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## **Središnja medicinska knjižnica**

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### **Human neuroblastoma SH-SY5Y cells treated with okadaic acid express phosphorylated high molecular weight tau-immunoreactive protein species**

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#### **Abstract**

*Background:* Early stages of Alzheimer's disease (AD) are characterized by high phosphorylation of microtubule-associated protein tau, which may result from the downregulation of protein phosphatases.

*New method:* In order to model phosphatase downregulation and analyze its effect on tau aggregation *in vitro*, we treated neuroblastoma SH-SY5Y cells with okadaic acid (OA), a protein phosphatase inhibitor, and examined high molecular weight phospho-tau species.

*Results and comparison with existing methods:* OA treatment led to the appearance of heatstable protein species with apparent molecular weight around 100 kDa, which were immunoreactive to anti-tau antibodies against phosphorylated Ser202 and Ser396. As these high molecular weight tau-immunoreactive proteins (HMW-TIPs) corresponded to the predicted size of two tau monomers, we considered the possibility that they represent phosphorylation-induced tau oligomers. We attempted to dissociate HMW-TIPs by urea and guanidine, as well as by alkaline phosphatase treatment, but HMW-TIPs were stable under all conditions tested. These characteristics resemble properties of certain sodium dodecyl sulfate (SDS)-resistant tau oligomers from AD brains. The absence of HMW-TIPs detection by antitotal tau antibodies Tau46, CP27 and Tau13 may be a consequence of epitope masking and protein truncation. Alternatively, HMW-TIPs may represent previously unreported phosphoproteins cross-reacting with tau.

*Conclusions:* Taken together, our data provide a novel characterization of an OA-based cell culture model in which OA induces the appearance of HMW-TIPs. These findings have implications for further studies of tau under the conditions of protein phosphatase downregulation, aiming to explain mechanisms involved in early events leading to AD.

#### **Highlights**

- OA treatment of SH-SY5Y cells induced 100 kDa tau-immunoreactive species
- Induced HMW species were reactive to anti-tau-pS202 and anti-tau-p396 antibodies
- HMW species were heat-stable and stable in SDS, urea and guanidine
- OA-induced HMW species could not be detected by several anti-total tau antibodies
- HMW proteins may represent tau oligomers or phosphoproteins cross-reacting with tau

**Keywords**: Alzheimer's disease, neurodegeneration, cell culture, SH-SY5Y, okadaic acid, oligomerization, phosphorylation, protein phosphatase, tau protein, immunoblot

#### **1. Introduction**

One of the neuropathological hallmarks of Alzheimer's disease (AD) is the aggregation of tau protein in neurofibrillary tangles (NFT) in the neurons and glia cells in the brain. Tau (tubulin-associated unit) is a microtubule-associated protein, expressed mainly in neurons of the central nervous system [1], where its primary function is to modulate microtubule dynamics [2]. In adult human brain, there are six tau splicing isoforms ranging from 352 to 441 amino acids (for review, see [3]). Isoforms differ by the absence or presence of one or two 29 amino acid inserts in the N-terminal part in combination with either three or four microtubule-binding (MTB) repeat regions in the C-terminal part (for review, see [4]).

Tau is normally a natively unfolded protein [5] with little tendency for aggregation, therefore an intriguing, but mostly unresolved question is to understand what causes wild-type tau molecules to oligomerize and subsequently form larger aggregates that form NFT. Pathways leading to tau aggregation are largely unclear. However, tau propensity to aggregate may be driven by posttranslational modifications, mainly phosphorylation and C-terminal truncation [6]. Despite a paucity of stable secondary structure, tau in solution seems to form a transient global fold in which N- and C-terminal arms fold over the central MTB repeat domain that contains the sequences with a tendency to form  $\beta$ -sheets and aggregate (known as the "paper clip" model [7]).

Phosphorylation decreases affinity of tau for microtubules and makes it more prone to aggregation (for review, see [8]). In AD brains more than thirty tau amino acid residues are specifically phosphorylated, including Ser202 and Ser396 [9]. Hyperphosphorylation of tau in AD seems to be a consequence of a disrupted balance between kinase and phosphatase activity [10]. Several protein phosphatases were found dysregulated in AD brain, including PP2A, PP1 and PP5, which are all capable of dephosphorylating tau *in vitro* (for review, see [11]. The main regulator of tau dephosphorylation in healthy human brain appears to be PP2A, since its activity accounted for about 70% of the tau dephosphorylation in the assay using brain extracts [12]. PP2A has been shown to dephosphorylate tau *in vitro* at multiple sites, including Ser202 and Ser396 [13]. PP2A affects tau phosphorylation levels not only directly, but also indirectly by regulating the activities of several tau kinases, most notably glycogen synthase kinase  $3\beta$  (GSK-3 $\beta$ ) and Ca<sup>2+</sup>/calmodulin-dependent protein kinase II (CaMKII) [14].

Studying the process of tau aggregation requires a suitable cell culture model. In this report we examined the potential of SH-SY5Y cell line treated with the phosphatase inhibitor okadaic acid (OA) as a model for studying the initial steps of tau aggregation. SH-SY5Y is a widely used human neuroblastoma cell line with an endogenous expression of tau and the capacity to differentiate into neuron-like cells [15-17]. Okadaic acid is a cell-permeable potent inhibitor of protein phosphatases PP2A, PP4, PP5 and PP1 [18, 19], which has been previously used to upregulate the levels of phospho-tau in cultured cells and to investigate the role of protein phosphatases in tau phosphorylation [20-24]. To investigate the potential of SH-SY5Y cells treated with protein phosphatase inhibitor OA as a model for studying the process of tau aggregation and neurodegeneration [25], we examined the effects of OA treatment on generation of high molecular weight tau. We observed that the incubation of SH-SY5Y cells with OA leads to the expression of a high molecular weight phospho-protein species immunoreactive to tau antibodies against phosphorylated Ser202 and phosphorylated Ser396.

#### **4. Materials and Methods**

#### *4.1. Cell culture*

Cells SH-SY5Y (ECACC, 94030304) were grown in Dulbecco's modified Eagle medium (DMEM, Gibco, Gaithersburg, MD, USA, cat. no. 31885-049) supplemented with 10% fetal bovine serum (FBS, Gibco, cat. no. 10270106), 1% L-glutamine (Gibco, cat. no. 25030024), 1% non-essential amino acids (Sigma-Aldrich, Darmstadt, Germany, cat. no. M7145) and 1% penicillin-streptomycin (Gibco, cat. no. 15140-122) at 37°C in humidified atmosphere with 5% CO2. Unless indicated otherwise, undifferentiated SH-SY5Y cells grown to 60-90% confluency were used. For differentiation into neuron-like type, we followed the protocol described by Encinas and colleagues [26] with minor modifications. Briefly, cells were seeded at density of 10,000 cells/cm<sup>2</sup> in the medium described above. The following day a medium containing  $10 \mu$ M all-trans retinoic acid was added to the cells and incubated for five days, the medium being replaced every other day. Cells were washed with serum-free medium and incubated in medium containing 1% FBS and 50 ng/ml brain-derived neurotrophic factor (BDNF, Sigma-Aldrich, cat. no. SRP3014) for two days. Cells were photographed using phase contrast microscopy (Zeiss, Oberkochen, Germany). Where indicated, cells were

treated with indicated concentration of OA (Abcam, Cambridge, UK, cat. no. ab120375) added from 100  $\mu$ M OA solution in dimethyl sulfoxide (DMSO). As a negative control, an equal amount of DMSO to the one used in OA solution was applied to the cells.

#### *4.2. Testing cell viability by MTT assay*

Cell viability was tested using MTT assay. In short, cells were grown in 96-well plates containing 100  $\mu$ l cell culture medium and treated with 30 nM OA, added from a 100  $\mu$ M OA solution in DMSO. For control, an equal amount of DMSO as used in OA solution was added to cells. After 8 h, 40 µl of MTT (3-(4,5-dimethylthiazol-2-yl)-2,5-diphenyltetrazolium bromide) was added to each well and samples were incubated at 37°C for 3 h, followed by the addition of 160 µl of DMSO. Absorbance at 570 nM was measured using BioRad XR680 microplate reader (BioRad Laboratories, Hercules, CA, USA).

#### *4.3. Preparation of cell lysates and protein extraction*

Upon harvesting, cells were washed in ice-cold Tris-buffered saline (TBS) buffer (20 mM Tris-HCl pH 7.4, 150 mM NaCl) and resuspended in Laemmli buffer (50 mM Tris-HCl pH 6.8, 2% sodium dodecyl sulfate [SDS], 2 mM ethylenediaminetetraacetic acid [EDTA], 10% glycerol, 1 "Complete Mini" protease inhibitor cocktail tablet per 10 ml [Roche Diagnostics, Basel, Switzerland cat. no. 11836170001, Roche Diagnostics, Basel, Switzerland], 1 mM phenylmethylsulfonyl fluoride [added from a 100x concentrated solution in DMSO], 10 mM sodium fluoride, 20 mM  $\beta$ -glycerophosphate, and 2 mM sodium orthovanadate). For experiments shown in **Figures 1E**, **2B**, and **3A** cells were resuspended in 2x Laemmli buffer without prior washing step.  $\beta$ -mercaptoethanol (2-5%) was present in the lysis buffer and samples were denatured at 90°C or 95°C for 5-15 min. For the experiment shown in **Figures 1A**, **1B**, **3D**, and **S1** cells were resuspended in ice-cold radioimmunoprecipitation assay (RIPA) buffer (20 mM Tris-HCl pH 7.4, 1% Triton X-100, 0.1% sodium dodecyl sulfate, 0.5% sodium-deoxycholate, 150 mM NaCl, 1 mM EDTA, 10% glycerol, protease and phosphatase inhibitors as described above), drawn through 23 G needle 8 times and incubated on ice for 30 min. Samples were centrifuged at 20,000 g for 10 min at 4°C, supernatant was collected, mixed with Laemmli buffer containing  $5\%$   $\beta$ -mercaptoethanol and incubated at 95°C for 5 min. For the experiment in **Fig. S1**, prior to adding Laemmli buffer, samples were incubated at 85°C for 10 min and centrifuged at 20,000 g for 10 min. For urea treatment, cells

were resuspended in 2x Laemmli buffer containing 8 M urea, samples were incubated at 65°C for 10 min, centrifuged at 20,000 g for 20 min at room temperature (RT) and the supernatant was collected.

#### *4.4. Cell extraction by guanidine-hydrochloride*

Cells harvested by trypsinization were washed in ice-cold phosphate-buffered saline (PBS) and resuspended in a solution containing 6 M guanidine-hydrochloride, 20 mM Tris-HCl pH 7.4 and 100 mM NaCl, incubated at RT for 10 min. Suspension was drawn through 23 G needle and incubated at RT for 20 min, followed by centrifugation at 20,000 g for 5 min at 4°C (**Fig. S3B**) or 20 min at RT (**Fig. 5B**). The supernatant was collected and the centrifugation step repeated. Guanidine hydrochloride forms a precipitate with SDS and is thus incompatible with SDS-PAGE [27]. To remove guanidine hydrochloride, proteins from the supernatant were precipitated by incubation in 90% ethanol at -20°C for 1 h. The samples were centrifuged at 20,000 g for 10 or 20 min at 4°C, the pellet was washed in absolute ethanol and air-dried. For the experiment shown in **Fig. 5B**, precipitated proteins were solubilized in 2x Laemmli buffer (100 mM Tris-HCl pH 6.8, 4% SDS, 4 mM EDTA, 10% glycerol, 1 Roche "Complete Mini" protease inhibitor cocktail tablet per 10 ml, 1 mM phenylmethylsulfonyl fluoride, 10 mM sodium fluoride, 20 mM  $\beta$ -glycerophosphate and 2 mM sodium orthovanadate) containing 8 M urea, incubated at 37°C for 20 min, and centrifuged at 20,000 g for 20 min at RT.  $\beta$ -mercaptoethanol was added to supernatant to a final concentration of 5 % and the centrifugation step repeated. For the experiment shown in **Fig. S3B**, precipitated proteins were solubilized in a volume of RIPA buffer, followed by the addition of a third of the volume of concentrated Laemmli buffer containing  $5\%$   $\beta$ mercaptoethanol, incubation at 95°C for 5 min, centrifugation at 20,000 g at RT and the collection of the supernatant.

#### *4.5. Heat-stable protein fraction*

To prepare heat stable fraction, we followed the protocol described by Petry *et al.* with some modifications [28]. Briefly, the cells harvested by Versene (Gibco, cat. no. 15040-066) were resuspended in 300  $\mu$ l lysis buffer (50 mM Tris-HCl pH 7.4, 1% Triton X-100, 0.25 % sodium-deoxycholate, 150 mM NaCl, 1 mM EDTA, 10% glycerol, 1 Roche "Complete Mini" protease inhibitor cocktail tablet per 10 ml, 1 mM phenylmethylsulfonyl fluoride, 10 mM

sodium fluoride, 20 mM  $\beta$ -glycerophosphate, and 2 mM sodium orthovanadate), incubated on ice for 10 min, drawn through 23 G needle eight times, incubated on ice for 20 min and centrifuged at 20,000 g for 20 min at 4°C. The supernatant was collected and incubated at 95°C for 10 min, followed by centrifugation at 20,000 g for 10 min at 4°C. Supernatant containing the heat-stable fraction (protein concentration of  $0.4 - 0.5 \mu g/\mu l$ ) was separated from the pellet. A volume of the heat-stable fraction  $(150 - 200 \mu l)$  was incubated with one third of the volume of concentrated Laemmli buffer (8% SDS, 200 mM Tris-HCl pH 6.8, 8 mM EDTA, 20% glycerol). The amount of the heat stable protein samples loaded on gel was 12 g of protein. The pellet was resuspended in a similar volume of radioimmunoprecipitation assay (RIPA) buffer and concentrated Laemmli buffer.

#### *4.6. Alkaline phosphatase treatment of protein lysates*

The cells were treated with 100 nM OA for 2 h, trypsinized, washed in PBS buffer and resuspended in lysis buffer (50 mM Tris-HCl pH 7.9, 1% Triton X-100, 0.25% sodiumdeoxycholate, 150 mM NaCl, 1 mM MgCl2, 1 Roche "Complete Mini" protease inhibitor cocktail tablet per 10 ml, 1 mM phenylmethylsulfonyl fluoride, 10 mM sodium fluoride, 20  $mM$   $\beta$ -glycerophosphate), incubated on ice for 10 min, drawn through 23 G needle eight times, incubated on ice for 20 min followed by 5 min centrifugation at 20,000 g at 4°C and the supernatant (protein concentration of 0.4  $\mu$ g/ $\mu$ l) was collected. To enrich heat stable proteins, the sample was incubated at 95°C for 10 min, followed by centrifugation at 20,000 g for 20 min at 4 $\degree$ C. The supernatant (10 µl) was incubated with 10 U of alkaline phosphatase (Promega, Fitchburg, WI, USA, cat. no. M2825) or water (control) at 37°C for 30 min, followed by addition of Laemmli buffer and 5 min incubation at 95°C. The whole reaction mixture was loaded on gel for analysis by immunoblot.

#### *4.7. Immunoblotting*

Proteins in cell extracts were separated on 7.5% (**Figures 1E**, **3C**, and **6**) or 10% (all other figures) SDS-PAGE gels and transferred onto a nitrocellulose membrane in the transfer buffer (25 mM Tris, 192 mM glycine), using the voltage of 50 V for the duration of 2 h. The membrane was blocked in 5% milk diluted in TBS or phosphate-buffered saline (PBS; 137 mM NaCl, 2.7 mM KCl, 10 mM Na<sub>2</sub>HPO<sub>4</sub>, 1.8 mM KH<sub>2</sub>PO<sub>4</sub>) and incubated with primary

antibody diluted in TBS or PBS containing 5% milk or 3% bovine serum albumin, overnight at 4°C. The primary antibodies and their working dilutions used were: anti-tau-pS396 (Abcam, cat. no. ab109390, 1:10,000 dilution), Tau46 (Sigma-Aldrich, cat. no. T9450, dilution 1:1,000), anti-tau-pS202 (CP13, mouse monoclonal, gift of Prof. Peter Davies, dilution 1:500), anti-total tau mouse monoclonal antibody CP27 (gift of Prof. Peter Davies, dilution 1:250), Tau13 (Santa Cruz Biotechnology, Santa Cruz, CA, USA, cat. no. sc-21796, dilution 1:250), polyclonal anti-tau (DAKO-Agilent, CA, USA, cat. no. A-0024, dilution 1:10000), anti-MAP2 (clone HM-2, Sigma-Aldrich, cat. no. M4403, dilution 1:1000), DC11 (generous gift from dr. Michal Novak, dilution 1:50), anti- $\beta$ -actin (Sigma-Aldrich, cat. no. A5441, dilution 1:10,000), and anti-GAPDH (glyceraldehyde 3-phosphate dehydrogenase, Cell Signaling, Leiden, Netherlands, cat. no. 2118, dilution 1:1,000 to 1:2,000). The secondary antibodies were horse radish peroxidase-linked IgG (Cell Signaling, cat. no. 7074 and 7076). Proteins were visualized using Super Signal West substrate (Thermo Scientific, Waltham, MA, USA, Pico cat. no. 34077 or Femto cat. no. 34095) and imager ChemiDoc XRS+ with ImageLab software (BioRad Laboratories, Hercules, CA, USA). Exposure times were selected to maximize the signal to noise ratio. In the **Fig. 1E**, where the signal of samples present on two separate pieces of membrane is compared, membranes were imaged simultaneously. In **Fig. 4**, heat-stable fraction and pellet samples were present and analyzed on the same membrane and imaged simultaneously. Where appropriate, equal loading of samples on gel was assessed using GAPDH levels. Figures are showing representative images of data that were generated from at least three independent samples, or two independent experiments performed on different days, each containing two independent samples generated from separate cell culture dishes, treatment, harvesting and protein preparation. Where appropriate, data were analyzed using ImageLab software (BioRad Laboratories, Hercules, CA, USA).

#### **2. Results**

*2.1. OA treatment of SH-SY5Y cells induced expression of 100 kDa proteins immunoreactive with anti-tau-pS202 and anti-tau-pS396 antibodies* 

To inhibit the activity of protein phosphatases, we incubated undifferentiated neuroblastoma SH-SY5Y cells that endogenously express tau with OA. Taking into consideration the OA

concentration necessary to inhibit the activity of different tau protein phosphatases and possible high local concentration of phosphatases in the cultures [18], we initially treated the cells with 100 nM OA (**Fig. 1A** and **1B**). A similar concentration had been used in previous studies [29-31]. Proteins from total cell protein lysates were separated by denaturing SDS polyacrylamide gel electrophoresis (SDS-PAGE). Tau phosphorylation at specific epitopes was assessed by immunoblot using phospho-specific tau antibodies. Using the antibody that recognizes tau phosphorylated at S396, we observed protein bands of the apparent molecular weight of 50-65 kDa in samples from both treated (OA+) and untreated (OA-) cells (**Fig. 1A**). Additionally, a protein co-migrating with the molecular weight of  $\sim$ 100 kDa appeared in samples from cells treated with OA. The same samples were analyzed with the antibody CP13 that recognizes tau phosphorylated at S202 (**Fig. 1B**). In cells treated with OA a protein with an apparent molecular weight of  $~100$  kDa was clearly visible, while in samples from untreated cells the signal was very weak or not detectable, indicating that under normal growth conditions endogenous tau is phosphorylated at S202 at low levels. The 100 kDa phospho-tau immunoreactive proteins were also induced upon cell treatment with lower OA concentrations of 30 nM OA for 8 h (**Fig. 1C**) and with 20 nM OA for 4 h (**Fig. S1**). Notably, MTT assay on cells treated with 30 nM OA for 8 h revealed that the viability of OA-treated cells was similar to control cells (**Fig. 1D**), suggesting that the appearance of 100 kDa protein is not due to OA-induced cell toxicity. Furthermore, a similar pattern of OA-induced 100 kDa phospho-tau immunoreactive protein was observed in SH-SY5Y cells differentiated into neuron-like cells (**Fig. 2**). Together, the data indicated that protein phosphatase inhibition by OA induced the formation of 100 kDa protein species immunoreactive to tau-pS202 and taupS396. Based on the molecular size of the reported protein and its immunoreactivity with anti-tau antibodies, this result is consistent with the possibility of tau oligomerization.

This high molecular weight tau-immunoreactive protein species (HMW-TIPs) detected with CP13 and anti-tau-S396 antibodies migrated in the SDS-PAGE with nearly identical electrophoretic mobility. To test whether both CP13 and anti-tau-pS396 recognized the same protein, we divided the lane containing protein lysate from OA-treated cells on the nitrocellulose membrane in half by cutting it vertically. We incubated one half with antibody CP13 and the other half with antibody against tau-pS396 and aligned the halves for imaging (**Fig. 1E**). We noticed that the protein band recognized by the CP13 antibody migrated slightly slower than the anti-tau-pS396 antibody-reactive band (**Fig. 1E**), indicating two distinct protein species. These distinct protein bands could represent an identical tau splicing

isoform containing different phosphorylation patterns or an additional posttranslational modification, such as protein truncation.

Next, we examined HMW-TIPs using anti-total tau antibodies that recognize tau regardless of its phosphorylation state. We analyzed OA-treated and -untreated samples using antibody Tau46 that recognizes an epitope in the C-terminal part of tau (aa 404-441). We observed that antibody Tau46 does not detect the 100 kDa protein, although it recognizes 50 kDa tau and a protein around 70 kDa (**Fig. 3A and Fig. 2B**), which presumably represents a low molecular weight MAP2 isoform [32]. To test whether the 70 kDa protein recognized by antibody Tau46 is a MAP2 isoform, we analyzed OA-treated samples with an anti-MAP2 antibody that detects all MAP2 isoforms (**Fig. S2**). In agreement with a previous report showing that SH-SY5Y cells primarily express a 70 kDa MAP2 isoform [33], the anti-MAP2 antibody recognized a protein of this size, similarly to Tau46. The 70 kDa band was clearly distinct from the 100 kDa protein visualized with antibody CP13 (**Fig. S2**, left panel).

Next, we used anti-total tau antibodies raised against other tau regions, Tau13 (epitope most likely within the first 35 aa), CP27 (whose epitope includes residues 130-150), and a rabbit polyclonal antibody raised against the C-terminal part of tau (aa 243-441). Despite the recognition of monomeric tau proteins, Tau13, CP27 and this polyclonal antibody did not detect HMW-TIPs (**Fig. 3B-D**). We further tested the reactivity of HMW-TIPs with antibody DC11 that has been reported to recognize truncated tau (aa 151-421) [34]. However, the pattern of DC11 antibody did not overlap with that of HMW-TIPs (**Fig. 3E**).

Taken together, we were unable to detect HMW-TIPs using several anti-total tau antibodies, which together cover tau epitopes in the N-terminal, central and C-terminal regions of tau, and additionally an antibody against a truncated version of tau. The lack of HMW-TIPs detection using these antibodies could be due to antigen masking within the oligomer or a combination of epitope masking and a different tau protein truncation.

#### *2.2. HMW-TIPs are found in a heat-stable fraction*

To investigate the properties of HMW-TIPs, we examined whether it could be found in a heat-stable fraction. Due to their internally disordered structure, tau and other MAP2 family proteins are resistant to heat-induced precipitation. After isolating a heat-stable fraction using the protocol described by Petry and collaborators [28], in which cells are lysed in a buffer containing 1% non-ionic detergent Triton X-100, 0.25% ionic detergent sodium-deoxycholate

and 150 mM NaCl, but no SDS, the cell lysate was boiled and insoluble proteins were precipitated by centrifugation at 20,000 g. As expected, antibody Tau46 detected 50 kDa tau and a 70 kDa protein, presumably a MAP2 isoform [32] in the heat-stable fraction, and no signal was observed in the pellet fraction (**Fig. 4A**, upper panels). In contrast, the levels of  $\beta$ actin were much lower in the heat-stable fraction than in the pellet (**Fig. 4A**, lower panels). Together, this indicated that the heat-stable fraction was enriched in heat-stable proteins. Using the antibodies anti-tau-pS396 and CP13, we found that HMW-TIPs were present in the heat-stable fraction (**Fig. 4B**). In conclusion, HMW-TIPs co-fractionated with heat-stable Tau46-reactive proteins, which is a characteristic of proteins with internally disordered structure, including tau.

#### *2.3. HMW-TIPs are stable under reducing and strong denaturing conditions*

Considering the possibility that HMW-TIPs represent tau oligomers, we tested conditions under which oligomers could be dissociated. Previous reports have shown that tau oligomers can be disrupted under reducing conditions [35,36], although tau crosslinking by disulfide bonds is not an absolute prerequisite for tau aggregation. Cell lysates in all the experiments were prepared under reducing conditions, therefore this indicated that the stability of the putative oligomer does not depend on disulfide bonds. Next, we examined whether putative oligomers could be dissociated by strong denaturing agent urea and guanidine hydrochloride. Immunoblot analysis of cell lysates using CP13 antibody revealed that HMW-TIPs did not dissociate in the presence of either 8 M urea (**Fig. 5A**) or 6 M guanidine hydrochloride (**Fig. 5B**), and a similar result was obtained using anti-tau-pS396 antibody (**Fig. S3**). Quantification of HMW-TIPs signal relative to GAPDH levels in samples prepared with and without 8 M urea revealed that there was no significant difference between anti-tau-pS396-reactive HMW-TIPs levels in samples from OA-treated cells extracted with and without urea (**Fig. S3A**), while the levels of CP13-immunoreactive HMW-TIPs were higher in cell lysates prepared with urea than without urea, presumably due to a better extraction in the presence of urea (**Fig. 5A**).

#### *2.4. Characterization of the HMW-TIPs species upon protein dephosphorylation*

Finally, to examine the impact of protein phosphorylation on the putative 100 kDa tau oligomer, lysates of cells treated with OA were incubated with alkaline phosphatase. Because

none of the tested anti-total tau antibodies recognized HMW-TIPs, this presented an obstacle in examining proteins upon dephosphorylation. Surprisingly, we observed that alkaline phosphatase treatment did not abolish detection of 55-60 kDa tau and HMW-TIPs by anti-taupS396 antibody (**Fig. 6**), although their electrophoretic mobility was slightly modified (**Fig. 6**, compare molecular weight of proteins in the samples treated and not treated with alkaline phosphatase). The change in electrophoretic mobility indicates that some amino acid residues were dephosphorylated, but not the S396 epitope. Under similar conditions of alkaline phosphatase treatment, CP13-dependent signal of HMW-TIPs was diminished (**Fig. S4**), indicating efficient dephosphorylation at the tau-S202 epitope. The fact that alkaline phosphatase treatment affects the migration rate of HMW-TIPs only to a small degree suggested that the putative oligomer does not dissociate into monomers upon partial protein dephosphorylation.

#### **3. Discussion**

We used SH-SY5Y cell culture model of protein phosphatase downregulation based on cell treatment with OA and showed that OA leads to the appearance of 100 kDa protein species immunoreactive to phospho-tau specific antibodies CP13 (anti-tau-pS202) and anti-taupS396.

Based on its molecular size, we initially considered the possibility that HMW-TIPs represent large tau splicing isoform containing exon 4A (so-called "big tau"), which contains an additional 254 aa insert in the N-terminal part [37] and is normally expressed predominantly in the peripheral nervous tissue [38]. This would be in accordance with a previous report showing that SH-SY5Y cells and their parental line SK-N-SH express a tau isoform around 100 kDa [16]. However, we were unable to detect HMW-TIPs using any of the antibodies used against total tau, namely Tau46, CP27, Tau13, and a rabbit polyclonal antibody. As "big tau" normally contains all of the epitopes recognized by these antibodies, with the possible exception of CP27, and these epitopes are not recognized by the anti-total tau antibodies we used, it is highly unlikely that HMW-TIPs represent "big tau".

Based on the observation that the molecular weight of the HMW-TIPs corresponds to the predicted size of two molecules of fetal tau, which is one of the most prominent tau isoforms in undifferentiated SH-SY5Y cells [17], we considered the possibility that it represents tau oligomers. In support of this possibility, a study using bimolecular fluorescence complementation showed that HEK293 cells expressing tau constructs fused to split Venus increase in fluorescence upon treatment of cells with 30 nM OA for 24 h, suggesting tau oligomerization [39]. The absence of oligomer detection by anti-total tau antibodies in our experiments could be a consequence of by epitope masking or a combination of epitope masking and protein truncation, which would render HMW-TIPs unrecognizable.

In attempt to dissociate putative oligomers by reducing and denaturing agents, we found that HMW-TIPs were stable in the presence of  $\beta$ -mercaptoethanol, SDS, urea and guanidine. The role of disulfide bridges in tau oligomerization is not entirely clear, as both reductionsensitive and reduction-resistant tau dimers have been previously observed [35, 36]. In our experiments, HMW-TIPs were detected in the presence of  $\beta$ -mercaptoethanol, indicating that the stability of a putative oligomer does not depend on disulfide bonds. Despite the fact that most protein complexes dissociate in the presence of SDS, some protein complexes remain assembled [40]. For example, SDS-resistant tau oligomers have been reported in AD brain [35]. Moreover, it has been previously reported that a pool of tau isolated from AD brain remained aggregated in high molecular mass structures even after the treatment with 8 M urea [41], whereas certain types of tau aggregates were stable in 6 M guanidine hydrochloride [42]. Taken together, we were unable to dissociate putative oligomers with SDS, urea, and guanidine. However, the observed stability of HMW-TIPs in the presence of these agents is compatible with the characteristics of some previously reported tau oligomers.

The present study showed that treatment of cell lysates with alkaline phosphatase resulted in partially dephosphorylated HMW-TIPs, leaving the S396 epitope phosphorylated and the putative oligomer assembled. In fact, an inefficient removal of phosphates by alkaline phosphatase from the tau PHF-1 epitope, which includes pS396 and pS404, has been already reported [43]. Our data suggest that exhaustive tau phosphorylation is either not essential for the stability of the putative oligomer, or is necessary for initial oligomer formation, but once formed, oligomers could be stable even upon dephosphorylation. Based on our present data it is also possible that, in addition to other sites, dephosphorylation of S396 is required for the disassembly of the putative oligomer.

Finally, OA affects several cellular processes [45] and inhibition of protein phosphatases PP1, PP2A, PP4 and PP5 by OA is lethal to most human cells [18]. To minimize the toxic effect of OA on cells, we used treatment with OA concentrations of 100-150 nM for a short period of 2-2.5 h. Moreover, we have also observed the induction of the 100 kDa anti-tau reactive protein using a lower OA concentration of 20 nM for 4 h and upon cell treatment with 30 nM OA for 8 h, which did not affect cell viability, suggesting that the appearance of the 100 kDa protein is not due to OA-induced cell toxicity. However, we cannot entirely exclude the possibility that OA-induced HMW-TIPs is not a direct consequence of tau-immunoreactive protein phosphorylation, but could arise due to an indirect effect of phosphatase inhibition by OA [44], or through a phosphatase-independent pathway [45].

In summary, OA treatment of SH-SY5Y cells lead to the appearance of HMW proteins immunoreactive to anti-tau-pS202 and -pS396 antibodies, which is in line with the possibility that HMW-TIPs represent tau oligomers. However, due to the inability to detect HMW-TIPs with anti-total tau antibodies and to dissociate putative oligomers under conditions tested, we cannot exclude the possibility that HMW-TIPs represent tau-unrelated proteins that are detected with anti-tau-pS202 and -pS396 antibodies due to cross-reactivity. The finding that anti-tau-pS396- and -pS202-immunoreactive HMW-TIPs exhibited almost identical, yet distinct electrophoretic mobilities in SDS-PAGE supports this possibility. Furthermore,

although HMW-TIPs were present in the heat-stable fraction, which is a characteristic of tau and other intrinsically disordered proteins [46], it should be noted that due to the presence of 1% Triton X-100 and 0.25% sodium-deoxycholate in the lysis buffer, our protocol may have extracted a broader range of proteins than buffers without detergents.

In conclusion, we provide strong evidence that OA treatment of neuroblastoma cells in culture induces a high molecular weight phospho-tau immunoreactive species. Although additional research is required to clarify further the identity of the reported proteins, our findings indicate that OA-treated SH-SY5Y cells represent a potentially valuable cell culture model for studying tau oligomerization under the conditions of protein phosphatase downregulation, as well as tau aggregation inhibitors and neuroprotective compounds [47-49]. Our findings may thus facilitate future studies aiming to explain molecular mechanisms involved in the early events leading to tau aggregation and neurodegeneration, which may eventually lead to effective therapy and prevention of neurofibrillary degeneration and AD.

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**Author Contributions. Declaration of authorship**. GŠ conceived and directed the project, coordinated experiments and co-wrote the paper. MB designed, performed and interpreted experiments and co-wrote the paper. MBL established the experimental model, performed and interpreted initial experiments and contributed with the data for the revision. TM performed and analyzed experiments under supervision of MB. PRH substantially contributed to the

interpretation of data for the work. All authors contributed to revising and editing the manuscript critically for important intellectual content. All authors approved the final version of the manuscript.

**Competing interests**. All authors have completed the Unified Competing Interest form at www.icmje.org/coi\_disclosure.pdf (available on request from the corresponding author) and declare no financial relationships with any organizations that might have an interest in the submitted work in the previous three years. All authors declare no other relationships or activities that could appear to have influenced the submitted work.

### **Abbreviations**



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#### **FIGURES AND FIGURES LEGENDS**

**Figure 1. OA treatment of SH-SY5Y cells induces expression of a high molecular weight protein species immunoreactive to phospho-tau**. Undifferentiated SH-SY5Y cells were treated with indicated concentrations of OA for an indicated time period. Around  $25 \mu g$ protein in total cells lysates were analyzed by immunoblotting using antibodies that detect tau-pS396 or tau-pS202 (CP13). GAPDH served as a loading control (**A-C**). Note a strong band migrating around 100-kDa in lysates of OA-treated cells (**A-C**). (**D**) Viability of cells treated with 30 nM OA for 8 h (number of independent samples  $n = 8$ ) and control cells ( $n =$ 4) was tested using MTT assay. Percent viability of control cells is shown. No significant difference in viability was found ( $t = 0.372$ ,  $df = 10$ ,  $p = 0.125$ ). (**E**) Nitrocellulose membrane with transferred proteins was cut vertically in the middle of the lane (scissors) and the halves were incubated with CP13 or anti-tau-pS396 antibodies. Membranes were placed side-by-side and imaged simultaneously.

**Figure 2. HMW-TIP is detected in OA-treated SH-SY5Y cells differentiated into neuron-like cells.** Differentiated SH-SY5Y cells visualized using phase contrast microscopy (**A**) were treated with 100 nM OA for two hours. Note the differentiated cells with visible neuritic processes. Scale bar = 50  $\mu$ m. Total cell lysates were analyzed by immunoblot using anti-tau-pS202 (CP13), anti-tau-pS396 and anti-total tau (Tau46) antibodies (**B**). Note a strong band migrating around 100-kDa in lysates of OA-treated cells. GAPDH is used as a loading control.

**Figure 3. HMW-TIPs are not detected by anti-total tau antibodies.** Where indicated, cells were treated with 100 nM (**A**, **D**) or 150 nM (**B**, **C**, **E**) OA for 2 - 2.5 hours. Total cell lysates were analyzed by immunoblotting using anti-total tau antibodies Tau46 (**A**), CP27 (**B**), Tau13 (**C**) and rabbit polyclonal antibody (**D**), and antibody DC11 against truncated tau (**E**). To visualize HMW-TIPs, same samples were analyzed by antibody CP13 (**A-C**) or anti-taupS396 (**D**, **E**). Note a CP13- and anti-tau-pS396- reactive band migrating around 100-kDa in lysates of OA-treated cells (**A-E**). Note that all anti-total antibodies detect a band migrating above 50 kDa that represents monomeric tau, but do not detect 100 kDa protein (**A-D**). Tau46 antibody additionally detects a 70 kDa band that presumably represents MAP2 isoform (**A**). GAPDH served as a loading control (**C**, **D**). Control and OA-treated samples in (**D**) were present on the same membrane. Note that the rabbit polyclonal anti-total antibody (right panel) detects a band migrating above 100 kDa tau (red arrow), which is clearly distinct from the OA-induced anti-tau-pS396-reactive band (left panel, OA+) migrating at 100 kDa (black arrow).

**Figure 4. HMW-TIP is present in a heat-stable fraction.** SH-SY5Y cells were treated with 100 nM (OA+) or DMSO (OA-) for 2:15 h. (**A**) Heat-stable (HS) and pellet fractions were analyzed by immunoblot using anti-total tau antibody Tau46 (upper panels) and anti- $\beta$ -actin (lower panels). Heat-stable fraction and pellet samples were analyzed on the same nitrocellulose membrane and imaged simultaneously. Note that 50 kDa tau and 70 kDa proteins (70 kDa protein presumably being a MAP2 isoform) detected by Tau46 antibody are enriched in the heat-stable fraction compared to the pellet. In contrast,  $\beta$ -actin is enriched in the pellet compared to heat-stable fraction. (**B**) The same samples of heat-stable (HS) fraction as in (**A**) were analyzed by immunoblot using anti-tau-pS396 and CP13 (anti-tau-pS202) antibodies. Note the presence of pS396- and CP13-immunoreactive 100 kDa bands in the heat stable fraction.

**Figure 5. HMW-TIP is stable under denaturing conditions.** Cells were treated with 150 nM (**A**) or 100 nM (**B**) OA for 2 h. Cell lysates were prepared in the absence or presence of 8 M urea (**A**) or with 6 M guanidine-hydrochloride (**B**). Immunoblotting was performed using anti-tau-pS202 antibody (CP13). GAPDH was used as a loading control. Note that the 100 kDa band is present in samples from OA-treated cells under all tested conditions. The graph shows signal intensity of HMW-TIP relative to GAPDH levels of the samples shown in **A**. Average values and standard deviation of replicate samples are shown.

**Figure 6. HMW-TIPs stability upon protein dephosphorylation.** Cells were incubated with 100 nM OA for 2 h. Heat-stable fraction was treated with alkaline phosphatase (AP) where indicated and analyzed by immunoblot using anti-tau-pS396 antibody. Note the persistent antibody detection of protein migrating around 100 kDa and changed electrophoretic mobility of proteins treated with alkaline phosphatase.

**Figure S1. Treatment of SH-SY5Y cells with low concentration of OA induces expression of 100 kDa tau-reactive protein.** Cells were treated with 20 nM OA for 4 h. Samples were analyzed by immunoblotting using antibodies that detect tau-pS202 (CP13 antibody, upper panel) and tau-pS396 (lower panel). Note CP13- and pS396-immunoreactive bands (arrows) migrating around 100-kDa in lysates of OA-treated cells.

**Figure S2. Anti-MAP antibody detects 70 kDa protein reactive to Tau46.** Cells were treated with 150 nM OA for 2 h and total cell lysates were analyzed by immunoblot using antibodies CP13, anti-MAP2 and Tau46. GAPDH was used as a loading control. Arrows mark the 100 kDa tau-reactive protein. Asterisk marks anti-MAP2- and Tau46 immunoreactive protein migrating at 70 kDa.

**Figure S3. Anti-tau-pS396-immunoreactive HMW-TIP is stable under denaturing conditions.** Cells were treated with 150 nM (**A**) or 100 nM (**B**) OA for 2 h. Cell lysates were prepared in the absence or presence of 8 M urea (**A**) or using lysis buffer containing 6 M guanidine-hydrochloride (**B**). Immunoblotting was performed using anti-tau-pS396 antibody. GAPDH was used as a loading control. Note that the 100 kDa band is present in samples from OA-treated cells under all tested conditions. The graph shows signal intensity of anti-taupS396 reactive HMW-TIP relative to GAPDH levels of the samples shown in **A**. Average values and standard deviation of replicate samples are shown. Note that there is no significant difference between "+OA -urea" and "+OA +urea" conditions ( $p = 0.34$ ).

**Figure S4. CP13-dependent HMW-TIPs signal diminishes upon protein dephosphorylation by alkaline phosphatase.** Cells were incubated with 100 nM OA for 2 h. Heat-stable fraction was treated with alkaline phosphatase (AP) and analyzed by immunoblot using CP13 antibody. Note the 100 kDa band representing HMW-TIPs and the disappearance of the signal upon alkaline phosphatase treatment.



 $\overline{\mathbf{A}}$ 











B













 $-AP$   $+ AP$ 



taupS396









 $\blacksquare$ 



+ Gu

