Research Article

Tau and Caspase 3 as Targets for Neuroprotection

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The peptide drug candidate NAP (davunetide) has demonstrated protective effects in various in vivo and in vitro models of neurodegeneration. NAP was shown to reduce tau hyperphosphorylation as well as to prevent caspase-3 activation and cytochrome-3 release from mitochondria, both characteristic of apoptotic cell death. Recent studies suggest that caspases may play a role in tau pathology. The purpose of this study was to evaluate the effect of NAP on tau hyperphosphorylation and caspase activity in the same biological system. Our experimental setup used primary neuronal cultures subjected to oxygen-glucose deprivation (OGD), with and without NAP or caspase inhibitor. Cell viability was assessed by measuring mitochondrial activity (MTS assay), and immunoblots were used for analyzing protein level. It was shown that apoptosis was responsible for all cell death occurring following ischemia, and NAP treatment showed a concentration-dependent protection from cell death. Ischemia caused an increase in the levels of active caspase-3 and hyperphosphorylated tau, both of which were prevented by either NAP or caspase-inhibitor treatment. Our data suggest that, in this model system, caspase activation may be an upstream event to tau hyperphosphorylation, although additional studies will be required to fully elucidate the cascade of events.

1. Introduction

NAP is an 8 amino acid peptide, which was identified as an active neuroprotective fragment of activity-dependent neuroprotective protein (ADNP) [1]. NAP was found to be neuroprotective in various in vivo and in vitro models of neurodegeneration [2]. NAP treatment was shown to reduce two major pathological markers—tau hyperphosphorylation [3–5] and caspase-3 activation/apoptosis [6, 7]. Hyperphosphorylated and aggregated tau, originally detected in Alzheimer’s disease (AD) brains by Grundke-Iqbal and colleagues [8], is a hallmark of a group of diseases, generally referred to as “tauopathies” which differ from each other by genetic background and by additional pathological and phenotypic characteristics [9].

Tau is a microtubule-associated protein (MAP) which promotes microtubule stabilization. The first study that reported disassembly of microtubules from AD brain due to the abnormal hyperphosphorylation of tau was by Iqbal et al. [10]. Hyperphosphorylated tau loses its microtubule affinity causing a change in microtubule dynamics towards disassembly [11–16] and further accumulation of aggregated tau. Alonso et al. originally showed that AD abnormal hyperphosphorylation of tau causes not only loss of function but also the gain of toxic function, with hyperphosphorylated tau blocking microtubule assembly in the presence of normal tau [17] and promoting the formation of normal tau containing tangles [18]. The most prevalent tauopathy is Alzheimer’s disease (AD), and other tauopathies include a selection of frontotemporal dementia/degeneration, with pure tauopathies like progressive supranuclear palsy (PSP) [19].

One of the hallmarks of tauopathies is the accumulation of neurofibrillary tangles (NFTs). NFT accumulation correlates with the severity of dementia and memory loss [20–22] and with neuronal degeneration in AD and PSP [23, 24]. Recent studies associated the spread of tauopathies with propagation of prion-like protein inclusions [25] and suggested transsynaptic spread of tau [26]. However, several studies showed that soluble defective tau is also correlated
2. Materials and Methods

All procedures involving animals were approved by the Animal Care Committee of Tel-Aviv University and further approved by the Israel Health Authorities.

2.1. Primary Neuronal Cultures. Primary neuronal cultures were produced from cortices obtained from 1-2-day-old Sprague-Dawley rat pups. Pups were decapitated, their skulls were removed, and their cortices were transferred to a Petri dish field with HBSS-HEPES buffer. Meninges were removed using tweezers under binocular. Neurons were produced with Worthington’s papain dissociation kit (Worthington, cat.LK003150) according to the manufacturer’s instructions. The cells were seeded in Neurobasal-A media (Invitrogen, cat.LK003150) according to the manufacturer’s instructions.

2.2. Antibodies. The following antibodies were used for immune-fluorescence: NeuN (Neuronal Nuclei, Chemicon, cat. MAB377), diluted 1:100, and GFAP (glial fibrillary acidic protein, Abcam, cat. 7260), diluted 1:500. The following antibodies were used for immunoblot analysis: p-tau202 (Anaspec, cat. 28017) 0.25 μg/mL, total tau (MBL international, cat. AT-5004) 0.5 μg/mL, and active caspase-3 (Abcam, cat. 2302) 1 μg/mL. Fluorophore and horseradish peroxidase conjugated secondary antibodies were purchased from Jackson ImmunoResearch Laboratories Inc.

2.3. Analysis of Culture Purity. Cells were seeded in 24 well plates on poly-D lysine- (PDL-) coated coverslips (Sigma-Aldrich, cat. p-6407) at a density of 0.5 × 10^5 cells/well. At 5–6 DIV, the cultures were fixed using a 4% paraformaldehyde (PFA) solution. Antigen retrieval, if needed, was performed by boiling the sample in sodium citrate buffer (pH 6) for 20 minutes. Permeabilization was performed with a 0.2% Triton-X solution. Nonspecific antigen binding was blocked by incubation in a 2% bovine serum albumin (BSA) solution, then primary antibody (Ab) diluted in primary antibody diluent (Biotec applications, cat. MSBA-AbDil) was added for 1 hour at RT. Cells were washed three times in 2% solution of BSA and incubated in secondary antibody solution for 30 minutes in the dark. After an additional wash, PBS + DAPI (4′,6-diamidino-2-phenylindole, a fluorescent stain that binds strongly to A–T-rich regions in DNA, Invitrogen) was applied for 5 minutes, in order to stain cell nuclei. The coverslips were put on an antifading buffer on a carrier slides. For analysis of culture purity, we used 3 cultures produced independently. We used Zeiss fluorescence microscope and

Table 1: Models of NAP protection against tau pathology or increased markers of apoptosis.

| Models of tau pathology detecting protective effects when treated with NAP |
|--------------------------------------------------------------------------|
| (i) Transgenic ADNP heterozygous mouse [4]                                |
| (ii) Transgenic human double-mutant tau mouse [5]                        |
| (iii) Triple transgenic mice expressing the amyloid (Aβ) precursor protein APP(Swe), presenilin PS1 (M146V), and tau (P301L) [45,46] |
| (iv) Mixed neuroglial primary cultures treated with Aβ (1–42, 2.5 μM) [3]|
| (v) Primary cultures of astrocytes [47]                                   |

| Models of apoptosis detecting protective effects when treated with NAP    |
|--------------------------------------------------------------------------|
| (i) A stroke model using spontaneously hypertensive rats which underwent permanent middle cerebral artery occlusion [48] |
| (ii) A rat model of diabetes (streptozocin toxicity) [7]                 |
| (iii) A rat model of epilepsy [49]                                        |
| (iv) PC-12 cells exposed to H2O2 [50]                                    |
| (v) Primary neuronal cultures subjected to ischemia/ reperfusion schedule of 3 h/3 h [39] |

with cognitive deficits [27–29], causing synaptic loss and gliosis before NFTs formation [30]. Tau is also a substrate of multiple caspases, which cleave it and promote its pathologic aggregation [31–33]. Cleaved and hyperphosphorylated tau is found in deposits in AD brains, and it was found that truncated tau promotes apoptosis [34]. The relationship between tau hyperphosphorylation and cleavage is not completely understood. Though some evidence suggests that caspase cleavage of tau is not necessary for tau hyperphosphorylation, this question should be further clarified [35,36]. Caspases are known to play a role in AD (for review, see [37]), and caspase effects on modifications of tau are of a great interest. Furthermore, treatment of an AD mouse model with a broad-spectrum caspase inhibitor was later shown to reduce tau pathology but not amyloid-β (Aβ) pathology [38].

To date, we have tested NAP effects on either tau hyperphosphorylation or on caspase activation in separate models, showing inhibition of tau pathology or inhibition of caspase activation (or other apoptotic markers) (Table 1). The data emerging from these studies raise the question whether there is a dependency between the effects of NAP on the two pathologies. In the current study, we used an in vitro model of ischemia in which NAP treatment reduced apoptosis [39]. As animal models of ischemia exhibit tau hyperphosphorylation [40–42], we investigated whether the in vitro ischemia model mimics the in vivo situation. Thus, the aim of the current study was to identify a possible interaction between caspase activity and tau hyperphosphorylation, and to test the effect of NAP on both events under ischemic conditions.
camera to obtain 10 pictures (×20 magnification) from each culture. Each picture contained a minimum of 30 cells. NeuN-positive and GFAP-negative cells were considered neuronal and were expressed as % of the total DAPI count.

2.4. OGD Treatment. Cells were seeded on PDL-coated plates employing two different plating conditions (Corning): (1) 10 cm Petri dishes, 6 × 10^5 cells per dish (for protein extraction), and (2) 48 well plates 0.5 × 10^5 cells/well (for viability tests). After 5-6 DIV, the condition media (CM) was collected, filtered, and kept at 37°C. Cultures were washed twice with 37°C PBS, and the experimental media was added (PBS for the OGD groups, fresh media for the control group). Treatments were added to the experimental media (QVD-OPH, a broad-spectrum caspase inhibitor [43], 20 μM diluted in DMSO (Biovision, cat. no. 1170) or 10^{-5} M NAP diluted in PBS or diluents only). The culture dishes were placed in an ischemic chamber (Billups-Rothenberg, Inc., cat. MIC 101) which was filled with a 5%CO_2/95%N_2 gas mixture and sealed, and the chamber was placed in an incubator for 2 hours; at 37°C. Two experimental modes were applied, in the first one (used for NAP protecting concentration curve), gassing was for 25 minutes followed by sealing for 2 hours, in the second paradigm (used for calibration of caspase inhibition and protein analysis), gassing was for 5 minutes only. O_2 percentage was monitored with O_2 meter (HUMIL International Corporation, cat. PO2-250) and did not exceed 1.2% any time during the OGD period. At the end of the OGD period, the culture dishes were removed from the chamber. For protein extraction, the experimental media was removed and protein was extracted as described below.

2.5. Viability Assay. Cells were seeded in 48 well plates at a density of 0.5 × 10^5 cells/well (as described above). Following the OGD period, the condition media (CM) which was removed at the initiation of the OGD treatment (see above) was reapplied to the corresponding treatments. The MTS assay was used to assess viability (Promega, cat. G3580). Absorbance at 490 nm was measured in a SpectraMAX 190 plate reader, (Molecular Devices, Inc.). The results were analyzed using SoftMax Pro software version/year (Molecular Devices, Inc.) and expressed as % of the control. The results summarize 3 independent experiments with 5 replicates per treatment per experiment. Data are presented as means/medians ± SE.

2.6. Immunoblots. Immediately after OGD treatment (following gassing mode of 5 minutes of gassing and 2 hours OGD period), experimental media was discarded, and the dishes were washed twice with PBS. Protein was extracted using RIPA buffer with anti-proteases (ROCHE, cat. 11 873 580 001) and anti-phosphatases. Protein concentration was quantified using Pierce BCA protein assay reagents (Pierce, cat. 23223, 23224) with bovine serum albumin (BSA) as a standard protein. Equal amounts of protein diluted in sample buffer were separated by sodium dodecyl sulfate (SDS) polyacrylamide gel electrophoresis (12% gels or 4–20% gradient gels, nUVview, cat. NG21-420). Proteins were transferred to nitrocellulose membranes using standard techniques, as before [44]. Membranes were blocked with 5% BSA in Tris-buffered saline containing 0.1% Tween-20 (TBST) and then incubated overnight with primary antibody. Membranes were washed 3 times with TBST and incubated with secondary Ab for 1 hour at RT. Pierce ELC substrate or Pierce super signal ECL was used to visualize proteins (cat. 32106, 34080). Signal density was detected with the DNR bioimaging system/MiniBIS pro (DNR Bio-Imaging Systems Ltd.) and TotalLab TL-100 software (Nonlinear Dynamics Ltd.).

2.7. Statistical Analysis. SPSS 19.0 for windows (http://www-01.ibm.com/software/analytics/spss/) was used for statistical analysis. For viability assessment, 3 independent experiments were conducted with 5 trials per treatment in each experiment. Data were analyzed using a one-way ANOVA with 5% selected the level of significance, and post hoc Scheffe and LSD corrections. The effect of NAP was evaluated in 14 independent experiments, of which in 4 experiments QVD-OPH effect was also tested, and the levels of significance were evaluated using ANOVA with post hoc LSD.

3. Results

3.1. Characterization of Culture Purity. Culture purity was analyzed using immunofluorescent staining with cell-specific markers. After 5-6 DIV (days in vitro), 95.9 ± 0.96% (SEM) of the cells (counted using nuclear DAPI stain) were positive for the neuronal NeuN marker and negative for the astrocytic marker, GFAP. A representative immunofluorescence stain in Figure 1 shows neuronal cells stained with NeuN in red, and astrocytes stained with GFAP in green.

3.2. NAP Protective Effect Following 2 Hours of OGD. Cell viability was evaluated following a 25-minute gassing period and an additional 2 hours of OGD. The results were calculated and expressed as % of the control group (Figure 2). Under this paradigm, ∼70% cell death was observed. Treatment with NAP increased cell viability immediately following OGD (F = 17.667, df = 7, P < 0.001). Post hoc pairwise comparisons (LSD) indicated that NAP concentrations of 10^{-7} M and 10^{-5} M significantly protected against ischemia (Figure 2, P ≤ 0.05).

3.3. Apoptosis Was the Main Type of Cell-Death Following 2 Hours of OGD. Using the broad spectrum caspase inhibitor QVD-OPH, we tested the extent of apoptotic death following 5-minute of gassing and 2 hours of OGD. Cell viability was evaluated and compared to the control group (Figure 3). Under this paradigm, ∼55% cell death was observed. Here, QVD-OPH (2 × 10^{-5} M) treatment significantly increased cell viability to the control levels. In the same experiment, NAP in the same experiment, NAP (10^{-5} M) treatment rescued from the apoptotic cell death (F = 16.090, df = 7, P < 0.001). Post hoc pairwise comparisons did not detect any
Figure 1: Culture purity: at 5-6 DIV, cells were fixed and immunofluorescence staining was performed using the astrocyte marker GFAP (green), the neuronal marker NeuN (red), and DAPI stain (blue) for nuclei. Quantification of neuronal cells was done using 10 random fields from each of 3 experiments (×20 magnification). 95.9 ± 0.96% of the total cells (DAPI stain) were recognized as neurons (NeuN positive, GFAP negative).

significant differences between the caspase inhibitor, NAP, and control no-ischemia groups.

3.4. Active Caspase-3 Expression Was Increased Following OGD and Reduced Due to NAP Treatment. The levels of active caspase-3 were evaluated immediately after the OGD period using immunoblots with a specific active caspase-3 antibody and further quantified and normalized to actin. As shown in a representative blot in Figure 4(a), OGD treatment increased active caspases-3 levels. This increase was prevented by NAP treatment (10⁻⁵ M) or QVD-OPH (caspase inhibitor) treatment (2 × 10⁻⁵ M). Figure 4(b) depicts the level of active caspase-3 expressed as % of control. A significant increase in active caspase-3 expression was induced by OGD and partially but significantly prevented by NAP treatment (F = 7.880, df = 2, P = 0.004).

3.5. Tau Hyperphosphorylation Was Increased Following 2 Hours of OGD; P-Tau Increase Was Prevented by Either NAP Treatment or QVD-OPH Treatment. Phospho-tau (p-tau) levels were detected by immunoblotting, with a p-tau202-specific antibody. A representative blot is shown in Figure 5(a). P-tau levels were quantified, normalized to total tau levels and expressed as % of the p-tau levels in the control culture (Figure 5(b)). Phospho-tau levels were significantly increased following the OGD insult. Both NAP treatment (10⁻⁵ M) and QVD-OPH (caspases inhibitor) treatment (2 × 10⁻⁵ M) prevented p-tau increase (ANOVA with post hoc LSD, P ≤ 0.05), (F = 5.633, df’ = 2, P = 0.012).

4. Discussion

Focusing on the protective properties of NAP, two central characteristics of neurodegeneration have been shown to be inhibited by NAP treatment both in vivo and in vitro: tau hyperphosphorylation [3–5] and apoptosis [6, 7, 51]. The main goal of the current study was to look for an association between these two pathological cascades. Our experimental setup used primary neuronal cultures (95.9 ± 0.96% purity), subjected to 2 hours of OGD with no reperfusion period. Under this paradigm, apoptosis was found to be responsible for all cell death in that it was inhibited by QVD-OPH. Our results showed that inhibition of caspase activity prevented tau hyperphosphorylation, leading us to conclude that in the current experimental
and cellular conditions, caspase activation is an upstream event to tau hyperphosphorylation. We further evaluated the effect of NAP on cell viability, caspase-3 activation, and tau hyperphosphorylation. Viability assays showed that NAP treatment rescued cells from apoptosis as demonstrated by the reduction in active caspases-3 following NAP treatment. NAP treatment also prevented tau hyperphosphorylation after the OGD. Combining the effect of NAP on active caspases-3 and tau hyperphosphorylation, it seems likely that caspases-3 or an upstream pathway is targeted by NAP activity in isolated neuronal cells that are metabolically stressed.

A careful look should be given to the concentrations by which NAP had a significant protective effect. Previous data, including ischemia-reperfusion experiments conducted in primary neuronal and neuroglial cultures, exhibited protection from cytochrome-c release, microtubule breakdown and reduction in MAP2 intensity using NAP concentrations of $10^{-15}$–$10^{-8}$ M [39, 52]. Though most studies used femtomolar concentration of NAP, Pascual and Guerri [53] used concentration of $10^{-7}$ M to show NAP protection in a model neurons cocultured with astrocytes obtained from prenatal ethanol-exposed fetuses. In our model, protection was detected when treating with NAP at concentrations $\geq 10^{-7}$ M. The differences in the potent NAP concentration could be explained by the combination of the acute insult applied to a relatively pure culture of immature neurons. Immature neurons (2–4 DIV) in a pure neuronal culture (99% purity) were shown to go through spontaneous apoptotic death which was prevented when astrocyte condition media was added to the neuronal cultures. In comparison to neuroglial mixed culture, pure neuronal cultures were also found to be more sensitive to excitotoxic insult that also plays a role in the ischemia induced cell death [54]. We have previously shown that neuronal protection in the absence of glia required increased concentrations of NAP as

**Figure 4:** An increase in active caspase-3 levels induced by 2 hours of OGD was diminished by NAP treatment ($10^{-5}$ M). Cultures were treated with $10^{-5}$ M NAP or with $20 \mu$M QVD-OPH (broad spectrum caspase inhibitor) exposed to OGD insult for 2 hours. Proteins were extracted and analyzed using immunoblot with a specific antiactive caspase-3 antibody. A representative blot of active caspase-3 antibody is exhibited in (a). (ANOVA with post hoc LSD, $P \leq 0.05$) (b).

**Figure 5:** 2 hours of OGD caused an increase in p-tau202 levels, prevented by either NAP or QVD-OPH treatment. Cultures were treated with $10^{-5}$ M NAP or with $20 \mu$M QVD-OPH (broad spectrum caspase inhibitor, indicated as OGD + Q) and exposed to OGD insult for 2 hours. Proteins were extracted and analyzed using immunoblot with a specific anti-p-tau202 and antitotal tau antibodies. A representative blot of p-tau202 and total tau is exhibited in (a). P-tau levels were quantified and normalized to total tau levels (ANOVA with post hoc LSD, $P \leq 0.05$) (b).
compared to mixed neuroglial cultures [55]. Furthermore, the original discovery of the NAP containing protein, ADNP, was as a glial protein providing neuroprotection. ADNP is essential for central nervous system development, expressed in specific brain tissue in the adult brain [1], and is also deregulated and may be overexpressed following acute brain damage [56] and in animal models of neurodegeneration [57, 58]. ADNP-like immunoreactivity is secreted from glial cells [59] further providing neuronal protection [4, 53, 60]. In this respect, NAP protected against ADNP deficiencies as outlined in the Introduction [4]. NAP was further shown to promote neurite outgrowth [61, 62] provide microtubule stabilization [63] and protect neurons in a broad spectrum of neuropathologies models [2]. Importantly, NAP also showed glial protection in vitro [64] and in vivo [7].

In summary, we demonstrated that NAP reduces caspase-3 activation, and tau hyperphosphorylation, suggesting that caspase-3 or upstream pathways may be targets for NAP activity. We believe these data provide additional insights regarding the molecular mechanism for NAP’s neuroprotective activity and the intricate interaction between microtubules/tau and apoptotic mechanisms.

Disclosure

Professor I. Gozes serves as the Founder Scientist and Director at Allon Therapeutics Inc., the company that develops davunetide (NAP), (http://www.allontherapeutics.com/).

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References

[1] M. Bassan, R. Zamostiano, A. Davidson et al., “Complete sequence of a novel protein containing a femtomolar-activity-dependent neuroprotective peptide,” Journal of Neurochemistry, vol. 72, no. 3, pp. 1283–1293, 1999.
[2] I. Gozes, B. H. Morimoto, J. Tiong et al., “NAP: research and development of a peptide derived from activity-dependent neuroprotective protein (ADNP),” CNS Drug Reviews, vol. 11, no. 4, pp. 353–368, 2005.
[3] N. Shiryaev, R. Pikman, E. Giladi, and I. Gozes, “Protection against tauopathy by the drug candidates NAP (Davunetide) and D-SAL: biochemical, cellular and behavioral aspects,” Current Pharmaceutical Design, vol. 17, no. 25, pp. 2603–2612, 2011.
[4] I. Vulih-Shultzman, A. Pinhasov, S. Mandel et al., “Activity-dependent neuroprotective protein snippet NAP reduces tau hyperphosphorylation and enhances learning in a novel transgenic mouse model,” Journal of Pharmacology and Experimental Therapeutics, vol. 323, no. 2, pp. 438–449, 2007.
[5] N. Shiryaev, Y. Jouroukhin, E. Giladi et al., “NAP protects memory, increases soluble tau and reduces tau hyperphosphorylation in a tauopathy model,” Neurobiology of Disease, vol. 34, no. 2, pp. 381–388, 2009.
[6] R. R. Leker, A. Teichner, N. Grigoriadis et al., “NAP, a femtomolar-acting peptide, protects the brain against ischemic injury by reducing apoptotic death,” Stroke, vol. 33, no. 4, pp. 1085–1092, 2002.
[7] A Idan-Feldman, Y. Schirer, E. Polyzoidou et al., “Davunetide (NAP) as a preventative treatment for central nervous system complications in a diabetes rat model,” Neurobiology of Disease, vol. 44, no. 3, pp. 327–339, 2011.
[8] I. Grundke-Iqbal, K. Iqbal, and Y. C. Tung, “Abnormal phosphorylation of the microtubule-associated protein r (tau) in Alzheimer cytoskeletal pathology,” Proceedings of the National Academy of Sciences of the United States of America, vol. 83, no. 13, pp. 44913–4917, 1986.
[9] V. M. Y. Lee, M. Goedert, and J. Q. Trojanowski, “Neurodegenerative tauopathies,” Annual Review of Neuroscience, vol. 24, pp. 1121–1159, 1999.
[10] K. Iqbal, I. Grundke-Iqbal, and T. Zaidi, “Defective brain microtubule assembly in Alzheimer’s disease,” The Lancet, vol. 2, no. 8504, pp. 421–426, 1986.
[11] A. Ebner, R. Godemann, K. Stamer et al., “Overexpression of tau protein inhibits kinesin-dependent trafficking of vesicles, mitochondria, and endoplasmic reticulum: implications for Alzheimer’s disease,” Journal of Cell Biology, vol. 143, no. 3, pp. 777–794, 1998.
[12] D. Terwel, I. Dewachter, and F. Van Leuven, “Axonal transport, tau protein, and neurodegeneration in Alzheimer’s disease,” NeuroMolecular Medicine, vol. 2, no. 2, pp. 151–165, 2002.
[13] D. N. Dreichsel, A. A. Hyman, M. H. Cobb, and M. W. Kirschner, “Modulation of the dynamic instability of tubulin assembly by the microtubule-associated protein tau,” Molecular Biology of the Cell, vol. 3, no. 10, pp. 1141–1154, 1992.
[14] G. T. Bramblett, M. Goedert, R. Jakes, S. E. Merrick, J. Q. Trojanowski, and V. M. Y. Lee - , “Abnormal tau phosphorylation at Ser396 in Alzheimer’s disease recapitulates development and contributes to reduced microtubule binding,” Neuron, vol. 10, no. 6, pp. 1089–1099, 1993.
[15] J. Biernat, N. Gustke, G. Drewes, E. M. Mandelkow, and E. Mandelkow, “Phosphorylation of Ser262 strongly reduces binding of tau to microtubules: distinction between PHF-like immunoreactivity and microtubule binding,” Neuron, vol. 11, no. 1, pp. 153–163, 1993.
[16] A. D. C. Alonso, T. Zaidi, M. Novak, I. Grundke-Iqbal, and K. Iqbal, “Hyperphosphorylation induces self-assembly of r into tangles of paired helical filaments/straight filaments,” Proceedings of the National Academy of Sciences of the United States of America, vol. 98, no. 12, pp. 6923–6928, 2001.
[17] A. D. C. Alonso, T. Zaidi, I. Grundke-Iqbal, and K. Iqbal, “Role of abnormally phosphorylated tau in the breakdown of microtubules in Alzheimer’s disease,” Proceedings of the National Academy of Sciences of the United States of America, vol. 91, no. 12, pp. 5562–5566, 1994.
against excitotoxic brain damage in the newborn mice: implications for cerebral palsy,” *Neuroscience*, vol. 173, pp. 156–168, 2011.

[49] I. Zemlyak, N. Manley, I. Vulih-Shultzman et al., “The microtubule interacting drug candidate NAP protects against kainic acid toxicity in a rat model of epilepsy,” *Journal of Neurochemistry*, vol. 111, no. 5, pp. 1252–1263, 2009.

[50] I. Gozes, R. A. Steingart, and A. D. Spier, “NAP mechanisms of neuroprotection,” *Journal of Molecular Neuroscience*, vol. 24, no. 1, pp. 67–72, 2004.

[51] Y. Sari, “Activity-dependent neuroprotective protein-derived peptide, NAP, preventing alcohol-induced apoptosis in fetal brain of C57BL/6 mouse,” *Neuroscience*, vol. 158, no. 4, pp. 1426–1435, 2009.

[52] I. Zemlyak, N. Manley, R. Sapolsky, and L. Gozes, “NAP protects hippocampal neurons against multiple toxins,” *Peptides*, vol. 28, no. 10, pp. 2004–2008, 2007.

[53] M. Pascual and C. Guerri, “The peptide NAP promotes neuronal growth and differentiation through extracellular signal-regulated protein kinase and Akt pathways, and protects neurons co-cultured with astrocytes damaged by ethanol,” *Journal of Neurochemistry*, vol. 103, no. 2, pp. 557–568, 2007.

[54] I. Zemlyak, S. Furman, D. E. Brenneman, and I. Gozes, “A Novel peptide prevents death in enriched neuronal cultures,” *Regulatory Peptides*, vol. 96, no. 1-2, pp. 39–43, 2000.

[55] R. Zaltzman, A. Alexandrovich, S. M. Beni, V. Trembovler, E. Shohami, and I. Gozes, “Brain injury-dependent expression of activity-dependent neuroprotective protein,” *Journal of Molecular Neuroscience*, vol. 24, no. 2, pp. 181–187, 2004.

[56] I. Zemlyak, S. Furman, D. E. Brenneman, E. Shohami, and I. Gozes, “A Novel peptide prevents death in enriched neuronal cultures,” *Regulatory Peptides*, vol. 96, no. 1-2, pp. 39–43, 2000.

[57] R. Fernandez-Montesinos, M. Torres, D. Baglietto-Vargas et al., “Activity-dependent neuroprotective protein (ADNP) expression in the amyloid precursor protein/presenilin 1 mouse model of Alzheimer’s disease,” *Journal of Molecular Neuroscience*, vol. 41, no. 1, pp. 114–120, 2010.

[58] I. Gozes, R. Zaltzman, J. Hauser, D. E. Brenneman, E. Shohami, and J. M. Hill, “The expression of activity-dependent neuroprotective protein (ADNP) is regulated by brain damage and treatment of mice with the ADNP derived peptide, NAP, reduces the severity of traumatic head injury,” *Current Alzheimer Research*, vol. 2, no. 2, pp. 149–153, 2005.

[59] S. Furman, R. A. Steingart, S. Mandel et al., “Subcellular localization and secretion of activity-dependent neuroprotective protein in astrocytes,” *Neuron Glia Biology*, vol. 1, pp. 193–199, 2004.

[60] R. A. Steingart and I. Gozes, “Recombinant activity-dependent neuroprotective protein protects cells against oxidative stress,” *Molecular and Cellular Endocrinology*, vol. 252, no. 1-2, pp. 148–153, 2006.

[61] V. L. Smith-Swintosky, I. Gozes, D. E. Brenneman, M. R. D’Andrea, and C. R. Plata-Salaman, “Activity-dependent neurotrophic factor-9 and NAP promote neurite outgrowth in rat hippocampal and cortical cultures,” *Journal of Molecular Neuroscience*, vol. 25, no. 3, pp. 225–237, 2005.

[62] W. A. Lagrèze, A. Pielen, R. Steingart et al., “The peptides ADNF-9 and NAP increase survival and neurite outgrowth of rat retinal ganglion cells in vitro,” *Investigative Ophthalmology and Visual Science*, vol. 46, no. 3, pp. 933–938, 2005.

[63] I. Gozes and I. Divinski, “NAP, a neuroprotective drug candidate in clinical trials, stimulates microtubule assembly in

the living cell,” *Current Alzheimer Research*, vol. 4, no. 5, pp. 507–509, 2007.

[64] I. Divinski, L. Mittelman, and I. Gozes, “A femtomolar acting octapeptide interacts with tubulin and protects astrocytes against zinc intoxication,” *Journal of Biological Chemistry*, vol. 279, no. 27, pp. 28531–28538, 2004.