse. Two-tailed Student’s t test, p > 0.05.
(I) The volume of the transduced brain area (GFP+) and (J) the number of transduced (GFP+) neurons was similar in GSK-3β-HK and WT mice. Data are presented as mean ± SEM, N = 14 mice: 7WT, 7HK, 3–5 sections per mouse, two-tailed Student’s t test, p > 0.05.

(K) Representative image of western blot from EC lysates from non-injected and AAV-injected WT and GSK-3β-HK mice probed for postsynaptic (PSD95) and presynaptic (synaptophysin, SYP) markers and Total Tau (ms + h). GAPDH was used as loading control.

(L) Quantification of western blots for Postsynaptic density protein 95 (PSD95), Synaptophysin (SYP), and (N) Total Tau (ms + h). Levels of synaptic markers (PSD95 and SYP) and total tau were not significantly different between WT and GSK-3β-HK mice at baseline and 12 weeks after AAV injection. Data are presented as mean ± SEM, N = 6 non-injected mice: 3WT, 3HK, two-tailed Student’s t test; N = 14 AAV-injected mice: 7WT, 7HK, two-tailed Student’s t test, p > 0.05. Scale bars: 1,000 µm (E), 20 µm (F, in vitro and in vivo).

Indicator for general neuronal abundance and hTau expression was comparable between the WT and GSK-3β-HK groups of mice (Figures 2C and 2D). Furthermore, the overall levels of post- and pre-synaptic markers, PSD95 and synaptophysin, respectively, were similar in WT and GSK-3β-HK mice both at baseline (PSD95: p = 0.7960; synaptophysin: p = 0.6304) and after AAV injections (PSD95: p = 0.9242; synaptophysin: p = 0.7402; Figures 3L and 3M), indicating no overt gain or loss of synapses upon GSK-3β reduction or as a result of AAV injections and hTau expression.

GSK-3β reduction diminishes tau spreading in vitro and in vivo

Next, we assessed if the reduction in GSK-3β changes the potential of tau to propagate between neurons in the brain. hTau propagation was quantified both in vitro in primary neuronal cultures from WT and GSK-3β-HK embryos and in vivo in adult WT and GSK-3β-HK mice unilaterally injected with AAV-CBA-eGFP-2a-hTau into the EC. We counted all neurons (in culture and in the EC) that were simultaneously positive for GFP and hTau (= transduced tau donor neurons) and neurons that were positive only for hTau (= tau recipient neurons). We observed in vitro that primary neurons derived from GSK-3β-HK embryos showed significant lower number of hTau recipient neurons compared with WT (0.68% ± 0.19% vs. 3.19% ± 0.72%, respectively; p = 0.0356; Figure 4A). Furthermore, and as expected from previous in vivo studies (Wegmann et al., 2015, 2019), transduced donor neurons near the injection site expressed both GFP and hTau (Figure 4B: green and red neuron “b” in lower panel) and, after 12 weeks, a small number of hTau recipient neurons (expressing hTau only but not GFP) were also present both in WT and GSK-3β-HK mice (Figure 4B: red neuron “a” in lower panel). Even though tau propagation was a rare event in WT mice and less than 1% of all hTau-containing neurons in the EC were identified as “tau recipient” neurons (7.9 ± 2.1 of 829 hTau positive neurons (0.94%)), this percentage was found to be significantly reduced in GSK-3β-HK mice, with the number of tau recipient neurons being zero in some GSK-3β-HK mice (1.7 ± 0.8 of 826 hTau positive neurons (0.21%); p = 0.0063; Figure 4B). These results indicate that a partial decrease in GSK-3β levels can significantly diminish tau trans-cellular propagation in the brain.

GSK-3β reduction decreases synaptic accrual of p-Tau in vivo

Levels of total tau and p-Tau (as reported by PHF-1 antibody) in total cell extracts from primary neuronal cultures did not show significant differences between GSK-3β-HK and WT neurons (total tau: p = 0.2437 and PHF-1: p = 0.2757. Figures 5A and 5B). It has been suggested that tau phosphorylation site-specific antibodies against pS202 and pS396/p404 and against misfolded tau label progressive stages of disease-associated tau changes (Augustinack et al., 2002). In agreement with these observations, immunofluorescence labeling of GSK-3β-targeted tau phosphorylation sites in brain sections from WT and GSK-3β-HK mice showed that of all the neuronal cell bodies expressing hTau in the EC, approximately 30% were positive for pS202 (CP13 antibody), 18% were positive for pS396/p5404 (PHF-1 antibody), and 8% were positive for misfolded tau (Alz50 antibody), with no significant differences between WT and GSK-3β-HK AAV-injected mice (CP13: p = 0.3626; PHF-1: p = 0.8224; Alz50: p = 0.6490. Figures 5C and 5D). The spread of NFT pathology through the AD brain is thought to rely, at least in part, on synaptic transmission of pathological tau proteins between neurons. Because a reduction in GSK-3β levels diminished the trans-cellular transmission of tau in the EC of GSK-3β-HK mice compared with WT, we investigated the effect of the overall decrease in GSK-3β expression on the kinase levels, specifically within the synaptic compartment. We
Figure 4. Reduction of GSK-3β diminishes tau propagation in vitro and in vivo

(A) Representative images of WT and GSK-3β-HK primary neurons immunostained for GFP and hTau (Y13) after addition of AAV to culture media. Arrowhead shows a tau-recipient neuron (red) amid AAV-transduced neurons (red and green). The percentage of tau recipient neurons was significantly reduced in primary neurons from GSK-3β-HK embryos compared with WT. Data are presented as mean ± SEM, N = 7: 4WT, 3HK, 6 wells of plated neurons per embryo. Two-tailed Student’s t test, *p < 0.05.

(B) Immunofluorescence images of the injected EC of WT and GSK-3β-HK mice showed neurons that had hTau (red) but no GFP (green) (“tau recipient neurons”, neurons in insert “a”) and neurons that had both GFP (green) and hTau (red) (“AAV-transduced neurons”; neurons in insert “b”). The percentage of tau recipient neurons (normalized to GFP + neurons) was significantly reduced in the EC of GSK-3β-HK compared with WT mice. Data are presented as mean ± SEM, n = 15: 7WT, 7HK, two-tailed Student’s t test, **p < 0.01. Scale bars: 20 μm (A) and 10 μm (B).
Figure 5. Reduction of GSK-3β leads to lower levels of p-Tau tau in synapses
(A) Representative western blot images using homogenates from WT and GSK-3β-HK primary neurons after treatment with hTau (AAV added to culture media). Immunoblots were probed with antibodies for GSK-3β, total tau, and PHF-1 (pS396, pS404). GAPDH was used as loading control.
(B) Protein quantification showed that despite significantly lower levels of GSK-3β in total cell homogenates, GSK-3β-HK neurons displayed similar overall levels of total tau and p-Tau (PHF-1) compared with WT neurons. Data are presented as mean ± SEM, n = 10: 5WT, 5HK. Two-tailed Student’s t test, **p < 0.001.

(C) Horizontal brain section showing an illustrative outline of the EC where AAV-transduced neurons can be easily identified by the presence of GFP labeling.
Brain sections of WT and GSK-3β-HK-injected mice were co-labeled for GFP and specific tau-phosphorylation sites targeted by GSK-3β: CP13: pS202, PHF-1: pS396/pS404, and Alz50: misfolded tau.

(D) Quantification of GFP + neurons that were also positive for CP13, PHF-1, and Alz50. About 27% of the transduced neurons were positive for CP13, 15% were positive for PHF-1, and only 6% were positive for Alz50. There were no significant differences between the number of neurons positive for any of the three markers between GSK-3β-HK and WT mice. Data are presented as mean ± SEM, n = 14: 7WT, 7HK, 3 sections per mouse. Two-tailed Student’s t test, p > 0.05.
measured the levels of GSK-3β, pGSK-3β-Tyr216 (active form of the enzyme) and -Ser9 (inhibited form), total tau, and p-Tau in both the cytosol (CYT) and in synaptoneurosome (SNS) preparations from the EC of WT and GSK-3β-HK mice after AAV CBA-eGFP-2a-hTau injection (Figure 5E).

Interestingly, GSK-3β-HK mice, compared with WT mice, not only had significantly lower levels of GSK-3β in both compartments (CYT and SNS) (CYT: p < 0.05 and SNS: p < 0.0001) but also exhibited a different distribution of GSK-3β between CYT and SNS. WT mice had higher levels of GSK-3β in SNS compared with CYT (p < 0.05), whereas there was no difference in GSK-3β levels between the two compartments in GSK-3β-HK mice (p > 0.05; Figure 5F). This suggests that neurons, in response to genetic reduction of total GSK-3β levels, may favor a decrease of synaptic over cytosolic GSK-3β, perhaps to preserve minimal required levels of this kinase in the cytosol for critical cellular functions. Total protein levels of pGSK-3β (-Tyr216 and -Ser9) were also significantly lower in GSK-3β-HK compared with WT mice (Figure 2B). Notably, the levels of active GSK-3β (-Tyr216) were significantly lower in both compartments (CYT and SNS) in GSK-3β-HK compared with WT mice (CYT and SNS: p < 0.0001; Figure 5F), but were not associated with a significant change in the compartmental distribution between the two groups (p > 0.05). Interestingly, the levels of active GSK-3β (-Tyr216) were significantly lower in the cytosol but not in synapses of GSK-3β-HK compared with WT mice (CYT: p < 0.05 and SNS: p > 0.05 for SNS; Figure 5F). These results further support the idea of a preferential reduction of total and active GSK-3β in synapses of GSK-3β-HK mice.

Overall levels of total tau were not significantly different between WT and GSK-3β-HK mice (Figures 2C and 2D), yet both genotypes had significantly more tau in the SNS fraction (42% and 36% higher levels in synapses, respectively; p < 0.01; Figure 5F). Importantly, p-Tau (PHF-1 antibody-detecting epitopes s396/p404 in mouse and human tau) levels were overall significantly lower in GSK-3β-HK compared with WT mice (p = 0.0233), whereby p-Tau levels were only slightly lower in CYT but significantly lower in SNS of GSK-3β-HK compared with WT mice (CYT: p < 0.05 and SNS: p < 0.05; Figure 5F).

Together, these results show that even though WT and GSK-3β-HK mice had similar amounts of total tau in the cytosolic and synaptic compartments, the 45% reduction of GSK-3β in the GSK-3β-HK mouse resulted in the selective decrease of GSK-3β—followed by reduction of tau phosphorylation—within the synaptic compartment. Notably, this seems to be a specific effect of the isoform-selective reduction of GSK-3β because no compensatory up- or downregulation of other relevant tau kinases (GSK-3α, pGSK-3α-Tyr216 (active form), CDK5, and FYN) could be detected in the brain of GSK-3β-HK mice (Figures 2A and 2B).

Partial GSK-3β reduction decreases tau aggregation in vitro

GSK-3β has been found to be associated with tangles in AD brains (Imahori and Uchida, 1997; Pei et al., 1997, 1999; Yamaguchi et al., 1996) as well as accumulated in pre-tangle neurons (Pei et al., 1999).

In our model, in vivo expression of hTau upon AAV CBA-eGFP-2a-hTau injections did not trigger the formation of brain β-sheet-containing tau aggregates, as reported by Thioflavin S, either in WT or in GSK-3β-HK mice, even after long-term expression (6 months post-injection, Figure 6A). This is in agreement with previous findings that tau aggregation into β-sheet-containing aggregates is not a prerequisite for propagation (Dujardin et al., 2014; Wegmann et al., 2019).

We therefore tested if a reduction of GSK-3β could potentially decrease tau aggregation using a more aggressive in vitro model based on the expression of aggregation-prone FTD-mutant tau in primary
Figure 6. Reduced levels of GSK-3β decrease tau aggregation in vitro

(A) Representative images of brain sections from WT and GSK-3β-HK mice 6 months after AAV-injection, stained with Thioflavin S (marker of β-sheet-rich amyloid-like aggregated proteins) and immunolabeled for NeuN (neuronal marker). No tau aggregates were observed.

(B) Schematic representation of tau aggregation assay in neurons: primary neurons from WT and GSK-3β-HK embryos were co-transduced with lentiviruses encoding tauRD(P301L)-CFP and tauRD(P301L)-YFP, and tau aggregation is initiated by subsequent treatment of the neurons with seeding-competent tau in brain extracts from tau transgenic mice (rTg4510). Tau aggregation in the neurons can be detected by fluorescent imaging as FRET signal.

(C) Timeline for tau aggregation assay: primary neurons from WT and GSK-3β-HK embryos are plated in 96-well plate (DIV0) and transduced with lentivirus (lenti-tauRD(P301L)-YFP and -CFP) (DIV1). Transduction is followed with treatment with rTg4510 brain lysate (DIV6) and imaging of FRET-positive tau aggregates (DIV8).

(D) Representative images and quantification of tau aggregates in primary neuronal cultures at DIV8. GSK-3β-HK neurons showed significantly fewer intracellular tau aggregates compared with WT neurons. Inserts show tau aggregates formed intracellularly in WT and HK neurons in culture. Data are presented as mean ± SEM, N = 22 embryos: 10WT, 12HK, 3–5 wells plated neurons per embryo. Two-tailed Student’s t test, ***p < 0.0001. Scale bars: 100 μm (A) and 20 μm (D).
neuronal cultures derived from WT and GSK-3β-HK embryos. In this in vitro model, primary neurons were transduced with lentiviral viruses (Lenti CBA.tauRDP301L-YFP/CFP) encoding the repeat domain of tau (tauRD) carrying a P301L FTD mutation fused to either CFP or YFP fluorescent protein, in order to monitor tauRD aggregation through fluorescence resonance energy transfer (FRET; Figure 6B) (Holmes et al., 2014; Nobuhara et al., 2017). Aggregation of tauRD was induced by the addition of brain lysates from tauP301L transgenic mice (tg4510 line) containing pre-aggregated, seeding-competent tau (Figures 6B and 6C). Six days after aggregation seeding (at DIV8), we found that the number of intracellular tau aggregates was significantly reduced in GSK-3β-HK compared with WT neurons (p < 0.001; Figure 6D). Importantly, viral particle number (number of neurons transduced) and neuronal cell density were comparable in all experiments and treatment groups. These results show that the reduction of GSK-3β not only significantly lowers synaptic tau phosphorylation and propagation in the brain but it also significantly inhibits tau aggregation in neurons in culture.

**DISCUSSION**

In the present work we show that AD-related tau pathological changes (hyperphosphorylation, aggregation, and propagation) can be diminished by the reduction of GSK-3β. Expressing unmodified human wild-type tau in WT and GSK-3β-HK mice using AAVs, we found that partial isoform-selective genetic decrease of GSK-3β could reduce synaptic tau hyperphosphorylation and propagation in the mouse brain. Furthermore, seeded tau aggregation was diminished in primary neurons from GSK-3β-HK embryos expressing aggregation prone FTD-mutant tau. Here, we show that the genetic decrease of GSK-3β reduces not only tau phosphorylation and aggregation but also neuronal propagation. Tau propagation across neural circuits is increasingly recognized to contribute to pathology in AD and other tauopathies (Gibbons et al., 2019; Mudher et al., 2017; Takeda, 2019a, 2019b; Walker et al., 2013). We conclude that a partial reduction of GSK-3β is sufficient to reduce pathological tau changes in the brain and therefore could be utilized to halt the tau-associated neurodegeneration.

We and others have previously shown aberrant activation of GSK-3β in human AD brains and AD transgenic mice (Blalock et al., 2009; DaRocha-Souto et al., 2012; Leroy et al., 2007; Serenó et al., 2009). Because there is a general consensus that pharmacological modulation of abnormal tau hyperphosphorylation is a promising therapeutic approach for AD (Engel, 2006; Engel et al., 2006; Le Corre et al., 2006; Nakashima et al., 2005; Noble et al., 2005; Pérez et al., 2003), the inhibition of GSK-3β seems an obvious target to achieve the reduction of pathological tau hyperphosphorylation (Cohen and Goedert, 2004). Pharmacological reduction of GSK-3 has been used by several groups, including our own, to reduce AD-like phenotypic changes in transgenic rodent and in vitro models (de Barreda et al., 2010; DaRocha-Souto et al., 2012; Engel, 2006; Houck et al., 2016; Muñoz-Montaño et al., 1997; Serenó et al., 2009; Wilson et al., 2020). However, the GKS-3 inhibitors used in those studies (e.g. thiadiazolidine derivatives and lithium, among others) are not selective for the GSK-3β isoform, and thus inhibition of other kinases could have potentially contributed to the observed results. The few thiadiazolidine derivative and lithium trials conducted in AD patients have yielded mixed results on potential benefits/side effects (Lovestone et al., 2015; Matsunaga et al., 2015; Mazanetz and Fischer, 2007; Muñoz-Montaño et al., 1997; Wilson et al., 2020). Thus, the development of a specific pharmacological inhibitor of GSK-3β is still an awaited feat (Gong and Iqbal, 2008). Efforts to develop selective, safe, tissue-specific, and dose-efficient GSK-3β inhibitors have been unsuccessful to date (Bhat et al., 2018; Matsunaga et al., 2019). However, the recent development of novel drug delivery strategies aimed at achieving adequate therapeutic concentrations to target organ or cells (i.e. nanoparticles or GSK-3β-cell knockouts), whereas avoiding potential off-target effects provides a new window of opportunity (reviewed in Bhat et al., 2018).

In the current study, we overcame the limitations of non-selective inhibition by using a mouse model with an isoform-selective reduction of GSK-3β activity through haploinsufficiency (GSK-3β-HK line).

Interestingly, GSK-3β-HK mice did not only show reduced tau pathological changes but also revealed new insights about the cellular distribution of GSK-3β in neurons. We found that GSK-3β, commonly referred to as mostly cytosolic (Bijur and Jope, 2001, 2003; Watcharasit et al., 2002), is present at even higher levels in synapses, pointing to its (yet insufficiently acknowledged) role in this cellular compartment. Even though a role of GSK-3β as a regulator of synaptic plasticity has been suggested (Bradley et al., 2012), particularly in long-term depression via NMDA receptors (Peineau et al., 2007, 2008, 2009), the physiological function of this enzyme in synapses remains largely unknown. GSK-3β-HK mice had a stronger reduction of GSK-3β...
levels in synapses compared with the cytosol, potentially explaining the partial protection against tau pathological changes in these mice. In fact, the function of GSK-3β may depend, at least in part, on its subcellular localization. For example, whereas nuclear and mitochondrial GSK-3β were reported to participate in pro-apoptotic signaling, cytosolic GSK-3β seems to mediate survival signals (Jacobs et al., 2012). In combination with our data, we postulate that synaptic GSK-3β levels may be reduced in GSK-3β-HK neurons in order to ensure essential amounts of GSK-3β in the cytosol.

In healthy neurons, tau is mostly restricted to axons where it binds to microtubules and promotes axonal transport (Avila et al., 2004; Buee et al., 2000; Papasozomenos and Binder, 1987). However, synaptic accumulation of p-Tau in AD (Brandt et al., 2005; Gendron and Petrucelli, 2009; Perez-Nievas et al., 2013; Tai et al., 2012) can lead to altered synaptic function (Hoover et al., 2010), correlates with preclinical stages of the disease, and seems to precede neuronal degeneration (Anvanitakis et al., 2007). Thus, the reduction of soluble hyperphosphorylated tau accumulation in synapses could protect against the tau-initiated neurodegenerative cascade in AD. Synaptic p-Tau could result either from local tau phosphorylation by synaptic GSK-3β activity or from cytosolic GSK-3β-mediated tau phosphorylation inducing p-Tau transport to the synapse.

In this study we were able to show that both soluble p-Tau accrual in synapses and tau protein spread between neurons in the brain can be successfully achieved through partial reduction of GSK3β. These findings support the idea that the presence of p-Tau in synapses may be directly linked to its ability to propagate trans-cellularly. We propose that GSK-3β activation lies upstream of neurodegenerative tau phenotypes in AD, not only by promoting tau phosphorylation and aggregation (Engel, 2006; reviewed in Mines et al., 2011) but also by promoting abnormal accrual of p-Tau into the synaptic compartment and trans-cellular spread in the brain, ultimately compromising the function and anatomical integrity of neurons and synapses in AD (see Figure 7 for proposed model).

Previous studies based on selective in vivo silencing of either GSK-3α or GSK-3β in transgenic mice suggested that the GSK-3α isoform contributes to both amyloid and tau pathology, whereas the GSK-3β isoform only modulates tau phosphorylation and neurofibrillary tangle formation (Hurtado et al., 2012). Our results show that GSK-3β-HK neurons (in vitro and in vivo) still render overall levels of tau phosphorylation that are similar to WT neurons and yet exhibit a targeted reduction of p-Tau levels in synapses that may confer protection against tau pathological changes.

After 12 weeks of human wild-type tau expression in vivo, we did not observe β-sheet-containing tau aggregates (Thioflavin-S staining) in WT or GSK-3β-HK mice, even in the presence of robust tau hyperphosphorylation (PHF1+) and misfolding (Alz50+). This observation, in agreement with our previous work and

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**Figure 7. Schematic of GSK-3β and p-Tau in the synapses**

GSK-3β and tau co-localize in the soma and neurites of neurons. Genetic reduction of GSK-3β in mice leads to overall lower levels of GSK-3β in all cellular compartments and also changes the relative abundance of the kinase in the cytosolic and synaptic compartments. In WT mice, GSK-3β accumulates in higher levels in the synapse compared with the cytosol, whereas in GSK-3β-HK mice both compartments have similar amounts of GSK-3β. GSK-3β-HK mice have the same amount of total tau in the synapses as WT mice but significantly lower amounts of active GSK-3β and p-Tau.
others (Dujardin et al., 2014; Mudher et al., 2017; Wegmann et al., 2019), provides further evidence that not necessarily aggregated but also soluble tau species can travel from a “donor” to a “recipient” cell. Our current results support the idea that a decrease in soluble synaptic tau phosphorylated at Ser396/404 may be linked to reduced tau spreading observed in GSK-3β-HK mice and primary neurons. Whether GSK-3β-mediated tau phosphorylation at this epitope directly promotes tau release and uptake (leading to tau spreading) remains to be further elucidated.

In conclusion, the current study shows that a partial isoform-selective decrease of GSK-3β is sufficient to diminish pathological phosphorylation and propagation of human tau in the mouse brain and is also effective against the aggregation of FTD-mutant tau in cultured neurons. We propose a mechanism, in which the reduction of GSK-3β in synapses lowers synaptic p-Tau, leading to (1) the reduction of tau trans-cellular propagation and (2) a decrease in seeded tau aggregate formation. Targeted manipulation of GSK-3β levels and/or its activity in the synapse may thus be a promising approach against tau-related pathological changes in the brain that are relevant in AD and other tauopathies.

Limitations of the study
We used AAV-mediated local expression of wild-type human tau expression to detect tau protein propagation and the effects of local tau overexpression. Our model has limitations for studying tau aggregation in vivo: AAV-injected WT and GSK-3β-HK mice did not develop β-sheet-containing tau aggregates upon expression of hTau in the EC, even at 6 months post-injection. To study the effect of GKS-3β reduction on tau aggregation, we therefore employed a previously established approach to measure tau aggregation propensity in cells through expression of the aggregation-prone repeat domain of FTLD-mutant tauP301S (Nobuhara et al., 2017). Of note, this tau construct does not occur in the etiology of AD and may be mechanistically different from wild-type tau aggregation in AD. However, this in vitro assay allowed us to demonstrate that the selective reduction of GKS-3β was protective against tau-induced pathological formation of neuronal aggregates.

Resource availability
Lead contact
Requests for additional information can be directed to the Lead Contact, Ana Claudia Amaral (aamaral@mgh.harvard.edu).

Materials availability
This study did not generate unique reagents.

Data and code availability
This study did not generate code.

METHODS
All methods can be found in the accompanying Transparent methods supplemental file.

SUPPLEMENTAL INFORMATION
Supplemental information can be found online at https://doi.org/10.1016/j.isci.2021.102058.

ACKNOWLEDGMENTS
We thank Dr. James R. Woodgett for the donation of the GSK-3β-HK mouse line and Dr. Peter Davies for the donation of the p-Tau antibodies (CP13, PHF-1, and Alz50) used in this study.

AUTHOR CONTRIBUTIONS
Conceptualization: A.C.A., B.P.N., S.W., and T.G.I.; Design of the Experiments: A.C.A and T.G.I.; Performance of the Majority of the Experiments: A.C.A. and M.S.T.C.; Methodology: A.C.A, E.H., Z.F., S.T., S.W., and T.G.I.; Additional Experimental Work: A.G.M., H.A.E., S.R.G., C.C., S.M., B.E., and P.R.; Construct Design and Production: E.H., Z.F., and S.W.; Funding Acquisition: T.G.I; Supervision: S.W. and T.G.I.
DECLARATION OF INTERESTS

Teresa Gómez-Isla participated as speaker in an Eli Lilly and Company-sponsored educational symposium and serves in an Eli Lilly Data Monitoring Committee. All other authors declare no competing interests.

Received: July 13, 2020
Revised: November 20, 2020
Accepted: January 7, 2021
Published: February 19, 2021

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Supplemental Information

Isoform-selective decrease of glycogen synthase kinase-3-beta (GSK-3β) reduces synaptic tau phosphorylation, transcellular spreading, and aggregation

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# Supplemental Information

## Table S1. Key Resources Table

| REAGENT or RESOURCE | SOURCE | IDENTIFIER |
|---------------------|--------|------------|
| **Antibodies**      |        |            |
| Rabbit anti-human tau (Y9) | Enzo Life Sciences | Cat# BML-TA3119-0025, RRID:AB_205263 |
| Mouse anti-Tau 13 | Covance | Cat# MMS-520R-500, RRID:AB_291452 |
| Mouse anti-Tau (HT7) | Thermo Fischer Scientific | Cat# MN1000, RRID:AB_2314654 |
| Rabbit anti-Tau (mouse and human) | Agilent | Cat# A0024, RRID:AB_10013724 |
| Mouse anti-CP13 (p-Tau S202, T205) | Peter Davies | RRID:N/A |
| Mouse anti-PHF-1 (p-Tau S396, S404) | Peter Davies | RRID:N/A |
| Mouse anti-Alz50 (misfolded tau) | Peter Davies | RRID:N/A |
| Rabbit anti-PSD95 (post synaptic density) | Cell Signaling Technology | Cat# 3409, RRID:AB_1264242 |
| Mouse anti-PSD95 (post synaptic density) | Thermo Fischer Scientific | Cat# MA1-045, RRID:AB_325399 |
| Mouse anti-Synaptophysin (pre-synaptic vesicles) | Millipore | Cat# MAB329, RRID:AB_94786 |
| Chicken anti-GAPDH | Millipore | AB2302, RRID:AB_10615768 |
| Mouse anti-GAPDH | Thermo Fischer Scientific | Cat# AM4300, RRID:AB_2536381 |
| Rabbit anti-GAPDH | Cell Signaling Technology | Cat# 2118, RRID:AB_561053 |
| Rabbit anti-GSK-3β | Cell Signaling Technology | Cat# 9315, RRID:AB_490890 |
| Mouse anti-GSK-3β | Cell Signaling Technology | Cat# 9832, RRID:AB_10839406 |
| Rabbit anti-GSK-3α | Cell Signaling Technology | Cat# 9338, RRID:AB_2114897 |
| Mouse anti-p GSK-3β/α-Tyr216/Tyr279 | Millipore | Cat# 05-413, RRID:AB_309721 |
| Rabbit anti-p GSK-3β/α-Tyr216/Tyr279 | Bioss | Cat# bs-2073R, RRID:AB_10857314 |
| Rabbit anti-pGSK-3β/α-Ser9 | Cell Signaling Technology | Cat# 5558, RRID:AB_10013750 |
| Chicken anti-GFP (green fluorescent protein) | Aves | Cat# GFP-1020, RRID:AB_10000240 |
| Mouse anti-NeuN (neuronal marker) | Millipore | Cat# MAB377, RRID:AB_2298772 |
| Rabbit anti-Fyn | Cell Signaling Technology | Cat# 4023, RRID:AB_10698604 |
| Mouse anti-Cdk5 | Millipore | Cat# 05-364, RRID:AB_2229170 |
| LICOR goat anti-rabbit 680 (IRDye680RD) | LICOR Biosciences | Cat# 926-68071, RRID:AB_10956166 |
| LICOR donkey anti-mouse 680 (IRDye680RD) | LICOR Biosciences | Cat# 926-68072, RRID:AB_10953628 |
| LICOR goat anti-rabbit 800 (IRDye800CW) | LICOR Biosciences | Cat# 926-32211, RRID:AB_62184 |
| Label | Type | Cat# | RRID | Notes |
|-------|------|------|------|-------|
| LICOR goat anti-mouse 800 (IRDye800CW) | LICOR Biosciences | 926-32210 | AB_621842 |
| LICOR donkey anti-chicken 680 (IRDye680RD) | LICOR Biosciences | 926-68075 | AB_10974977 |
| Alexa Fluor 488 Goat anti-chicken | Thermo Fischer Scientific | A-11039 | AB_253409 |
| Cy3 goat anti-mouse | Abcam | ab97035 | AB_10680176 |
| Cy3 goat anti-rabbit | Abcam | ab6939 | AB_955021 |
| Bacterial and Virus Strains | | | | |
| AAV CBA-eGFP-2a-hTau | Wegmann et al., 2019 | | N/A |
| Lenti CBA.tauRDP301L-YFP/CFP | Nobuhara et al., 2017 | | N/A |
| Chemicals, Peptides, and Recombinant Proteins | | | | |
| Odyssey Blocking Buffer (PBS) | LICOR Biosciences | 927-40000 | N/A |
| DAPI Fluoromount-G | SouthernBiotech | 0100-20 | N/A |
| Experimental Models: Cell Lines | | | | |
| Primary mouse cortical neurons | Laboratory of Dr. Teresa Gomez-Isla | | N/A |
| Experimental Models: Organisms/Strains | | | | |
| Mouse: GSK-3β-HK | MGH mouse breeding facility | | N/A |
| Software | | | | |
| Axiovision Imaging Software | Carl Zeiss | | SCR_002677 |
| ZEN Digital Imaging for LSM 800 | Carl Zeiss | | SCR_013672 |
| NIS-Elements for Nikon A1 Confocal Microscope | Nikon Instruments | | SCR_014329 |
| LICOR Image Studio Software | LICOR Biosciences | | SCR_015795 |
| CAST Stereology Software | Olympus | N/A | N/A |
| ImageJ | National Institutes of Health | | SCR_003070 |
| Zotero | https://www.zotero.org | | SCR_013784 |
| GraphPad Prism 6 | GraphPad Prism | | SCR_002798 |
| Other | | | | |
| Zeiss Axiocam 100 Inverted Microscope | Carl Zeiss | | N/A |
| Zeiss LSM 800 Microscope | Carl Zeiss | | N/A |
| Nikon A1 Confocal Microscope | Nikon Instruments | | N/A |
| BX51 Microscope | Olympus | | N/A |
| Stereotactic Frame | David Kopf Instruments | | N/A |
Transparent Methods

Ethics statement
The generation, care, and use of animals as well as all experimental procedures were approved by the Institutional Animal Care and Use Committee (IACUC) of the Massachusetts General Hospital (MGH). The animals’ living conditions, including housing, feeding, and nonmedical care, were maintained by the house internal animal facility (Center for Comparative Medicine) at MGH. Mice of both genders were included in the experiments. Wild-type and GSK-3β-HK mice tissues were prepared and used for biochemical and immunostaining assays at MGH.

Animals
GSK-3β-HK mouse line described in Hoeflich et al. (2000) was kindly donated by Dr. James R. Woodgett. Mice were bred in-house and the mutation of GSK-3β was confirmed by tail DNA genotyping following previously published protocols (Hoeflich et al., 2000). Groups of 8-10-month-old gender-matched WT and GSK-3β-HK mice (N= 14 per genotype, 16 males and 12 females) underwent intracranial injections as described below. Brains from 8-10-month-old gender-matched non-injected WT and GSK-3β-HK mice (N= 5 per genotype, 5 males and 5 females) were also collected for baseline biochemical and immunostaining assays.

AAV production and intracranial injections
Production and intracranial injection of AAV were performed as described elsewhere (Wegmann et al., 2017, 2019). In brief, for efficient targeting of neurons in the CNS, the AAV vector used in this study was serotype AAV 8. The target protein expression is driven by the ubiquitous CBA promoter. The target gene encodes the fluorescent protein eGFP (enhanced green fluorescent protein as a transduction marker) and wild-type human tau separated by a translation interrupting 2a peptide (AAV8-CBA-eGFP-2a-4R-hTau). Vector was injected into the left entorhinal cortex (EC) of WT and GSK-3β-HK mice (volume: 1.5μl per animal and titer: ~6X10^{12} infectious particles/ml). Vector was produced at Mass Eye & Ear (MEEI) Vector Core, Boston. Animals were anesthetized with isoflurane (~2%) and positioned on a stereotactic frame (David Kopf Instruments) with the following coordinates calculated from bregma: -4.5 mm anteroposterior, mediolateral: -4.5 mm, and dorsoventral: -1.7 mm from the brain surface. Mice were euthanized twelve weeks after AAV injection.

Primary neuron cultures
Primary neuronal cultures were derived from freshly dissected cortices from WT (N=3) and GSK-3β-HK (N=4) E14-16 embryos. Neurons were seeded at a density of 0.6x10^{5} cells per well on culture dishes pretreated coated with Poly-D-lysine. Fortau propagation studies, neurons were transduced by direct addition of AAV (eGFP-2a-4R-hTau) particles to the culture medium at 7 days in vitro (7 DIV) as previously described (Wegmann et al., 2019). At 14 DIV, primary neuron cultures were carefully washed with PBS (without agitation) and fixed with 4% PFA for 15 min. Neurons were then washed with PBS, permeabilized with 0.2% Triton X-100 in PBS for 15 min, blocked with 5% normal goat serum (NGS) in PBS (RT), and then incubated with the primary antibodies overnight at 4°C in 5% NGS/PBS. For detection of tau recipient neurons (neurons that acquired hTau protein from transduced donor neurons), primary antibodies against GFP and hTau (Tau13) were used (see Key Resources Table for antibodies description). Appropriate secondary antibodies (Life Technologies, 1:1000) were applied in 2% NGS/PBS for 1h at RT in the dark. After washing the cells in PBS twice for 15 min, they were stained with DAPI (Roche, #10236276001, 1:1000), and washed again in PBS. Neurons were imaged using an Axiovert inverted microscope (Zeiss). For tau phosphorylation analyzes, primary neurons from WT (N=5) and GSK-3β-HK (N=5) E14-16 embryos were treated as described above (tau propagation study) until 14 DIV (7 days after the addition of AAV (eGFP-2a-4R-hTau) particles to the culture medium at 7 DIV). At 14 DIV primary neurons were collected and lysed in cold PBS with protease and phosphatase inhibitor cocktail. The cells were transferred to a 2ml Eppendorf, homogenized, and incubated for 30 min on ice. Each sample was then centrifuged, and the supernatants were collected. Protein concentrations were determined using a Pierce BCA Protein Assay Kit. Whole cell lysates were electrophoresed on 4-12% Bis-Tris Novex mini gels (Invitrogen, #MAN0003679) in MES running buffer for SDS-polyacrylamide gel electrophoresis (Invitrogen). Gels were transferred onto nitrocellulose membranes (Thermo Fisher Scientific) and blocked for 1h in Blocking Buffer (Odyssey, Li-Cor, #927-40000). The membranes were probed with antibodies for anti-total tau, PHF-1, and
Lentiviral vectors encoding the mutant tau repeat domain (tauRD) constructs tauRD P301L-CFP and tauRD P301L-YFP were generated as previously described (Nobuhara et al., 2017) and added to neurons in culture. In short, primary neurons derived from E14-16 cortices of WT (N=10) and GSK-3β-HK (N=12) embryos were co-transduced with lentiviruses encoding tauRD P301L-CFP and tau RD P301L-YFP on DIV1 and treated with rTg4510 brain extract on DIV6. Prior to application, rTg4510 brain extracts were diluted with culture medium to a final concentration of 100 µg/mL total protein and filtered through a 0.2-µm membrane filter, and 50 µL were added to each well (5 µg total protein/well). Primary neurons from WT and GSK-3β-HK mouse embryos were cultured in eight wells, of which three were randomly selected for quantification. Five 10X images were taken from each well (approximately 25% of each well) in an Axiovert inverted microscope (Zeiss) and the number of intracellular tau aggregates were counted using ImageJ (NIH). Cells were stained for DAPI and NeuN and the number of nuclei and neurons counted to verify consistent plating and transfection across both conditions.

GSK3β (See Key Resources Table for complete information on antibodies). GAPDH was used as a loading control. Appropriate secondary antibodies (goat anti-rabbit-IRDye680, goat anti-rabbit-IRDye800, goat anti-mouse-IRDye680, goat anti-chicken-IRDye680, and goat anti-mouse-IRDye800 (Li-Cor)) were added at a 1:5000 dilution for 1h at RT and membranes were visualized using the Li-Cor Imaging system. Western blot images were then quantified using Image Studio Lite.

**Tau aggregation assay in primary neurons**

Lentiviral vectors encoding the mutant tau repeat domain (tauRD) constructs tauRD P301L-CFP and tauRD P301L-YFP were generated as previously described (Nobuhara et al., 2017) and added to neurons in culture. In short, primary neurons derived from E14-16 cortices of WT (N=10) and GSK-3β-HK (N=12) embryos were co-transduced with lentiviruses encoding tauRD P301L-CFP and tau RD P301L-YFP on DIV1 and treated with rTg4510 brain extract on DIV6. Prior to application, rTg4510 brain extracts were diluted with culture medium to a final concentration of 100 µg/mL total protein and filtered through a 0.2-µm membrane filter, and 50 µL were added to each well (5 µg total protein/well). Primary neurons from WT and GSK-3β-HK mouse embryos were cultured in eight wells, of which three were randomly selected for quantification. Five 10X images were taken from each well (approximately 25% of each well) in an Axiovert inverted microscope (Zeiss) and the number of intracellular tau aggregates were counted using ImageJ (NIH). Cells were stained for DAPI and NeuN and the number of nuclei and neurons counted to verify consistent plating and transfection across both conditions.

**Synaptoneurosomes preparation and analysis of tau content by Western blotting**

Mice were euthanized by CO2 asphyxiation and perfused through the heart with 20 ml of PBS. Brains were quickly removed, washed in PBS, and immediately frozen in liquid nitrogen and stored at -80°C until ready to be batch processed. Left EC from AAV-injected and from non-injected mice were processed to separate synaptoneurosomes from cytosolic fraction as previously described (Tai et al., 2012). Briefly, left EC was homogenized in Buffer A (25mM HEPES ph7.5, 120mM NaCl, 5mM KCl, 1mM MgCl2, 1mM CaCl2) with phosphatase inhibitor (Roche 04906845001) and protease inhibitor (Roche 11697498001). The homogenate was then passed through two 80um filters (Millipore Nylon Filter, NY8002500) and 70µL were separated for the total fraction. 70µL of distilled water and 23µL of 10% SDS were added to the separated homogenate and passed through a 27 1/2G needle 3 times to shear DNA. The remaining homogenate was passed through 5um filters (PALL Acrodisc, 4650) and centrifuged at 1000g for 10 min at 4°C. The supernatant was collected and placed in the ultra-centrifuge at 100,000g (38,000 rpm) for 45 min at 4°C. Meanwhile, the pellet was resuspended in 70µL of buffer B (50mM Tris HCL, 1.5% SDS, 1mM DTT). The pellet resuspension and the total homogenate fraction were then boiled for 5 min and centrifuged at 15min at 15,000g at 4°C. The supernatant of the pellet resuspension was collected as the synaptoneurosome fraction (SNS) and the supernatant of the homogenate fraction was collected as the total fraction. Finally, the supernatant was collected from the ultra-centrifuged samples as the cytosolic fraction (CYT). Fraction purity was confirmed by Western blot using a PSD95 specific antibody. Brain lysates from the different fractions (25 µg of protein) were electrophoresed on 4-12% Bis-Tris Novex mini gels (Invitrogen, #MAN0003679) in MES running buffer for SDS-polyacrylamide gel electrophoresis (Invitrogen). Gels were transferred onto nitrocellulose membranes (Thermo Fisher Scientific) and blocked for 1h in Blocking Buffer (Odyssey, Li-Cor, #927-40000). The membranes were probed with the following antibodies: anti-GFP, total tau, human tau, PHF-1, GSK3β, phospho-GSK-3β-Tyr216, phospho-GSK-3β-Ser9, GSK-3c, phospho-GSK-3α-Tyr279, CP13, Alz50, CDK5, Fyn, PSD95, and synaptophysin (See Key Resources Table for complete information on antibodies). GAPDH was used as a loading control. Appropriate secondary antibodies (goat anti-rabbit-IRDye680, goat anti-rabbit-IRDye800, goat anti-mouse-IRDye680, goat anti-chicken-IRDye680, and goat anti-mouse-IRDye800 (Li-Cor)) were added at a 1:5000 dilution for 1h at RT and membranes were visualized using the Li-Cor Imaging system. Western blot images were then quantified using Image Studio Lite.

**Analysis of tau content by immunohistochemistry**

Brain tissue to be used in immunohistochemistry was collected after mice were euthanized by CO2 asphyxiation and perfused with 20 ml of PBS and 20 ml of 4% PFA/PBS. Brains were removed and cryoprotected in a series of glycerol solutions (10% glycerol with 2% dimethylsulphide (DMSO) in PBS for 24 h and then transferred to a solution of 20% glycerol with 2% DMSO in PBS for 24h). After cryoprotection, brains were flash-frozen in 2-methylbutane at -75°C and stored at -80°C until sectioning (Rosene et al., 2012).
Sectioning was accomplished using a freezing microtome set to a thickness of 40 µm, and horizontal sections were collected in ten interrupted series of sections so that the interval between sections within a given series was 400 µm. Series of free-floating sections were stored in 30% glycerol/PBS at -80°C until ready to be batch processed. One series of sections was selected for thionin staining and stereological analysis, which were immediately mounted on gelatin-subbed slides and allowed to air dry. The sections were then rehydrated in descending graded ethanol solutions (100% X2, 95%, 70% 50%, and distilled water) for 25 seconds per bath, and stained with thionin for 2 minutes. Slides were then dehydrated in ascending ethanol solutions (distilled water, 50%, 70%, 95%, 100% X2; 1 min per bath), cleared in three xylene baths for 5 min each, and coverslipped with Permount (Fisher Scientific). The thionin series was used to quantify neuronal numbers and regional volumes, while the remaining series were used for immunohistochemistry (IHC). Sections for IHC were stored in a solution of 30% glycerol/PBS at -20°C until ready to be batch processed. For labeling of proteins of interest (such as hTau, total tau, p-Tau, GSK-3β, and others), sections were thawed and washed in PBS, permeabilized with 0.1% Triton-X, and blocked in 5% NGS/PBS. Primary antibodies were diluted in 5% NGS and were incubated overnight at 4°C (see Key Resources Table for a list of antibodies). The following day, sections were washed in PBS and incubated with secondary antibodies, washed in PBS, mounted in glass slides and coverslipped using mounting medium with DAPI (SouthernBiotech). Images were obtained using an Olympus BX51 microscope and a Nikon A1 confocal microscope. Images were processed in Nikon Elements and ImageJ (NIH).

Stereological neuronal cell counts
The number of hTau donor cells, hTau recipient cells, p-Tau (CP13- and PHF-1-positive), and misfolded tau (Alz50-positive) containing neurons in the EC was determined by counting all [GFP+], [hTau+/GFP+], [CP13+/GFP+], [PHF1+/GFP+], and [Alz50+/GFP+] neurons in the left EC using the CAST software (Olympus). The cell numbers were counted in three to four brain sections per mouse and in five to seven mice per group. For the total number of neurons in the left (injected) and right (non-injected) ECs, we outlined layers 1-6 of the medial and lateral entorhinal cortices and performed stereological counting of 20% of the neurons in the ROIs, followed by extrapolation to 100%. Counting was done on an Olympus BX51 light microscope equipped with a 20X objective.

Quantification and Statistical Analysis
Statistical analysis was performed using GraphPad Prism6. The applied statistical tests, post hoc analyses, and sample sizes (n) are defined in each figure legend. A p value < 0.05 was considered statistically significant. Sample size was chosen based on similar experiments in previously published studies (Wegmann et al., 2015, 2019). Unless otherwise noted, all data are presented as mean ± SEM.

Supplemental References
Rosene, D.L., Roy, N.J., and Davis, B.J. (1986). A cryoprotection method that facilitates cutting frozen sections of whole monkey brains for histological and histochemical processing without freezing artifact. J. Histochem. Cytochem. 34, 1301–1315.