Reversible Paired Helical Filament-Like Phosphorylation of Tau Is an Adaptive Process Associated with Neuronal Plasticity in Hibernating Animals

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Introduction

Formation of paired helical filaments (PHFs) is one of the critical neuropathological hallmarks of Alzheimer’s disease (AD). Although the microtubule-associated protein tau in a hyperphosphorylated form has been established as primary constituent though the microtubule-associated protein tau in a hyperphosphorylated form is a major hallmark of Alzheimer’s disease and related disorders. The process of tau phosphorylation, thought to be of critical importance for PHF formation, and its potential link to neurodegeneration, however, is not understood very well, mostly because of the lack of a physiological in vivo model of PHF-like tau phosphorylation. Here we describe the formation of highly phosphorylated tau, containing a number of PHF-like epitopes in torpor during hibernation. PHF-like phosphorylation of tau was not associated with fibril formation and was fully reversible after arousal. Distribution of PHF-like tau followed a consistent pattern, being most intense in the entorhinal cortex, hippocampus, and isocortical areas. Within the hippocampus, a particularly high labeling was seen in CA3 pyramidal cells. Somewhat lesser reactivity was present in CA1 neurons while dentate gyrus granule cells were not reactive. Formation of PHF-like tau in CA3 neurons was paralleled by the regression of synaptic contacts of the mossy fiber system terminating on CA3 apical dendrites. Mossy fiber afferentation was re-established during arousal, concomitantly with the decrease of PHF-like tau in CA3 neurons.

These findings implicate an essential link between neuronal plasticity and PHF-like phosphorylation of tau. The repeated formation and degradation of PHF-like tau might, thus, represent a physiological mechanism not necessarily associated with pathological effects. Hibernation will, therefore, be a valuable model to study the regulation of PHF-like tau-phosphorylation and its cell biological sequelae under physiological in vivo conditions.

Key words: Alzheimer’s disease; hibernation; natural hypothermia; PHF; phosphorylation; plasticity; synapse; tau

Neurofibrillary pathology [paired helical filaments (PHFs)] formed by the microtubule-associated protein tau in a hyperphosphorylated form is a major hallmark of Alzheimer’s disease and related disorders. The process of tau phosphorylation, though to be of critical importance for PHF formation, and its potential link to neurodegeneration, however, is not understood very well, mostly because of the lack of a physiological in vivo model of PHF-like tau phosphorylation. Here we describe the formation of highly phosphorylated tau, containing a number of PHF-like epitopes in torpor during hibernation. PHF-like phosphorylation of tau was not associated with fibril formation and was fully reversible after arousal. Distribution of PHF-like tau followed a consistent pattern, being most intense in the entorhinal cortex, hippocampus, and isocortical areas. Within the hippocampus, a particularly high labeling was seen in CA3 pyramidal cells. Somewhat lesser reactivity was present in CA1 neurons while dentate gyrus granule cells were not reactive. Formation of PHF-like tau in CA3 neurons was paralleled by the regression of synaptic contacts of the mossy fiber system terminating on CA3 apical dendrites. Mossy fiber afferentation was re-established during arousal, concomitantly with the decrease of PHF-like tau in CA3 neurons.

These findings implicate an essential link between neuronal plasticity and PHF-like phosphorylation of tau. The repeated formation and degradation of PHF-like tau might, thus, represent a physiological mechanism not necessarily associated with pathological effects. Hibernation will, therefore, be a valuable model to study the regulation of PHF-like tau-phosphorylation and its cell biological sequelae under physiological in vivo conditions.

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are most early and most severely affected. Failures of synaptic plasticity are, thus, assumed to represent early events in the course of AD (Ashford and Jarvik, 1985; Cotman and Anderson, 1988; Flood and Coleman, 1990; Geddes and Cotman, 1991; Swaab, 1991; Mesulam, 1999) that eventually lead to alteration of tau phosphorylation. The potential link between synaptic plasticity, synaptic detachment, and the regulation of tau phosphorylation, however, has not been addressed directly under physiological in vivo conditions.

In the present study, we have used the hibernation cycle, a physiological model of adaptation associated with an extraordinary high degree of structural neuronal plasticity, to analyze the potential relationship between synaptic plasticity and alterations in tau phosphorylation. It has been shown previously, that during torpor, a natural state of hypothermia, synaptic contacts between mossy fibers and hippocampal pyramidal neurons undergo dramatic regressive changes that are fully reversible very rapidly during euthermy (Popov and Bocharova, 1992; Popov et al., 1992). Here, we demonstrate that this rapid, reversible, and repeated regression of synaptic and dendritic components on CA3 neurons is associated with a reversible PHF-like phosphorylation of tau at a similar time course. These findings implicate an essential
link between neuronal plasticity and PHF-like phosphorylation of tau, one of the major hallmarks of AD. Hibernation will, thus, represent a model to study the regulation of PHF-like tau phosphorylation and its cell biological sequelae under physiological in vivo conditions.

Materials and Methods

Animals. The 36 adult European ground squirrels (Spermophilus citellus) used in this study were either captured from a dense population near Vienna (Millesi et al., 2001), born in the laboratory from females that were caught pregnant, or bred in outdoor enclosures in Haren, The Netherlands (Hut et al. 2002a). The animals were kept in Lucite cages (length × width × height = 48 × 28 × 50 cm) with a wooden nestbox attached (15 × 15 × 15 cm). Wood shavings were used as bedding material, and food (rabbit breeding chow, Teurlings, Waalwijk, The Netherlands) and water were supplied ad libitum. The animals were kept in a climate-controlled room at a relative humidity of 60% throughout the experiment. Hibernation was induced by gradually lowering ambient temperature from 20 to 7°C and changing light conditions from a 12 hr light/dark cycle to continuous dim red light (< 1 lux) in autumn. Individual torpor-arousal patterns were assessed by measuring nestbox temperatures every minute with a computer-based recording system or by recording outside nestbox activity (Oklejewicz et al., 2001; Hut et al., 2002b). Registration of torpor-arousal patterns was validated by using customized abdominal temperature loggers (Tidbit; Onset; Hut et al. 2002a,b) that registered body temperature every 48 min. This study was approved by the Animal Experiments Committee of the University of Groningen (BG02198).

Experimental design. In total, 36 animals were studied, 32 for immunohistochemistry and an additional 4 for Western blotting. The animals were killed in four different stages within the torpor and arousal periods in hibernation: torpor short (TS; n = 6), torpor long (TL; n = 6), arousal short (AS; n = 6), and arousal long (AL; n = 7). Eleven animals were killed after becoming continuously euthermic after hibernation (EU; n = 11). Hibernating animals showed hypothermic periods of 11.01 ± 0.18 d (mean ± SEM) and euthermic periods (including arousal) of 21.13 ± 0.38 hr before killing. Brain material of the hibernating animals was collected after 1.53 ± 0.06 hr (AS), 8.27 ± 0.05 hr (AL), 2.34 ± 0.21 d (TS), and 7.11 ± 0.1 d (TL) after the onset of the previous (final) arousal to euthermic body temperatures. Body temperatures after start of perfusion confirmed that the groups represented the specific phases of hibernation: rectal temperatures were 30.9 ± 1.6°C (AS), 34.5 ± 0.3°C (AL), 9.8 ± 1.5°C (TS), and 8.21 ± 0.3°C (TL). Arousal was induced by gentle handling at room temperature for 3–5 min. Arousal induction was performed at least 10 weeks after onset of hibernation. At the time of brain material collection, the duration of the previously experienced torpor phase did not differ between the groups.

Brain material of nonhibernating animals (EU) was collected 6–7 d after cessation of hibernation, initiated by an increase in ambient temperature from 7 to 25°C in early spring (Hut et al., 2002b). Body temperature after start of perfusion was 36.5 ± 0.4°C (mean ± SEM). All animals were killed with 2 ml of 6% pentobarbital 10 min before perfusion (immunohistochemistry, electron microscopy) or decapitation (Western blotting).

Immunohistochemistry. Animals were transcardially perfused with 4% paraformaldehyde in phosphate buffer. Brains were equilibrated with 30% sucrose in phosphate buffer and cut coronally on a freezing microtome into 25 μm sections. Series of free-floating sections from 32 animals of all groups (five to six animals per group) were applied to the immunoperoxidase labeling of PHF-like phosphorylated tau protein (AT8, 1:2000; Innogenetics, Zwijndrecht, Belgium), MAP2 (mouse clone HM-2; 1:1000; Sigma, Taufkirchen, Germany), PSA–NCAM (mouse clone 2B–28B; 1:500; AbCys S.A., Paris, France), and synaptophysin (rabbit; 1:1600; Dako, Hamburg, Germany).

In brief, all sections were processed with a streptavidin–biotin technique and nickel-enhanced diaminobenzidine as chromogen as previously described (Härtig et al., 1995). For double immunofluorescence of AT8 staining and MAP2, sections were incubated overnight with a mixture of AT8 (1:400) and rabbit anti-MAP2 (1:250; Chemicon, Temecula, CA). Immunoreactivities were visualized with a cocktail consisting of Cy3-conjugated goat anti-mouse IgG and Cy2-tagged goat anti-rabbit IgG (Dianova, Hamburg, Germany). Controls were performed by omitting the primary antibodies resulting in the absence of any cellular staining.

Relative strength of immunoreactivity in the hippocampal stratum lucidum (PSA–NCAM, synaptophysin, MAP2) and over somata of CA3 pyramidal neurons (AT8) was comparatively determined by densitometric measurements on sections processed in parallel with nickel-enhanced diaminobenzidine. Ten regions of interest (50 × 50 μm) were randomly selected for each region, and the optical density, corrected for background, was obtained using the image processing and analyzing system (Münster, Germany) connected to a xrs camera (SL Microtest, Jena, Germany) attached to a Zeiss Axioskop microscope. Five sections were analyzed for each animal (group size: n = 5 or 6 animals).

Electron microscopy. For electron microscopy, animals were transcardially perfused with 0.1 ml PBS, followed by 300 ml of ice-cold fixative containing 4% paraformaldehyde and 2.5% glutaraldehyde in 0.1 ml PBS, pH 7.4. After postfixation in the same solution overnight (4°C), 70 μm vibratome sections were made (Vibratome series 1000), rinsed in cacodylic buffer, pH 7.2, and subsequently postfixed for 2 hr at room temperature with 1% OsO4 in 0.1 ml cacodylic buffer, pH 7.2. After a short washing in aqua bidest, tissue was dehydrated in a graded ethanol series (30, 50, 70, and 90% each 10 min, 100% × 2, each 60 min, 100% propylene oxide × 2, each 7 min). Dehydrated sections were finally embedded in Epon 812 following routine procedures and cured for 48 hr at 60°C. Pieces of layer V prefrontal cortex were excised, and ultrathin sections cut on a Reichert Ultracut S were contrasted for 20 min with 5% uranyl acetate in aqua bidest and 2 min in Reynolds lead citrate. Sections were examined with a Zeiss (Oberkochen, Germany) EM 912 electron microscope.

Western blotting. Fresh brain material of four animals was collected for Western blotting. Three animals were hibernating, two matching the time that the AL group were killed (>7 hr after arousal induction, after >7 d of torpor) and one matching the time that the TL group were killed (>7 d of torpor). The fourth animal was killed during continuous euthermic in summer (EU group).

The animals were killed with 2 ml of 6% pentobarbital and decapitated. Brains were removed immediately and submerged in ice-cold PBS. Neocortex and hippocampus were dissected and homogenized in lysis buffer (20 mM Tris·HCl, pH 7.2, 2 mM MgCl2, 100 mM NaCl, 5 mM NaF, 1 mM Na3VO4, 0.5% NP-40, 1 mM DTT, 100 μg/ml PMSF, 2 μg/ml Leupeptin, 2 μg/ml Pepstatin A, and 600 μM okadaic acid (Sigma); ratio tissue to buffer: 1:5). After centrifugation (5000 × g, 30 min, 4°C), lysates were filled up with glyceral to a concentration of 50% (v/v). Protein contents were determined by the Bradford assay. Proteins were separated on 10 or 5% to 15% gradient SDS polyacrylamide gels using 30 μg of total protein per well and subsequently transferred to a polyvinylidene difluoride (PVDF) transfer membrane (PolyScreen; PerkinElmer Life Sciences, Boston, MA). Membranes were washed once in PBS, blocked in PBS containing 2% BSA (w/v), and probed with mouse anti-MAP2 (1:1000) or rabbit anti-synaptophysin (1:5000). PHF-like phosphorylated tau was detected by using the following mouse monoclonal antibodies (numbering of amino acid residues based on the 441 amino acids of human tau) (Goedert et al. 1989a): AT8 (Ser202/Thr205; 1:1000; Innogenetics, Zwijndrecht, Belgium); AT100 (Ser212/Thr214; 1:1000; Innogenetics), AT180 (Thr231; 1:1000; Innogenetics), AT270 (Thr181; 1:1000; Innogenetics), PHF-1 (Ser396/Ser404; 1:2000; courtesy of P. Davies, Albert Einstein College of Medicine, Bronx, NY) and 12E8 (Thr231; 1:1000, Dako, Hamburg, Germany).
Results
Hibernation induces PHF-like phosphorylation of tau

Effects of hibernation on the phosphorylation stage of tau were analyzed by Western blotting of cortical and hippocampal extracts and immunohistochemistry. A mobility shift of tau was observed after the transition from euthermic to torpor, as visualized by the phosphorylation-independent detection of tau with the antibody BR134 (Fig. 1). In the euthermic ground squirrels, this antibody detects bands at 68, 70, and 72 kDa. During torpor, the 68 kDa band is greatly diminished with the appearance of immunoreactive bands of >72 kDa. This mobility shift is accompanied by an enhancement of immunoreactivity for all phosphorylation-dependent antibodies tested, detecting six distinct PHF-like tau phosphoepitopes on tau (AT8, AT100, AT180, AT270, PHF-1, and 12E8) and tau, unphosphorylated at Ser198, Ser231, and Ser235. The relative abundance of four-repeat to three-repeat tau isoforms was determined by RT-PCR using forward primer TJF (GGC TAC AGC AGC CCC GGC TC) and reverse primer TJR. The cycling profile consists of an initial denaturation step 5 min at 94°C repeated 35 times, an annealing step 40 sec at 58°C, and a final elongation step 7 min 72°C. The PCR product was gel-extracted and ligated blunt into cloning vector pZErO-2 (Invitrogen). Clones containing insert were screened by colony-PCR, and the longest, most abundant, tau isoform was sequenced. The cDNA sequence was translated into amino acid sequence and was aligned with rat and human tau amino acid sequences. This cDNA corresponds to the longest human isoform containing exon 2, 3, and 10 because rodent tau isoforms are shorter than corresponding human isoforms, and in adult rodent brains this isofrom is most abundant. The pattern of isoforms and their relative expression levels were unaffected during the hibernation cycle.

To identify and assign the major ground squirrel tau isoform on Western Blot we cloned and sequenced the most abundant tau cDNA. This cDNA corresponds to the longest human isoform containing exon 2, 3, and 10 because rodent tau isoforms are shorter than corresponding human isoforms, and in adult rodent brains this isoform is most abundant. The pattern of isoforms and their relative expression levels were unaffected during the hibernation cycle.

Figure 1. Phosphorylation of tau protein in the neocortex and hippocampus in an euthermic animal (EU), in long torpor (TL), and after long arousal (AL). Immunoblots were reacted for phosphorylation-independent detection of tau (BR134), specific PHF-like tau phosphoepitopes (AT8, AT100, AT180, AT270, PHF-1, 12E8) and tau, unphosphorylated at Ser198, Ser231, and Ser235 (Tau-1).

After enzymatical dephosphorylation, a pattern of five isoforms was resolved (Fig. 2). The strongest BR134-immunoreactive band is the tau isoform with the lowest electrophoretic mobility in the ground squirrel and aligns with human recombinant tau isoform comprising exon 2 and exon 3. Presumably this band represents the longest ground squirrel tau isoform containing exon 2, 3, and 10 because rodent tau isoforms are shorter than corresponding human isoforms, and in adult rodent brains this isoform is most abundant. The pattern of isoforms and their relative expression levels were unaffected during the hibernation cycle.

To identify and assign the major ground squirrel tau isoform on Western Blot we cloned and sequenced the most abundant tau cDNA. This cDNA corresponds to the longest human isoform containing exon 2, 3, and 10. The ground squirrel amino acids sequence has two deletions in exon 1, a 11 amino acid deletion, which is typical for rodent tau sequences and a unique 2 amino acid deletion. This explains the faster electrophoretic mobility of the longest tau protein isoform after dephosphorylation in squirrel compared with human. RT-PCR of a tau cDNA fragment comprising exon 10 shows a more prominent occurrence of exon 10 containing tau mRNA in ground squirrel brain than in
human brain (Fig. 3). The cladogram derived from comparison of tau protein sequences of different mammals shows that the deduced amino acid sequence of ground squirrel has strongest similarities to rodents like mice and rats but has also similarities to human sequence (Fig. 4). To simplify the analysis we compared variable amino acids in ground squirrel sequence to human and rat sequence only. We found 14 amino acid substitutions and one gap of 11 amino acids, which are identical between ground squirrel and rat, whereas 19 amino acids are alike between ground squirrel and human, and another 17 amino acids and a two amino acid gap are unique for ground squirrel.

Furthermore, the amino acid sequence shows that the epitopes of phosphorylation-dependent and independent antibodies used to label tau protein in ground squirrel are sufficiently conserved.

When the distribution of phospho-tau was analyzed immunohistochemically, strongest reactivity for AT8 was found in the ventral hippocampus, entorhinal cortex, and isocortex (Fig. 5). A particularly high labeling was seen in hippocampal CA3 pyramidal cells. To a lesser extent reactivity was also present in CA1 pyramidal neurons, whereas dentate gyrus granule neurons were not reactive (Fig. 6). Marked labeling was also seen in subcortical areas, in particular in hypothalamic and epithalamic nuclei, whereas thalamic nuclei were only marginally reactive.

Both Congo red and thioflavin S staining for the detection of fibrillar aggregates were negative. Electron microscopic investigation of the cortex from torpid animals did not reveal any evidence for the formation of neurofibrillary aggregations within the dendritic compartment (Fig. 7).

Reversibility of tau phosphorylation after arousal

Reversibility of tau phosphorylation was of particular interest in the present paradigm, because incomplete reversibility would likely result in the formation of PHFs. As seen on Western blots (Fig. 1) and immunohistochemical preparations (Figs. 5, 6), increased tau phosphorylation was fully reversible after arousal. Already a few hours after animals were awake, a reversibility of the mobility shift of tau, associated with a diminished reactivity for all phosphorylation-dependent anti-tau antibodies, was observed (Fig. 1).

The process of reversible generation of phospho-tau epitopes and their subcellular distribution was analyzed in more detail in the hippocampus (Fig. 6). Immunoreactivity for AT8 was typically found in the somatodendritic compartment of CA3 pyramidal neurons and to a lesser extent also in CA1 and CA4 neurons, where it developed gradually. At shorter periods of torpor, labeling with AT8 became first detectable in the apical dendrite of pyramidal cells while the perikarya were still devoid of reactivity (Fig. 6). Conversely, after arousal, immunoreactivity first disappeared in the perikarya, whereas it remained somewhat longer in dendrites (Fig. 6). Reactivity for any phosphorylation-dependent antibody used in the study completely disappeared after longer periods of arousal (Fig. 6).

PHF-like phosphorylation–dephosphorylation of tau in the hibernation cycle is synchronized with a regression–re-establishment of afferentation

Cellular and subcellular pattern of phosphorylated tau present under the condition of torpor might be of particular relevance to the process of neuronal connectivity and plasticity. This relationship was specified further in the hippocampal mossy fiber system that reproducibly underwent cyclic changes during hibernation (Popov and Bocharova, 1992; Popov et al., 1992; Hut et al., 2001). The mossy fiber system arises in the dentate gyrus granule cells that remained completely devoid of AT8 reactivity during the whole hibernation cycle and terminates in the stratum lucidum on apical dendrites of CA3 pyramidal cells. The latter displayed a high AT8 reactivity during torpor.

As shown in the present study, the mossy fiber system can be identified by the presence of PSA-NCAM that in ground squirrels is not restricted to developmental stages but persists in the adult euthermic animal. During torpor, expression of PSA-NCAM dis-
appears almost completely with the exception of a few granule cells (Fig. 8A1, A2). After arousal, PSA-NCAM expression in the mossy fiber system gradually reappears (Fig. 8A3, A4). Correspondingly, immunoreactivity for synaptophysin in the CA3-stratum lucidum, where the mossy fibers contact CA3 apical dendrites disappears and reappears with the hibernation cycle (Figs. 6B, 7). Cyclic changes of synaptophysin immunoreactivity were also found by Strijkstra et al. (2003) based on a monoclonal antibody (Hut et al., 2001). These periodic changes during the cycle in presynaptic PSA-NCAM and synaptophysin are synchronized with the disappearance and reappearance of postsynaptic microtubule-associated protein (MAP2) in CA3 cells (Figs. 8C, 9), as reported by Hut et al. (2001). Our findings indicate a gradual regression of mossy fiber synapses and corresponding postsynaptic elements during torpor, associated with the accumulation of phosphorylated tau at postsynaptic sites, which completely reverses after arousal.

A synopsis of the synchronized changes in presynaptic and postsynaptic makers and phosphorylated tau obtained during the hibernation cycle on immunohistochemical preparations is displayed in Figure 10. In parallel with synaptic regression on CA3 cells, the accumulation of phosphorylated tau develops, reaching highest amounts during deep torpor. Highly phosphorylated tau decreases again during arousal and eventually disappears parallel to the process of re-expression of synaptophysin and MAP2.

Discussion

One of the first who clearly recognized the plasticity-related nature of Alzheimer pathology was Ramón y Cajal (1928). He also was the first who tried to mimic the pathology of AD, putting research on this disorder on an experimental basis. Among his pupils he prompted a number of studies intending to analyze sequelae of a disturbed metabolism induced by manipulations such as hibernation (Ramón y Cajal, 1904; Tello, 1904), starvation, and exposure to cold (Donaggio, 1906), combined with thyroidectomy or parathyroidectomy (Balli, 1906; Lewy, 1923; Rasdolsky, 1926; Alexander, 1934; Stern and Elliott, 1949). Even if these early studies failed to reproduce true neurofibrillary degeneration of AD, they were able to induce some kind of neurofibrillary changes (Balli, 1906; Donaggio, 1906; Lewy, 1923; Rasdolsky, 1926) associated with argyrophilic (Ramón y Cajal, 1904; Tello, 1904; Rasdolsky, 1926). In the light of more recent data on thyroid disease as a potential risk factor for AD (Kalmijn et al., 2000) and on the effects of starvation on tau phosphorylation (Yanagisawa et al., 1999; Planel et al., 2001), we re-evaluated a potential link between the hibernation cycle and neurofibrillary pathology made up by a hyperphosphorylated form of the microtubule-associated protein tau.

Hibernation is a behavioral strategy used by several mammalian species to minimize energy expenditure under inhospitable environmental conditions. During hibernation, overall metabolic rate as well as heart and respiratory rate are greatly reduced (Wang, 1978; Geiser and Kenagy, 1988), by tolerating a body temperature near the ambient temperature in a regulated hypothermic state. Low metabolic rate during torpor is accompanied by dramatically reduced neuronal functions. EEG measurements of torpid hibernators have shown that almost no brain activity is present (Walker et al., 1977; Krilowicz et al., 1988; Daan et al., 1991). Correspondingly, hibernation elicits negative effects on memory retention in conditioned tasks (Millesi et al., 2001). It was hypothesized that because of these potentially deleterious effects, hibernators interrupt the torpor state regularly to return.
These regularly occurring euthermic phases, called “arousal episodes”, are apparently necessary to protect against mechanisms that otherwise would lead to complete memory loss. They last 4–24 hr, depending on species and size (French, 1985). Arousals, however, are expensive in terms of energy (Wang, 1978, 1989; Kenagy et al., 1989). The adaptive significance of the hibernation cycle makes this an ideal model to investigate cellular sequelae of neuroplasticity and its potential association with the phosphorylation state of tau in a physiological setting.

During torpor, when brain temperature decreases to \(-15^\circ C\), electroencephalographic activity is strongly reduced (Krilowicz et al., 1988; Strijkstra et al., 1999; Gabriel et al., 1998). Because activity is a measure of use and neuronal connections remain functional through regular use, this decrease may negatively affect the maintenance of neuronal connections (Kavanau, 1997). Evidence for a reduced neuronal connectivity during the hibernation cycle has been provided for different brain regions in a variety of hibernating animals (Malinsky and Malinska, 1975, 1988; Malinsky, 1983; Malinsky and Polach, 1992; Popov and Bocharova, 1992; Strijkstra and Daan, 1997; Hut et al., 2002b). A cycle of synaptic regression during torpor and subsequent reinnervation in phases of arousal has been particularly well characterized for synaptic contacts between mossy fibers and hippocampal pyramidal neurons in ground squirrels (Popov and Bocharova, 1992; Popov et al., 1992; Hut et al., 2001).

We have been able to confirm and extend these studies on cyclic changes of the CA3 pyramidal cell afferentation by mossy fibers that are associated with changes in PSA-NCAM expression. PSA-NCAM is the developmentally regulated polysialylated form of the neural cell adhesion molecule NCAM, a cell surface glycoprotein that is involved in neuronal migration and neurite outgrowth (Seki and Arai, 1993, 1999; Durbec and Cremer, 2001). PSA-NCAM expression is related to bouton formation and remodeling, which accompanies synaptic formation (Seki and Arai, 1999), a process of critical importance for hippocampal plasticity in learning and memory (Fox et al., 1995; Muller et al., 1996; Murphy et al., 1996) and, as shown in the present study, for re-establishing neuronal connectivity during arousal.

One cellular mechanism that contributes to the regulated suppression of metabolism and thermogenesis during hibernation is reversible phosphorylation of enzymes and proteins that limits rates of flux through metabolic pathways (Storey 1987, 1997; Mehrani and Storey, 1997; Storey and Storey, 1997).
MacDonald and Storey, 1998, 1999; Chen et al., 2001). Reversible phosphorylation during hibernation also affects synaptic membrane proteins (Shchipakina et al., 1995), a process known to be involved in synaptic plasticity (Walaas and Greengard, 1991). Here we demonstrate that this mechanism of reversible protein phosphorylation also affects the microtubule-associated protein tau, thereby generating a condition that in the adult human brain is associated with neurofibrillary degeneration.

One remarkable feature is the apparent high molecular weight of tau extracted from euthermic animals ranging from 68 to 72 kDa, which is even shifted in the torpor animals. This slow electrophoretic mobility is reminiscent of PHF-tau and not present in biopsy-derived human tau or freshly prepared rodent tau. The retarded electrophoretic mobility in the euthermic and torpor state is solely attributable to tau phosphorylation, because dephosphorylation shifts tau from 72 to 63 kDa. A likely explanation is the presence of two additional potential phosphorylation sites in ground squirrel compared with human.

A high phosphorylation of tau at some PHF-like epitopes, e.g., at Ser-202, is also seen during normal development of the mammalian (Goedert et al., 1993) and non-mammalian (Rössner et al., 1994) brain, where it is downregulated during maturation. In the human brain, the switch from the highly phosphorylated to the less phosphorylated state occurs around birth (Goedert et al., 1993) and, thus, coincides with synaptogenesis, which is similar to what is seen in the present paradigm. The developmental regulation of tau, however, extends to the level of alternative mRNA splicing, giving rise to different isoform pattern. In fetal human brain there is a single isoform, but in adult brain there are six isoforms (Goedert et al., 1989a,b). At variance with this developmental shift of mRNA splicing, the pattern of six isoforms seen in the brain of adult ground squirrels was not affected by the hibernation cycle.

The present results, thus, indicate that regulating tau (hyper)phosphorylation is preserved in the adult mammalian brain as a naturally occurring process associated with specific requirements on neuroprotection and plasticity. It apparently reflects a physiological mechanism and is not necessarily associated with pathological effects. The necessity, however, of regular arousal phases to protect against permanent memory loss might indicate the potentially deleterious sequelae of this process if it lasts too long. The present results, thus, support the suggestion that hyperphosphorylation of tau reflects a protective mechanism in a unfavorable environment (Ihara, 2001). Recent studies investigating tau phosphorylation in relation to neuronal susceptibility for apoptosis have indeed suggested that a modest increase in tau phosphorylation correlates with increased protection of neurons against cell death (Lesort et al., 1997; Nagy and Esiri, 1997; Arendt et al., 1998b; Esclaire et al., 1998; Mills et al., 1998; Yardin et al., 1998).

By hibernation, animals may save up to 90% of the energy that would otherwise be required (Geiser, 1988; Wang, 1989; Heldmaier and Ruf, 1992; McKee and Andrews, 1992). In torpor, when the phosphorylation stage of tau is highest, the metabolic rate is strongly suppressed, often to <5% of the normal euthermic rate. This situation reflects a “vita minima”, where energy supply and requirements are both low but still balanced. Homeostatic control is preserved and brain damage does not occur. Hibernation is, therefore, associated with tolerance to deprivation of oxygen and glucose (Frerichs and Hallenbeck, 1998). In this
Hyperphosphorylation might confer tau resistance to pro- 


teases and could, thus, be a mechanism to stabilize its structure. 


Stabilization of cytoskeletal proteins might be a mechanism to “freeze” the dynamic structure during a "vita minima", prevent-


ing its degradation and preserving it for rapid activation in arousal phases. If synaptic connectivity is compromised during torpor and re-established very quickly in a similar way as before, a mecha-


anism must exist that "marks" those sites that will be occupied by synapses again. This mechanism might potentially involve stabi-


lized cytoskeletal proteins and need to be particularly well developed in neurons with a high neuroplastic potential. This would explain the hierarchy of neuronal vulnerability against neurofi-


brillary degeneration that has previously been shown to follow the pattern of the neuroplastic potential in the adult brain (Arendt et al., 1998a).


Most importantly, PHF-like phosphorylation of tau in the present model is fully reversible through a mechanism that oper-


ates naturally in the mammalian brain. This paradigm might, 


thus, be useful to study the physiological regulation of tau phos-


phorylation and dephosphorylation critically involved in the 


process of neurofibrillary degeneration in AD and related 


conditions.


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