Tau Phosphorylation Sites Work in Concert to Promote Neurotoxicity In Vivo

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INTRODUCTION

Phosphorylation of the microtubule-associated protein tau is elevated in neurodegenerative disease tissue, and tau-positive neurofibrillary pathology is characteristic of several disorders, including Alzheimer’s disease and other tauopathies (Feany and Dickson, 1996; Lee et al., 2001). This abnormal hyperphosphorylation of tau induces an increase in the apparent molecular mass of the protein, which is observed in disease states (Lee et al., 1991). Proline-directed kinases phosphorylate tau at serine-proline (SP) or threonine-proline (TP) motifs, many of which reside in the proline-rich regions flanking the microtubule binding domains of human tau. These phosphorylation events have been linked particularly to tau-induced neurodegeneration because a group of antibodies that specifically react to phosphorylated SP and TP sites recognize tau found in the brain tissue of Alzheimer’s disease patients but not autopsy-derived tau from nondemented age-matched individuals. This group of monoclonal antibodies recognizes single or two closely spaced phosphorylated SP/TP sites, including pT181 (AT270) (Goedert et al., 1994), pS202/pT205 (AT8) (Goedert et al., 1995), pT212/pS214 (AT100) (Zheng-Fischhofer et al., 1998), pT231/pS235 (TG3) (Jicha et al., 1997a), and pS396/pS404 (PHF1) (Otvos et al., 1994). Phosphorylation at SP/TP sites is a striking feature of tau from patients with Alzheimer’s disease and other tauopathies; consequently, these sites are commonly referred to as “disease-associated” sites. Aside from their development as specific diagnostic markers of tauopathies (for review, see Gong et al., 2005), monoclonal antibodies that recognize particular phosphopeptides of tau have also been used to study the correlation between phosphorylation and disease severity (Augustinack et al., 2002) and to delineate the hierarchical pattern of tau phosphorylation (Zheng-Fischhofer et al., 1998; Shea and Cressman, 1999).

To understand how phosphorylation contributes to tau-induced neurodegeneration, many laboratories have focused on the kinases and phosphatases that regulate tau phosphorylation. Adopting a different strategy, we used a Drosophila tauopathy model (Wittmann et al., 2001) to investigate the direct targets of these enzymes, namely, the SP and TP sites of tau, and we found that they are critical regulators of toxicity in vivo. We showed that blocking phosphorylation of all 14 SP/TP sites in tau by mutating them to alanine markedly inhibited tau-induced neurodegeneration in Drosophila (Fulga et al., 2007; Steinhilb et al., 2007). Conversely, mutation of all SP/TP sites to glutamate generated the pseudophosphorylated tau\textsuperscript{E14} construct, which is significantly more toxic than tau\textsuperscript{WT} (Khurana et al., 2006; Dias-Santagata et al., 2007; Fulga et al., 2007).

In this report, we used our Drosophila tauopathy model to investigate how phosphorylation of individual SP/TP sites impacts tau-induced neurotoxicity. We generated phosphorylation-incompetent forms of tau in which single phosphorylation sites, or a pair of sites, were replaced by alanines. Our results revealed that although SP/TP sites are critical mediators of neurotoxicity, there is no single site whose phosphorylation is essential for neurodegeneration. These findings suggest that SP/TP sites function together to promote neurotoxicity in vivo.
MATERIALS AND METHODS

Constructions, Genetics, and Stocks

The UAS-tauWT and UAS-tauΔC transgenic Drosophila lines have been described previously (Wittmann et al., 2001; Steinhilb et al., 2007). The vector pBStau24 (Wittmann et al., 2001) served as a template for the generation of alanine mutations at the 14 Ser/Thr-prosites of human tau by using the QuickChange site-directed mutagenesis kit (Stratagene, La Jolla, CA). The codons for T111, T153, T175, T181, S199, S202, T205, T212, S214, T217, S235, S239, S404, S406, and S422 were mutated to alanine individually or in pairs, and the resulting constructs were subcloned into the pUAST vector by using the restriction enzymes KpnI and NotI. The authenticity of all constructs was confirmed by sequencing, and lines with expression level equivalent to tauWT were determined by quantitative Western blot analysis. The following transgenic strains are described in Flybase (http://flybase.bio.indiana.edu/), and they were obtained from the Bloomington Drosophila Stock Center (Indiana University, Bloomington, IN) (das-GALA, UAS-p53), N. Dyson (Massachusetts General Hospital) (UAS-Rbf), J. K. Hariharan (Massachusetts General Hospital) (UAS-dap), C. Lehner (University of Bayreuth) (UAS-CycE), and T. Xu (Yale University) (UAS-Tsc2AΔRT). Crosses were maintained on standard cornmeal-based Drosophila medium at 25°C. Flies were analyzed at 1 to 3 d of age.

Western Blot

Heads from adult flies at 1 d posteclosion were homogenized in Laemmli buffer (Sigma-Aldrich, St. Louis, MO), boiled for 10 min at 100°C, and centrifuged at 13,000 × g for 2 min to remove insoluble debris. Proteins were resolved by SDS-polyacrylamide gel electrophoresis (PAGE) by using a 10% separating gel (Camberx, East Rutherford, NJ), transferred to nitrocellulose (Bio-Rad, Hercules, CA), blocked in 5% milk in phosphate-buffered saline with 0.1% Tween 20, and immunoblotted using one of the following antibodies: polyclonal anti-C terminal tau (Dako North America, Carpinteria, CA) at 1:10,000 dilution; AT8 (Innogenetics, Zwijndrecht, Antwerp, Belgium) at 1:1,000 dilution; Tau1 (Chemicon International, Temecula, CA) at 1:10,000 dilution; TG3 (P. Davies, Bronx, NY) at 1:100 dilution; PHF1 (P. Davies) at 1:500 dilution; pSer212 (BioSource International, Camarillo, CA) at 1:40,000; pSer214 (BioSource International) at 1:40,000; AT270 (Pierce-Endogen, Rockford, IL) at 1:1,000,000 dilution; and AT100 (Innogenetics) at 1:1,000 dilution. For each blot, the same number of fly heads was used for each genotype, Ponceau S stains were performed to ensure equivalent total protein loading and even transfer, and blots were reprobed for total tau (Dako North America). Western blots were repeated a minimum of three times with different animals, and representative blots are shown. The appropriate anti-mouse or anti-rabbit horseradish peroxidase-conjugated secondary antibody was applied and signals were detected by chemiluminescence (Pierce-Endogen).

Histology

Heads from adult flies at 1 to 3 d posteclosion were fixed in Formalin, embedded in paraffin, and 4-μm frontal sections were prepared. Serial sections were cut through the entire brain and placed on a single glass slide, and immunostaining was performed as described previously (Khurana et al., 2006). MC1 and Aii50 were used at dilutions of 1:1, and they were detected by AlexaFluor 488-conjugated secondary antibodies.

Phosphatase Treatment

Drosophila head homogenates were prepared in 1X lambda phosphatase buffer (New England Biolabs) containing a protease inhibitor cocktail (Roche Diagnostics, Indianapolis, IN) and 200 μg of total protein were incubated with 800 U of lambda protein phosphatase for 3 h at 37°C. Homogenates were then subjected to SDS-PAGE and immunoblot analysis as described above.

RESULTS

Generation of Individual SP/TP Transgenic Tau Mutants

To determine the relative contribution of individual SP/TP phosphorylation sites to tau toxicity, we generated eight transgenic Drosophila lines expressing mutant forms of human tau in which the following serine and/or threonine residues were replaced by alanine: tau111/153 (T111A, T153A); tau175/181 (T175A, T181A); tau199/202/205 (S199A, T205A, S202A); tau212/214/235 (T212A, T214A, S235A); tau396/404 (S396A, S404A); and tau422 (S422A) (Figure 1). Many of these sites, when phosphorylated, are recognized by a group of monoclonal antibodies that preferentially react with tau found in patients with neurodegenerative disorders. These antibodies include AT8, AT100, AT270, TG3, and PHF1, and the phosphopeptides recognized by each of them are schematized in Figure 1. Although not an SP site, we also generated tau214 (S214A), because Thr212 and Ser214 create the phosphoepitope recognized by the AT100 antibody, one of the most specific correlates of toxicity in human disease (Zheng-Fischhofer et al., 1998).

The Toxicity of Individual SP/TP Tau Mutants Is Similar to That of TauWT

To test whether abolishing phosphorylation at individual sites would affect tau-induced neurotoxicity, wild-type and mutant forms of human tau were expressed in the Drosophila retina under the control of the GMR-GAL4 driver (Figure 2). In contrast to the normal fly eye (Figure 2A), expression of wild-type human tau (tauWT) in the retina results in toxicity that can be visualized as a moderately rough eye phenotype caused by disruption of the ordered arrangement of ommatidia (Shulman and Feany, 2003; Figure 2B). We have recently reported that expression of a phosphorylation-incompetent tau (tauΔP), in which 14 disease-associated SP/TP sites have been replaced by alanine, has a marked effect on toxicity, almost completely blocking tau-induced toxicity in vivo (Fulga et al., 2007; Steinhilb et al., 2007; Figure 2C). In contrast, expression of equivalent levels of each of the individual point mutants in the retina resulted in a moderately rough eye (Figure 2, D–L) similar to that induced by tauWT (Figure 2B). These results indicate that blocking phosphorylation of individual SP/TP sites has no detectable effect on tau-induced toxicity in vivo. Because the phosphorylation state of tau depends on the cellular context within the nervous system (Grammenoudi et al., 2006), it is possible that individual site contribute differently to toxicity in other types of neurons. Unfortunately, the modest toxicity of wild-type human tau (Wittmann et al., 2001) limited our ability to assess the toxicity of our mutants in the brain.
Mutation of individual SP/TP sites does not significantly alter tau toxicity in vivo. Compared with the organized structure of a normal eye (A), expression of tauWT in the eye (under the control of the GMR-GAL4 driver) induces retinal degeneration and a rough eye phenotype (B). TauAP, in which all 14 SP/TP sites have been mutated to alanine, is indistinguishable from that induced by tauWT.

Figure 2. Mutation of individual SP/TP sites does not significantly alter tau toxicity in vivo. Compared with the organized structure of a normal eye (A), expression of tauWT in the eye (under the control of the GMR-GAL4 driver) induces retinal degeneration and a rough eye phenotype (B). TauAP, in which all 14 SP/TP sites have been mutated to alanine, is indistinguishable from that induced by tauWT. Control is GMR-GAL4/+. To examine whether single site phosphorylation affects the appearance of disease-specific conformations of tau, we used the antibodies MC1 and Alz50, which recognize abnormal conformations of tau associated with Alzheimer’s disease and related disorders (Hyman et al., 1988; Carmel et al., 1996; Vincent et al., 1996). Immunostaining of retinal sections with MC1 yielded no signal for nontransgenic control flies (Figure 3A), but it detected equivalent levels of immunoreactivity in tauWT (3B), and flies expressing the individual points mutants. Results from tau202/205 (3C), tau231/235 (3D), and tau396/404 (3E) are shown as representative examples. We obtained similar results using the conformation-specific antibody Alz50 (Figure 3, F–J). These data suggest that blocking phosphorylation at individual sites does not preclude tau from adopting disease-associated conformations. This finding contrasts with what we observed with the phosphorylation-incompetent mutant tauWT, in which mutation of all disease-associated SP/TP sites significantly reduces the propensity of tau to assume disease-specific conformations (Steinhilb et al., 2007). Thus, although SP/TP phosphorylation of tau is required for the generation of abnormal conformations and for neurotoxicity (Steinhilb et al., 2007), there is no single phosphorylation site, or pair of sites, that accounts for these disease-related features.

Neurotoxicity Induced by SP/TP Tau Mutants Is Mediated by TOR Signaling and Cell Cycle Activation

Our model provides a relatively rapid and flexible system in which to investigate the mechanisms of tau-induced neurodegeneration, and our recent findings suggest that expression of tau in Drosophila induces the sequential activation of the target of rapamycin (TOR) signaling pathway and of the cell cycle, leading to neurodegeneration (Khurana et al., 2006). Because cell cycle activation depends upon tau phosphorylation (Khurana et al., 2006), we tested whether the mutation of single phosphoepitopes would affect the ability of TOR signaling, the cell cycle or of apoptotic pathways to mediate tau neurotoxicity.

Blocking TOR signaling by coexpression of a constitutively active form of the TOR inhibitor Tsc2 (Tsc2AKT+) rescued tau induced toxicity in our model (Khurana et al., 2006) and suppressed to similar extents the rough eye phenotypes of tauWT (Figure 4A), tau202/205 (Figure 4B), tau231/235 (Figure 4C), and tau396/404 (Figure 4D) transgenic flies. In contrast, overexpression of Rheb, a positive modulator of the TOR pathway, enhanced tau-induced toxicity in flies expressing tauWT (Figure 4E), tau202/205 (Figure 4F), tau231/235 (Figure 4G), and tau396/404 (Figure 4H).

Expressing the Cdk2 inhibitor Dacapo (Dap, Drosophila homologue of human p21/p27) together with the E2F1 inhibitor retinoblastoma factor-1 (Rbf1) synergistically blocks the G1/S transition of the cell cycle, and it suppresses tau-induced neurodegeneration (Khurana et al., 2006). Coexpression of Dap and Rbf1 substantially suppressed the tau-induced retinal toxicity of tauWT (Figure 4I) and of the individual SP/TP mutants. Results with tau202/205 (Figure 4J), tau231/235 (Figure 4K), and tau396/404 (Figure 4L) are shown as representative examples. Conversely, coexpression of cyclin E, which catalyzes the G1/S transition together with Cdk2, strongly enhanced to equivalent levels tau toxicity in transgenic flies expressing tauWT (Figure 4M), tau202/205 (Figure 4N), tau231/235 (Figure 4O), and tau396/404 (Figure 4P).

Finally, coexpression of the apoptosis inhibitor p35 in tau transgenic animals suppressed the rough eye phenotype of tauWT (Jackson et al., 2002; Figure 4Q), tau202/205 (Figure 4R), tau231/235 (Figure 4S), tau396/404 (Figure 4T) and of the other individual SP/TP mutants.

In summary, modulating TOR signaling, cell cycle activation, or apoptosis modified to similar extents the neurodegenerative phenotypes induced by wild-type and SP/TP-mutant forms of tau. Thus, although in our model cell cycle activation and neuronal death depend on tau phosphorylation (Khurana...
Hyperphosphorylation of Tau Is Triggered by Largely Independent Events in Transgenic Flies

Several studies have used in vitro kinase assays to examine the process of tau phosphorylation, and they have suggested that the generation of disease-associated forms of tau requires a complex sequence of phosphorylation steps (Zheng-Fischhofer et al., 1998; Shea and Cressman, 1999). Expression of phosphorylation-incompetent tau mutants in Drosophila provided us with a system to address this question in vivo. If the steps leading to tau phosphorylation are temporally ordered and interdependent, we expect that inhibiting phosphorylation at a critical SP/TP site will prevent downstream phosphorylation events that require prior modification of that residue.

To examine the phosphorylation pattern of each tau construct, we prepared homogenates from transgenic flies expressing either tauWT or each of the individual SP/TP mutants for Western blot analysis. Nontransgenic controls showed no reactivity for any of the antibodies used (Steinhilb et al., 2007 and data not shown). As seen in Figure 5A, all extracts showed substantial reactivity with the AT270 antibody, with the exception of those derived from tau175/181-expressing flies (Figure 5A, first row, lane 2), as expected because AT270 detects phosphorylation at T175/T181 (Goe-der et al., 1994). Similarly, the TG3 antibody recognized tau in all extracts with the exception of tau231/235 (Figure 5A, fourth row, lane 6), as anticipated because phosphorylation at these two sites defines the TG3 epitope (Jicha et al., 1997a). The PHF1 antibody recognizes pS396/pS404 (Otvos et al., 1994); thus, it did not react with extracts derived from tau396/404 transgenic animals (Figure 5A, fifth row, lane 7), but it identified all the other mutant forms of tau. These results suggest that blocking phosphorylation of individual SP/TP residues in vivo does not prevent phosphorylation of the remaining sites.

We also examined the connection between the AT8 and Tau1 epitopes. Although the AT8 recognition site includes the remaining sites. Phospho-specific antibodies pThr212 and pSer214 recognize the AT100 antibody (Figure 5B, first row, lanes 4 and 10). Phospho-specific antibodies pThr212 and pSer214 recognize each individual site, and were used to investigate the relationship between these two phosphorylation events in vivo. As shown in Figure 5B, our data suggest that phosphorylation of T212 and S214 are interdependent because mutation of S214 to alanine partially inhibited phosphorylation at T212 (second row, lane 10). However, the T212A mutation did not appreciably alter phosphorylation at S214 (third row, lane 4). Interestingly, blocking phosphorylation at S199/T217 had a significant impact on AT100 immunoreactivity. As shown in Figure 5B, extracts from tau199/217-expressing flies were not recognized by AT100 (first row, lane 3). Analyses with the pThr212 and pSer214 antibodies re-
revealed that tau<sup>199/217</sup> is phosphorylated at T212 (Figure 5B, second row, lane 3) but not at S214 (third row, lane 3), suggesting that in vivo phosphorylation of S199 and/or T217 is essential for S214 phosphorylation and for the appearance of the highly disease-selective AT100 epitope. In addition, our results establish that, in vivo, phosphorylation of other individual SP/TP sites is not required for AT100 generation.

**Blocking Phosphorylation of a Select Group of SP/TP Sites Does Not Inhibit Tau-induced Toxicity**

We recently showed that mutation of all 14 SP/TP sites to alanine generated the phosphorylation-incompetent tau<sup>AP</sup> construct, which has markedly reduced toxicity in Drosophila compared with tau<sup>WT</sup> (Fulga et al., 2007; Steinhilb et al., 2007). Conversely, the pseudophosphorylated construct tau<sup>E14</sup>, in which all SP/TP sites are mutated to glutamate, is much more toxic than tau<sup>WT</sup> (Khurana et al., 2006; Dias-Santagata et al., 2007; Fulga et al., 2007). These observations demonstrate that SP/TP phosphorylation is critical for tau-induced neurodegeneration, but our data thus far show that neurotoxicity cannot be suppressed by blocking phosphorylation at a single site, or pair of sites (Figure 2). Based on these results, we suspected that a larger group of sites might be working together to induce toxicity. To test this hypothesis, we generated transgenic lines expressing tau<sup>AP5</sup>, a con-
Blocking phosphorylation at five SP/TP sites does not inhibit tau toxicity. (A) The tauAP mutant has the following five mutations: S202A, T205A, T212A, T231A, and S235A. This manipulation disrupts reactivity with the AT8, AT100, and TG3 antibodies. (B) TauAP-expressing flies have the same retinal toxicity as tauWT flies. (C) Western blot analysis confirms that tauAP-expressing flies are not reactive for AT8, AT100, and TG3, whereas phosphorylation at other disease-associated sites, including T181 (AT270), S214 (pSer214), and S396/S404 (PHF1) is maintained. Reactivity with the Tau1 antibody is also observed. TauWT and tauAP expression levels are equivalent, as demonstrated by immunoblot by using a polyclonal total tau antibody. Driver is GMR-GAL4.

Phosphorylation at S396/S404 Has a Substantial Effect on the Electrophoretic Mobility of Tau

It is well established that aberrantly phosphorylated tau, characteristic of disease states, exhibits a shift toward lower mobility on SDS-PAGE (Baudier and Cole, 1987; Flament and Delacourte, 1989; Greenberg and Davies, 1990). Phosphatase treatment studies revealed that the decreased electrophoretic mobility of disease-associated tau is mainly due to phosphorylation (Lee et al., 1991; Ksiezak-Reding et al., 1992). Consistent with these reports, we found that treatment of fly brain homogenates with λ-phosphatase increased the mobility of tauWT ((Steinhilb et al., 2007) and Figure 7, compare lanes 3 and 4). Moreover, mutation of all SP/TP sites to alanine in tauAP increased its electrophoretic mobility compared with tauWT (Steinhilb et al., 2007; Figure 7, lanes 1 and 3). Phosphatase treatment increased the migration of tauAP modestly (Steinhilb et al., 2007; Figure 7, lanes 1 and 2), indicating that the small shift in molecular weight observed in tauAP before treatment is due to phosphorylation at non-SP/TP sites.

To determine whether single SP/TP phosphorylation events substantially affect tau mobility in vivo, we tested all individual SP/TP mutants and found that tauS396/404 had an increased electrophoretic mobility compared with tauWT (Figure 7, compare lanes 3 and 5). We confirmed that this difference in mobility was a consequence of phosphorylation, by treating with lambda phosphatase (lane 6). In contrast to tauS396/404, the electrophoretic mobility of all other individual SP/TP point mutants was equivalent to that of tauWT.
analysis of events. T. phosphorylation in vivo relies on a hierarchical sequence of tau phosphorylation sites affecting neurotoxicity in an animal model, and whether human disease pathogenesis. We tested whether inhibiting tau phosphorylation at the five SP/TP sites abolishes reactivity with three important disease-specific antibodies without affecting tau-induced toxicity (Figure 6). Thus, we have established that inhibiting tau phosphorylation at restricted SP/TP sites in vivo does not reduce toxicity, suggesting that multiple sites work together to promote neurodegeneration.

Abnormal conformations of tau, recognized by specific antibodies such as Alz50 and MC1, have been associated with degeneration and can be detected before the appearance of neurofibrillary pathology (Hyman et al., 1988; Jicha et al., 1999a). Notably, tau phosphorylation may help stabilize these aberrant conformations (Carmel et al., 1996). In agreement with this hypothesis, we showed that inhibiting phosphorylation of all SP/TP sites significantly decreases the generation of disease-associated conformations of tau (Steinhilb et al., 2007). In contrast, transgenic flies expressing individual SP/TP tau mutants have similar levels of abnormally folded protein as tauWT-expressing flies (Figure 3), further underscoring the correlation between hyperphosphorylation, conformational changes, and tau-induced neurotoxicity.

Markers consistent with aberrant activation of the cell cycle and alterations in TOR signaling have been described in Alzheimer’s disease tissue (An et al., 2003; Li et al., 2004; Griffin et al., 2005), and these two pathways are critical mediators of tau-induced toxicity in our model (Khurana et al., 2006). Here, we show that the neurodegenerative phenotype of individual SP/TP mutants also relies on the activation of TOR signaling and the cell cycle (Figure 4), which indicates that blocking individual phosphorylation sites does not alter the mechanisms of tau-induced neurotoxicity.

Several studies suggest that tau phosphorylation at specific sites relies on a sequential and interdependent order of events (Kimura et al., 1996; Zheng-Fischhofer et al., 1998; Shea and Cressman, 1999; Augustinack et al., 2002). However, we show that individual SP/TP mutations do not completely prevent phosphorylation from occurring at other sites, showing that in vivo phosphorylation of individual SP/TP sites do not absolutely require phosphorylation at other SP/TP (Figure 5). However, a degree of interdependence is shown by partial reduction in AT8 immunoreactivity in flies expressing tauWT (Figure 5A and Supplementary Figure S1). In addition, full phosphorylation at serine 212 requires phosphorylation of the non-SP/TP site serine 214 (Figure 5B).

Historically, the AT8 and Tau1 antibodies were considered to be complementary in that AT8 reacted with phosphorylated serines 199 and/or 202 and 205, whereas Tau1 recognized this same stretch of amino acids in an unphosphorylated form (Biernat et al., 1992; Goedert et al., 1995). Because the time of the original mapping, the relationship between these antibodies has been shown to be less coincident (Szendrei et al., 1993; Shea and Cressman, 1999). Our results are in agreement with these latter observations. Although the tau202/205 mutant is not recognized by AT8, as expected, its reactivity with Tau1 is not elevated compared with that of tauWT (Figure 5). Instead, we show that the tau199/217 mutant has substantially increased reactivity with

**Figure 7.** Phosphorylation of S396/S404 significantly alters the electrophoretic mobility of tau. Drosophila head homogenates derived from transgenic flies expressing wild type and mutant forms of tau were incubated in the presence or absence of lambda phosphatase, followed by prolonged electrophoretic separation (to more easily detect differences in migration by SDS-PAGE) and by Western immunoblotting with a polyclonal tau antibody. Driver is GMR-GAL4.

tauWT, as can be appreciated for tau202/205, included in Figure 7 (lanes 7 and 8) as a representative of all other mutants. We also determined that blocking phosphorylation at the five SP/TP sites in tauWT had only a modest effect on tau migration compared with the effect of S396A/S404A mutations (compare lanes 3, 5 and 9). As shown in Figure 7, phosphatase treatment caused all of the tau proteins to migrate at the same molecular weight (lanes 2, 4, 6, 8, and 10), suggesting that phosphorylation is responsible for the major weight differences observed by SDS-PAGE.

Thus, in contrast to other individual SP/TP sites, blocking phosphorylation at S396/S404 resulted in a significant increase in the electrophoretic mobility of tau, which suggests that phosphorylation of S396 and/or S404 in vivo may contribute substantially to the mobility shift of hyperphosphorylated tau.

**DISCUSSION**

Hyperphosphorylation of tau has been linked extensively to neurodegeneration (Lee et al., 2001), and we and others have previously shown that SP/TP phosphorylation is critical for toxicity in a Drosophila model of human neurodegenerative disorders, including Alzheimer’s (Jackson et al., 2002; Nishimura et al., 2004; Khurana et al., 2006; Dias-Santagata et al., 2007; Fulga et al., 2007; Steinhilb et al., 2007). However, despite the extensive attention devoted to single phosphorylation sites throughout the literature, we now show that in vivo phosphorylation of individual SP/TP sites work together to promote neurodegeneration.

The significance of individual phosphorylation sites and the temporal and spatial appearance of specific phosphoepitopes of tau have been primarily investigated using purified proteins in in vitro kinase reactions or in tissue culture models using transient overexpression (Zheng-Fischhofer et al., 1998; Shea and Cressman, 1999). A major advantage of our Drosophila model is that, in a time- and cost-effective manner, we can ask biologically relevant questions in the context of a whole nervous system that mimics aspects of human disease pathogenesis. We tested whether inhibiting tau phosphorylation at each of the major disease-associated sites affects neurotoxicity in an animal model, and whether tau phosphorylation in vivo relies on a hierarchical sequence of events.

Mutating all SP/TP sites to alanine significantly ameliorates tau toxicity (Fulga et al., 2007; Steinhilb et al., 2007) and replacing them by phospho-mimetic glutamates exacerbates tau-induced neurodegeneration (Khurana et al., 2006; Dias-Santagata et al., 2007; Fulga et al., 2007). Although these findings support the hypothesis that SP/TP phosphorylation plays a fundamental role in neurotoxicity, we now show that mutating individual SP/TP sites to alanine has no clear effect on tau-induced degeneration in vivo. As represented in Figure 2, when expressed in the Drosophila retina, each of the individual SP/TP point mutants induces the same moderately rough eye as tauWT. Furthermore, blocking phosphorylation at five SP/TP sites abolishes reactivity with three important disease-specific antibodies without affecting tau-induced toxicity (Figure 6).

Thus, we have established that inhibiting tau phosphorylation at restricted SP/TP sites in vivo does not reduce toxicity, suggesting that multiple sites work together to promote neurodegeneration.

Abnormal conformations of tau, recognized by specific antibodies such as Alz50 and MC1, have been associated with degeneration and can be detected before the appearance of neurofibrillary pathology (Hyman et al., 1988; Jicha et al., 1999a). Notably, tau phosphorylation may help stabilize these aberrant conformations (Carmel et al., 1996). In agreement with this hypothesis, we showed that inhibiting phosphorylation of all SP/TP sites significantly decreases the generation of disease-associated conformations of tau (Steinhilb et al., 2007). In contrast, transgenic flies expressing individual SP/TP tau mutants have similar levels of abnormally folded protein as tauWT-expressing flies (Figure 3), further underscoring the correlation between hyperphosphorylation, conformational changes, and tau-induced neurotoxicity.

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![Figure 7](image-url)
Tau1, suggesting that Tau1 is more specific for the unphosphorylated residues S199 and/or T217 than S202/T205.

The AT100 antibody is one of the most specific correlates of neurodegeneration, because it recognizes tau present in the brains of patients with Alzheimer’s disease, but it does not tend to cross-react significantly with normal tissue (Zheng-Fischhofer et al., 1998). Thus, in an attempt to understand disease pathogenesis, much attention has been devoted to elucidating the steps required for the formation of this epitope. AT100 specifically reacts with two phosphorylated residues, pT212 and pS214 (Zheng-Fischhofer et al., 1998). Although some studies suggest that the process of AT100 generation first requires phosphorylation of T212 by glycogen synthase kinase (GSK)3β followed by phosphorylation at S214 by protein kinase A (PKA) (Scott et al., 1993; Zheng-Fischhofer et al., 1998; Jicha et al., 1999b), other reports suggest that prephosphorylation of tau at Ser214 by nonproline-directed kinases, including PKA, greatly enhances the ability of GSK to phosphorylate tau at T212 (Singh et al., 1995; Gong et al., 2005). Using our Drosophila model, in which kinases and phosphatases are endogenously regulated, we find that mutating either T212 or S214 to alanine does not prevent phosphorylation at S214 or T212, respectively (Figure 5), strongly suggesting that, in vivo, phosphorylation of T212 does not require prior phosphorylation of S214, and vice versa.

Interestingly, we show that inhibiting phosphorylation at S199/T217 prevents phosphorylation at the non-SP/TP site S214, which completely blocks formation of the AT100 epitope (Figure 5). Although S199 and T217 lie outside the recognition motif of AT100, our finding is consistent with other reports suggesting that phosphorylation of these sites is important for AT100 reactivity. Interestingly, previous in vitro studies found that a recombinant tau construct containing the T217A mutation could not form the AT100 epitope when treated with purified kinases (Zheng-Fischhofer et al., 1998; Yoshida and Goedert, 2006). Also, in autopsy-derived brain tissue from Alzheimer’s disease patients, AT100 immunostaining is associated with phosphorylation of S199 (Maurage et al., 2003). Together, our results suggest that, in addition to the phosphorylation of its recognition motif (pT212/pS214), tau phosphorylation at S199 and/or T217 is a prerequisite for AT100 generation in vivo.

Finally, we demonstrate that the decrease in electrophoretic mobility characteristic of hyperphosphorylated tau depends on phosphorylation at S396/S404 (Figure 7). We also show that blocking phosphorylation at five SP/TP sites in the tau1475 mutant does not as significantly affect tau mobility, consistent with the notion that a few site-specific phosphorylation events may contribute disproportionately to altered tau migration. Interestingly, previous in vitro studies had argued that the reduced electrophoretic mobility of disease-associated tau might not due to the particularly high degree of phosphorylation, but rather to phosphorylation at specific residues. Similar observations from other groups support a significant effect of C-terminal phosphorylation on the electrophoretic mobility of tau (Steiner et al., 1990; Scott et al., 1993).

In summary, our data suggest that there is no single SP/TP phosphorylation site in tau that preferentially endows the protein with neurotoxicity, suggesting that the sites work together to mediate tau toxicity in vivo. These observations are important because, according to in vitro data, the two proline-directed protein kinases that have been most convincingly implicated in tau pathology, GSK-3β and cdk5, can target multiple SP/TP sites in tau (for review, see Gong et al., 2005). Moreover, blocking GSK-3β and cdk5 activities has been shown to be protective in cell culture systems and animal models of neurodegeneration (Zheng et al., 2002; Mudher et al., 2004; Noble et al., 2005). Thus, together with our previous studies demonstrating the critical role of SP/TP phosphorylation in tau neurotoxicity (Khurana et al., 2006; Dias-Santagata et al., 2007; Fulga et al., 2007; Steinhilb et al., 2007), our current findings underscore the therapeutic potential of kinase inhibitors that can target multiple proline-directed phosphorylation sites for the treatment of Alzheimer’s disease and other tauopathies.

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