Relevance of Phosphorylation and Truncation of Tau to the Etiopathogenesis of Alzheimer’s Disease

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Microtubule (MT) associated protein tau is abnormally hyperphosphorylated and aggregated into paired helical filaments (PHFs), which manifest as neurofibrillary tangles (NFTs) in the brains of individuals with Alzheimer’s disease (AD) and related tauopathies. Hyperphosphorylation and truncation of tau have been linked to the progression of the disease. However, the nature of phosphorylation and truncation of tau in AD brain are not very clear. In the present study we investigated the association of phosphorylation and truncation with high-molecular weight oligomers of tau (HMW-tau) in post-mortem AD brain by western blots. We found that tau from AD brain appears as a smear from low molecular weight (LMW) to HMW tau species in western blots developed with pan-tau antibodies. Similar level of LMW-tau was found in AD and control brains, whereas HMW-tau was found in AD brain only. HMW-tau was hyperphosphorylated at multiple sites and not unphosphorylated at Ser46 or Ser198/199/202. HMW-tau was weakly labeled by tau antibodies 43D against a.a. 6–18 and HT7 against a.a. 159–163 of tau, whereas, the C-terminal antibodies, tau46 and tau46.1, strongly labeled HMW-tau. The ratio of HMW-tau/LMW-tau detected by tau antibodies increased as the epitope of the tau antibodies ranges from N-terminal to C-terminal. The level of tau truncated at Asp421 was increased in AD brain, but was poorly associated with the HMW-tau. These findings suggest that tau pathogenesis involves both hyperphosphorylation and dominantly N-terminal truncation of tau in AD.

Keywords: Alzheimer’s disease, tau, hyperphosphorylation, truncation, tau pathogenesis

INTRODUCTION

Alzheimer’s disease (AD) is characterized by extracellular deposits of β-amyloid plaques (Glenner et al., 1984) and intracellular neurofibrillary tangles (NFTs) consisting of abnormally hyperphosphorylated aggregates of the microtubule (MT) associated protein tau (Grundke-Iqbal et al., 1986a,b). The number of NFTs in the brain correlates to the severity of dementia symptoms in AD patients (Alafuzoff et al., 1987; Arriagada et al., 1992; Riley et al., 2002). Tau is a cytosolic phosphoprotein, the major function of which is to stimulate and stabilize MT assembly from tubulin subunits. Normal phosphorylation of tau controls its function in the regulation of MT dynamic, which is involved in neuronal polarity, axonal growth and axonal transportation. In AD brain, tau is abnormally hyperphosphorylated (Księżak-Reding et al., 1992; Köpke et al., 1993), which leads
to the loss of biological activity, the gain of toxic activity, and the aggregation into paired helical filaments (PHFs; Iqbal et al., 1986; Alonso et al., 1994, 1996, 2001; Lucas et al., 2001; Fath et al., 2002; Jackson et al., 2002; Pérez et al., 2002).

In addition to phosphorylation, tau is post-translationally modified by ubiquitination (Mori et al., 1987; Perry et al., 1987), SUMOylation (Dorval and Fraser, 2006; Luo et al., 2014), glycation (Ledesma et al., 1994; Yan et al., 1994), acetylation (Min et al., 2010; Cohen et al., 2011), glycosylation (Wang et al., 1996b), O-GlcNAcylation (Arnold et al., 1996; Liu et al., 2004), nitration (Horiguchi et al., 2003), and truncation (Novak et al., 1993). Among them, truncation of tau has been shown to promote tau aggregation (Kovacech and Novak, 2010; Wang et al., 2010).

As a natively unfolded/poorly-folded protein, tau is very sensitive to protease digestion. Numerous studies have shown that tau is a substrate for calpain (Johnson et al., 1989), caspase (Gamblin et al., 2003), thrombin (Arai et al., 2005), cathepsin (Bednarski and Lynch, 1996), Puromycin-specific aminopeptidase (PSA; Karsten et al., 2006), and asparagine endopeptidase (Zhang et al., 2014) in vitro and in vivo. These studies have shown that tau truncation plays an important role in both tau aggregation and neurodegeneration (Kovacech and Novak, 2010). In AD brain, several specific truncations of tau have been identified (Wang et al., 2010). Truncation of tau at Asp421 (D421) and Glu391 (E391) has been shown to make tau prone to aggregation (Kovacech and Novak, 2010). However, the truncation of tau in the oligomeric forms in AD brain has not been well characterized.

Tau aggregated into oligomers and PHF in AD brain, which is visualized as a smear in western blots (Grundke-Iqbal et al., 1986a,b; Lee et al., 1991; Köpke et al., 1993). In the present study we investigated the phosphorylation and truncation of aggregated tau in post-mortem AD brain tissue by western blots. We found that high molecular weight tau oligomers (HMW-tau) from AD brain were hyperphosphorylated at multiple sites and truncated dominantly at the N-terminus.

MATERIALS AND METHODS

Human Brain Tissue

Frozen frontal cortices from autopsied and histopathologically confirmed AD and age-matched normal human brains (Table 1) were obtained without identification of donors from the Sun Health Research Institute Donation Program (Sun City, AZ, USA). Brain samples were stored at −80°C until used. The use of autopsied frozen human brain tissue was in accordance with the National Institutes of Health guidelines and was exempted by the Institutional Review Board (IRB) of New York State Institute for Basic Research in Developmental Disabilities because “the research does not involve intervention or interaction with the individuals” nor “is the information individually identifiable”.

Western Blots

The brain tissue was homogenized in cold buffer consisting of 50 mM Tris-HCl, pH 7.4, 8.5% sucrose, 2.0 mM EDTA, 10 mM β-mercaptoethanol, 1.0 mM orthovanadate, 50 mM NaF, 1.0 mM 4-(2-aminoethyl) benzenesulfonyl fluoride hydrochloride (AEBSF), and 10 µg/ml each of aprotinin, leupeptin, and pepstatin and stored at −80°C for western blots analysis. Brain homogenates were diluted in 2× Laemmli SDS sample buffer at 1:1 ratio, followed by boiling for 5 min. Samples were subjected to 7.5% SDS-PAGE and electrically blotted onto polyvinylidene fluoride membrane (Millipore) in transfer buffer (25 mM Tris-Glycine, pH 8.8, 20% methanol) at 0.5 amper for 2 h. The membrane was subsequently blocked with 5% fat-free milk-TBS for 30 min, incubated with primary antibodies (Table 2) in TBS overnight, washed with TBST (TBS with 0.1% Tween20), incubated with HRP-conjugated secondary antibody for 2 h at RT, washed with TBST, and incubated with the ECL western Blotting Substrate (Thermo scientific) and exposed to HyBlot CL® autoradiography film (Denville Scientific, Inc., Holliston, MA, USA). Specific immunostaining was quantified by using the Multi Gauge software V3.0 from Fuji Film.

Statistical Analysis

Data points were compared by unpaired two-tailed Student’s t-test (for data with normal distribution) or Mann-Whitney test (for data with non-normal distribution) and one-way analysis of variance (ANOVA) followed by Tukey’s multiple comparisons test. The data are presented as the mean ± SD. To reveal the association between high molecular weight aggregated tau and position of the epitope of the antibodies, Spearman correlation analysis was performed. p < 0.05 was considered statistically significant.

RESULTS

HMW-tau Is Present in AD Brain Homogenates

The aggregated tau forms oligomers and PHFs in AD brain, which are seen as a smear on western blots.

| Case | Age at death (Year) | Gender | PMI (h) | Braak stage | Tangle score |
|------|---------------------|--------|---------|-------------|-------------|
| AD (n = 17) | 80.59 ± 6.70 | 10 | 2.52 ± 0.65 | 5.65 ± 0.49 | 13.57 ± 2.05 |
| Con (n = 17) | 82.59 ± 5.36 | 10 | 2.59 ± 0.53 | 2.18 ± 0.88 | 3.14 ± 1.89 |
| AD (n = 10) | 76.60 ± 3.60 | 4 | 2.57 ± 0.71 | 5.70 ± 0.48 | 14.27 ± 1.40 |
| Con (n = 10) | 79.80 ± 4.94 | 5 | 2.38 ± 0.55 | 1.70 ± 0.67 | 2.43 ± 1.86 |

4PMI, postmortem interval; 5Neurofibrillary pathology was staged according to Braak and Braak (1991); 6Tangle score was a density estimate and was designated as none, sparse, moderate or frequent (0, 1, 2 or 3 for statistics), as defined according to CERAD Alzheimer’s disease criteria (Mirra et al., 1991). Five areas (frontal, temporal, parietal, hippocampal and entorhinal) were examined, and the scores were combined for a maximum of 15. 7Cases used in Figure 4.
TABLE 2 | Primary antibodies employed in this study.

| Antibody         | Type  | Species | Specificity       | Reference/Source (catalog/lot number) |
|------------------|-------|---------|-------------------|---------------------------------------|
| Anti-pT181-tau   | Poly- | R       | p-tau (T181)      | Invitrogen                            |
| Anti-pS199-tau   | Poly- | R       | p-tau (S199)      | Invitrogen (44734Q)                    |
| AT8              | Mono- | M       | p-tau (S202/T205) | Thermo Scientific (MN1020)            |
| Anti-pT205-tau   | Poly- | R       | p-tau (T205)      | Invitrogen (44738G)                    |
| Anti-pT212-tau   | Poly- | R       | p-tau (T212)      | Invitrogen (44740G)                    |
| Anti-pS214-tau   | Poly- | R       | p-tau (S214)      | Invitrogen (44742G)                    |
| Anti-pT217-tau   | Poly- | R       | p-tau (T217)      | Invitrogen (44744)                    |
| 12E8             | Mono- | M       | p-tau (S262/356)  | Dr. D. Schenk                         |
| Anti-pS396-tau   | Poly- | R       | p-tau (S396)      | Invitrogen (44752G)                    |
| Anti-pS404-tau   | Poly- | R       | p-tau (S404)      | Invitrogen (44–758G)                   |
| R145d            | Mono- | M       | 3R-tau            | Millipore (05–803/JBC1863429)         |
| RD4              | Mono- | M       | 4R-tau            | Millipore (05–804/2073108)            |
| Tau-1            | Mono- | M       | Up-tau (S199/202) | Binder et al. (1998)                  |
| QCB23070         | Poly- | R       | Up-tau (S46)      | Gond et al. (1994)                    |
| 43D              | Mono- | M       | Pan-tau (a.a. 6–18)| Liu et al. (2009)                     |
| HT7              | Mono- | M       | Pan-tau (a.a. 159–163)| Thermo Scientific (MN10000)        |
| Tau 46.1         | Mono- | M       | Pan-tau (a.a. 428–441)| Millipore (05–838-MI)               |
| Tau 46           | Mono- | M       | Pan-tau (a.a. 403–421)| Invitrogen (19–4603)                 |
| Tau C3           | Mono- | M       | Truncated tau at D421| Invitrogen (AHB0061)                |
| Anti-GAPDH       | Poly- | R       | GAPDH             | Santa-Cruz (sc-25778)                 |

Abbreviations: Mono-, monoclonal; p-, phosphorylated; Up-, unphosphorylated; Poly-, polyclonal; M, Mouse; R, Rabbit.

**FIGURE 1 |** High-molecular weight tau (HMW-tau) is selectively present in Alzheimer's disease (AD) brain. (A) Frontal cortical homogenates from 17 control and 17 AD cases were analyzed by western blots developed with a pan-tau antibody, 92e. (B) Blots were analyzed by densitometry. The levels of total tau, HMW-tau, low molecular tau (LMW-tau), and ratio of the HMW-tau/LMW-tau are presented as scattered dots with mean ± SD. **p < 0.01; ****p < 0.0001.

(Hundke-Iqbal et al., 1986a,b; Lee et al., 1991). To learn the nature of tau in AD and control brains, frontal cortical homogenates from 17 controls and 17 AD cases from two cohorts (Table 1) were analyzed by western blots developed with a pan-tau antibody, 92e. We observed several bands of tau from 50 kDa to 65 kDa immuno-recognized by 92e in control brains (Figure 1A). Tau in AD brain homogenates appeared as a smear (Figure 1A). We quantified the level of total tau, HMW-tau (65 kDa upwards) and low molecular weight (LMW-tau; 65 kDa downwards), as indicated in Figure 1A, by densitometry. We found that the total level of tau was ~5-fold increase in AD brains (Figures 1A,B). The HMW-tau, which is considered as aggregated tau, was only seen in AD brain (Figure 1B). Lower molecular weight tau (LMW-tau) was increased by about 2.5-fold in AD brain (Figure 1B). The ratio of HMW-tau/LMW-tau was 0.78 in AD and ~0.04 in control cases. These results suggest that AD brain is characterized by a smeared aggregation of tau.

**HMW-tau Is Abnormally Hyperphosphorylated at Multiple Sites in AD Brain**

Tau is hyperphosphorylated in AD brain at multiple sites (Köpke et al., 1993; Liu et al., 2009). To determine association of phosphorylation with HMW-tau, we analyzed tau phosphorylation in AD and control brain homogenates by western blots developed with several site-specific and phosphorylation dependent tau antibodies (Table 2). We found that phosphorylation of HMW-tau in control brains was barely detectable (Figure 2A). In contrast in AD brains, HMW-tau was phosphorylated at all phosphorylation sites studied, which include Ser181, Ser199, Ser202/Thr205, Thr205, Thr212, Ser214, Thr217, Ser262/356, Ser396, Ser404 and Ser422 (Figures 2A,B). Compared with control brains, phosphorylation level of LMW-tau was increased at all studied phosphorylation sites, except Ser199, in AD brains (Figures 2A,C). These results
suggest that hyperphosphorylated tau is strongly associated with HMW-tau.

To investigate whether non-hyperphosphorylated tau is also present in the HMW-tau smears, the western blots of above brain homogenates were developed with QCB23070 against tau unphosphorylated at Ser46 and Tau-1 against tau unphosphorylated at Ser198/199/202, respectively. We found that the level of tau recognized by either QCB23070 or Tau-1 was decreased in AD brain (Figure 3). Furthermore, HMW-tau in AD brain was recognized by neither QCB23070 nor Tau-1,
suggestion that HMW-tau is phosphorylated at both Ser46 and Ser198/199/202. This finding confirms the involvement of hyperphosphorylation of tau in AD pathogenesis.

**N-terminal Truncation of Tau Is More Associated than Its C-terminal Truncation with HMW-tau in AD**

Several truncations of tau have been identified in AD brain, especially in NFTs (Novak et al., 1991; Basurto-Islas et al., 2008). To learn the N- and C-terminal truncations of HMW-tau, we analyzed tau in the above 10 AD and 10 control cases (see Table 1) by western blots developed with phosphorylation-independent monoclonal antibodies against different regions of the tau molecule, including 43D (a.a. 6–18), HT7 (a.a. 159–163), RD4 (a.a. 275–291), RD3 (a.a. 267–274 and 306–313), Tau46 (a.a. 404–421) and Tau46.1 (a.a. 428–441; Figure 4A). Consistent with above studies, no HMW-tau was observed in control brain by any of the antibodies used in the present study, except Con 2 by RD4 (Figure 4B). Level of LMW-tau in AD brain was similar to that in control brain detected with 43D and HT7, ~2-fold increased with RD3, Tau46 and Tau46.1, and slightly decreased with RD4 (Figures 4B,C). These data suggest that the levels of total LMW-tau and LMW-3R-tau are increased and LMW-4R-tau is decreased in AD brain. HMW-tau was present in AD brains detected with all the antibodies (Figures 4B,C). HMW-tau immuno-reacted weakly with N-terminal antibody 43D and HT7, and strongly with C-terminal antibodies Tau46 and Tau46.1 (Figure 4B). The level of HMW-tau was gradually increased with antibodies epitopes range from N-terminal to C-terminal (Figure 4B). The ratio of HMW/LMW was dramatically increased in AD brain compared with control brains (Figure 4C). These results suggest that the N-terminus may be truncated in the HMW-tau.

Subsequently, we compared the ratio of HMW-tau/LMW-tau in AD brains detected by these six antibodies against different tau epitopes. We found that the ratios of HMW-tau/LMW were significantly different, except between HT-7 and RD4. The ratio of HMW-tau/LMW-tau showed a significant increase as the epitopes of tau antibodies range from N-terminal to C terminal (Figure 4D), suggesting that truncation of tau at the N-terminus increases its aggregation. Furthermore, the ratio of HMW-tau/LMW-tau with Tau6.1 (a.a. 428–441) was decreased as compared with that detected by Tau46, suggesting that truncation of the last 14 amino acids of tau may also increase in the HMW-tau.

**D421 Truncation Is Increased in AD Brain, but Not Associated with HMW-tau**

Truncation of tau at Aps421 (tauD421) has been identified in AD brain, especially in NFTs, and it is believed that this truncation may enhance the aggregation of tau into filaments (Abrahama et al., 2000; Gambin et al., 2003). To determine the relevance of D421 truncation in tau aggregation in AD brain, we analyzed the brain homogenates by western blots developed with Tau-C3, which specifically recognizes tauD421. By analyzing blots developed for short and long exposure time (Figure 5A), we found only a weak signal and no difference in the level of tauD421 in HMW-tau (Figure 5B); a ~2-fold increase was, however, seen in LMW-tau in AD (Figure 5B). The ratio of HMW-tauD421/LMW-tauD421 was decreased in AD brain (Figure 5B). These data suggest that C-terminal D421 truncation is probably not a significant player in HMW-tau in AD.
DISCUSSION

Conversion of monomeric tau to oligomeric and filamentous aggregates is apparently central to tau pathogenesis in AD and related tauopathies (Iqbal et al., 2016). Tau in AD, especially in NFTs is hyperphosphorylated and truncated. In the present study, by analyzing post-mortem brain tissues, we determined the relevance of hyperphosphorylation and truncation of tau in HMW-tau aggregates in AD brain. We found HMW-tau only in AD brain, but not in control brain. The HMW-tau...
was hyperphosphorylated at Thr181, Ser199, Ser202, Thr205, Thr212, Ser214, Thr217, Ser262, Ser396, Ser404 and Ser422. Tau unphosphorylated at Ser46 or Ser198/199/202 was not found in the HMW-tau aggregates. HMW-tau lacked the extreme N-terminal portion of tau suggesting that N-terminal truncated tau should be more relevant than the C-terminal extreme N-terminal portion of tau suggesting that N-terminal found in the HMW-tau aggregates. HMW-tau lacked the Tau unphosphorylated at Ser46 or Ser198/199/202 was not hyperphosphorylated at Thr181, Ser199, Ser202, Thr205, Thr212, Ser214, Thr217, Ser262, Ser396, Ser404 and Ser422. Tau truncated at D421, tau441 was increased in AD brain, but not significantly associated with the HMW-tau as detected by western blots. The level of tau with extreme C-terminus, a.a. 428–441 was reduced in HMW-tau. These findings suggest that hyperphosphorylated and N-terminal truncated and not C-terminal truncated tau is expected.

In AD brain, tau pathology starts from the entorhinal cortex, spreads to the hippocampus and frontal and temporal cortices, and finally to all isocortical areas, but the cerebellum is spared from tau lesions. Six stages of disease propagation are distinguished with respect to the location of the tangle-bearing neurons and the severity of changes by Braak. Braak stages I and II are used when NFT involvement is confined mainly to the transentorhinal region of the brain, stages III and IV when there is also involvement of limbic regions such as the hippocampus, and V and VI when there is extensive neocortical involvement (Braak and Braak, 1991). In the present study, we analyzed tau in frontal cortex by extensive neocortical involvement (Braak and Braak, 1991).

Stage I–IV, at which frontal cortex may not develop tau lesion. However, all AD cases are Braak Stage V–VI, in which tau pathology spreads to all isocortical areas, including frontal cortex. We found a great level of HMW-tau and truncation of tau species in AD brain detected by various anti-tau antibodies. Furthermore, we speculate a higher level of HMW-tau and truncated tau in entorhinal cortex and the hippocampus, brain regions in which more advanced tau pathology would be expected.

As a phospho-protein, the biological activity of tau is regulated by its degree of phosphorylation. Tau441, the longest tau isoform in the human brain, has 80 Ser/Thr and 5 Tyr potential phosphorylation sites. Normal human brain tau contains 2–3 moles phosphates/mole of the protein, while the abnormally hyperphosphorylated tau from AD brain has a phosphorylation stoichiometry of 9–10 (Köpke et al., 1993). To date, more than 40 phosphorylation sites of tau from AD brain have been identified (Wang and Liu, 2008). Compared with the age- and post-mortem match controls, the level of tau phosphorylation at multiple sites is increased significantly in AD brain (Liu et al., 2009). In the present study, we found that the level of phosphorylation of tau at all phosphorylation sites examined is dramatically increased and is greatly associated with HMW-tau in AD brain, suggesting a great relevance of the hyperphosphorylation with the aggregation of tau.

Hyperphosphorylated tau from AD brain (AD p-tau), the major protein subunit of PHF, is able to self-assemble into PHF and is unable to promote the assembly of MTs in vitro (Alonso et al., 1994). Dephosphorylation of AD p-tau by protein phosphatases restores its biological activity to promote assembly of MTs and inhibits its self-polymerization into PHF/straight filaments (SF; Wang et al., 1996a; Wang J. Z. et al., 2007). Phosphorylation of recombinant tau by PKA or Dyrk1A suppresses its capability of promoting MT assembly (Liu et al., 2007) and phosphorylation by the kinases in rat brain extract induces the self-assembly of tau into tangles of PHF and SF (Alonso et al., 2001). Furthermore, AD p-tau sequestrates normal tau into tangles of filaments (Alonso et al., 1996), indicating its prion-like property. Dephosphorylation of AD P-tau with alkaline phosphatase abolishes its ability to aggregate with normal tau and prevents tangle formation (Alonso et al., 1996). Furthermore, propagation of AD p-tau in vivo is altered by dephosphorylation with PP2A, suggesting hyperphosphorylation determines its prion-like spreading (Hu et al., 2016).

Tau protein is natively unfolded or intrinsically disordered and has little tendency for aggregation (Mukrasch et al., 2009). However, the tau molecule shows a preference for changing its global conformations to form a paperclip-like conformation by folding the N- and C-terminal portions back on the MT binding repeats (Jeganathan et al., 2006), which might prevent tau from aggregation. Tau fragments truncated at both N- and C-termini that contain MT binding repeats have a higher tendency for aggregation (Wang Y. P. et al., 2007), probably resulting from the disruption of the paperclip structure. Thus, both the N- and C-termini appear to have an inhibitory effect on the aggregation potential of tau. The N-terminal fragments of
tau inhibit tau441 polymerization by interacting with a specific C-terminal sequences and stabilizing a soluble conformation of tau (Horowitz et al., 2006). In the present study, we found that in the HWM-tau from frontal cortex of AD was barely immunorecognized by 43D, suggesting that the HMW-tau lacks the N-termiu and that the N-terminal truncation of tau may promote tau aggregation to form NFTs in vivo. Interestingly, we recently found that 43D effectively inhibits tau pathology in 3XTg-AD mice (Dai et al., 2017). Since normal tau can be sequestered by hyperphosphorylated tau from AD brain (Alonso et al., 1996), we speculate that 43D suppresses tau pathology by acting on normal tau which is-associated with pathological tau.

Tau is a substrate for various proteases (Wang et al., 2010). It can be cleaved by caspase 6 at Asp113 and Asp402, by caspase 3 at Asp25 and Asp421, by chymotrypsin at Tyr197, by an unknown throbmin-like cytosolic protease at Lys257, by asparaginyl endopeptidase at Asn255 and Asn368, and by calpain at Lys44 and Arg230 (Wang et al., 2010). Glu391 is cleaved by an unknown protease (Wang et al., 2010). PSA proteolyses residues stepwise from the N-terminus of tau (Wang et al., 2010; Wang and Mandelkow, 2016). Among all truncations of tau, truncations at Glu (E391) and Asp (D421) have been mostly reported in AD brain. Both truncations were reported to make tau proteins more prone to aggregation than the full-length tau (Abraha et al., 2000; Berry et al., 2003; Gamblin et al., 2003; Yin et al., 2006). Tau4E391 not only greatly increases the rate of in vitro filament formation, but also increases the amount of the tau protein assembled into fibrillar structures (Abraha et al., 2000). PHFs isolated from AD brain and treated with pronase (to remove the fuzzy coat) contain tau fragment tau151-391 (Wischik et al., 1988), which is prone to aggregation. Rats transgenic for tau151-391 develop neurofibrillary pathology and also display hyperphosphorylation and production of HMW-tau species (Zilka et al., 2006). However, in the present study, we observed a great level of C-terminal portion of tau in HMW-tau detected by antibodies Tau46 and Tau46.1, suggesting that majority of tau in AD brain is not truncated at the C-terminal. It is reported that pseudophosphorylation of tau at Ser422 suppresses D421 or E391 truncation (Guillozet-Bongaarts et al., 2006). In AD brain, tau is hyperphosphorylated at Ser396, Ser404 and Ser422, which may inhibit these two C-terminal truncations. However, tau4D421 was increased and the level of tau with intact C-terminal end as detected by tau46.1 was significantly lower in HMW-tau than that of tau recognized by Tau46, suggesting that this C-terminal truncation may be involved in tau aggregation.

In conclusion, in the present study we found that HMW-tau are hyperphosphorylated at multiple sites, and unphosphorylated tau, such as at Ser46 and at Ser198/199/202, is absent in these aggregates. The N-terminal truncated tau is associated with HMW-tau more than the C-terminal truncated tau. These findings provide a novel insight into the association of the hyperphosphorylation and truncation of tau with tau aggregation into oligomers in AD.

**AUTHOR CONTRIBUTIONS**

FL, WH and KI: conception of the research. YZ, JS, DC, ZG and FL: performing experiments. YZ, WH and FL: analyses and interpretation of results. WH and FL: drafting of the manuscript. WH, C-XG and KI: critical revision of the manuscript.

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Conflict of Interest Statement: The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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