Keywords: Ocular injury; Abusive head trauma; Inflicted head injury

Abbreviations: AHT: Abusive Head Trauma; BFHI: Blunt Force Head Injury; BS: Brain Swelling; COD: Cause of Death; DOD: Date of Death; DH: Disc Hemorrhage; EBS: Endbulb Swelling; GHII: Global Hypoxic-Ischemic Injury; IOP: Intraocular Pressure; LC: Lamina Cribrosa

Introduction

Abusive head trauma (AKA: Shaken Baby Syndrome, Shaken Impact Syndrome; Inflicted Head Injury; Whiplash Shake Syndrome, Non-Accidental Head Trauma) remains a major societal problem and is among the most controversial and complex medical-legal issues worldwide by virtue of the difficulty of achieving clear proof of criminal intent by suspected perpetrators [1,2]. In the United States, trial by jury initiated by the local district attorney is the final arbiter. Trials typically involve testimonies from multiple witnesses including first responders, family members and medical experts for both the prosecution and defense.

Ocular injury, especially retinal hemorrhages, have been documented in 40-75% of reported AHT cases [3-5]. Other features of fatal cases often include somnolence on admission to emergency rooms, subarachnoid and subdural hemorrhage, skull and other fractures, apnea and hypoxia related to severe intracranial pressure (ICP) elevation inhibiting blood flow to the brain resulting in severe brain edema with fontanelle widening. Upper cervical cord injury from exaggerated head movement including cervical cord compression during shaking has been suggested as the primary trigger for central nervous system (CNS) hypoxia. Intraocular and anterior optic nerve sheath hemorrhages, commonly observed, presumably result from basal cistern blood migrating forward under the influence of high intracranial pressure through the subdural and subarachnoid spaces around the nerve in continuity with the basal cistern.

Neuropathology examination at autopsy often demonstrates a necrotic “respirator brain” as well as multiple other features [6-8]. Typical ocular pathology findings besides optic nerve sheath and retinal hemorrhages, sometimes include a macular ridge and hemorrhagic retinal schisis, and even retinal dehiscence [6]. Papilledema in some is a likely consequence of elevated intracranial pressure (ICP). Intraocular choroidal and retinal venous congestion and retinal venous congestion secondary to high ICP inhibiting blood return to the cavernous sinus could contribute to retinal and vitreous hemorrhages and possibly to elevation of IOP, seldom if ever measured pre-mortem in AHT cases. Recent scanning electron microscopy (SEM) studies offer an explanation for the reportedly common macular hemorrhagic retinal schisis due to substantial vitreo-retinal attachments peculiar to infants [9].

APP-A4 (AKA: Amyloid beta A4 protein, β pre-amyloid human protein, Alzheimer precursor protein, Alzheimer protein, neurofibrillary protein), came to high scientific interest in the late 1980s and is a widely available and sensitive neuropathology marker for identifying axonal injury consistent with severe shearing injury to myelinated and non-myelinated human CNS white matter [10-12]. APP-A4, currently being investigated for its role in Alzheimer disease, is a type-I transmembrane protein with a complex multiple domain genetic organization, now recognized as an evolutionarily conserved protein participating in multiple essential intracellular processes in all vertebrates. The human APP gene is expressed in glial and neuronal cells, but also in almost all tissues that have been examined [13]. Evidence suggests that a fragment of APP-A4 after α-secretase cleavage displays neuroprotective and trophic properties and reduces the degeneration of dendrites in experimental injury. For example, post-traumatic administration of APPα reduced the number of apoptotic neurons in a mouse model of brain injury [14]. Retinal ganglion cells (RGCs) manufacture the precursors in their cytoplasm and move them to sites of axonal injury in the rapid orthogonal transport system, somewhat akin to first responders in civil disasters.11

Materials and Methods

The selection of the first three cases was based on our observation of specific, not previously reported features, of optic nerve injury in AHT. The fourth case illustrates an important impact of premortem autopsies and forensic neuropathology in identifying inflicted head trauma.
previously mentioned but not illustrated symmetry across companion eyes we found common to our larger unpublished series [8]. We do not have access to multiple sets of autopsy eyes from children expiring for non-suspected AHT as they are not routinely sent out for ophthalmic pathology examination. We believe demonstrating that symmetrically heavy staining by APP-A4 of retinal ganglion cells across companion eyes and dissimilar findings in companion optic nerves constitute adequate controls for our purposes and conclusions. Background staining in non-injured anterior optic nerve segments was always present, but tapered rapidly posterior to the lamina usually absent by 10mm posterior to the globe.

An Appendix describes the APP-A4 technique. The cost per slide at our institution is $35.00/each (Table 1).

### Table 1: Case Data, Modified from: Reichard RR, et al. [8].

<table>
<thead>
<tr>
<th>#</th>
<th>Age (mo)</th>
<th>Sex</th>
<th>DOD COD</th>
<th>MOD</th>
<th>RH</th>
<th>DH</th>
<th>ONH</th>
<th>SDH</th>
<th>BS</th>
<th>Sex</th>
<th>Sex Ddays</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>10 mo M</td>
<td>40 years IHC stability</td>
<td>APP-A4 + OU</td>
<td>11/6/1976</td>
<td>AHT</td>
<td>Fe+ OU RT OD</td>
<td>Yes</td>
<td>No</td>
<td>No</td>
<td>Yes</td>
<td>Yes</td>
</tr>
<tr>
<td>2</td>
<td>45 mo F</td>
<td>APP-A4+ OU at anterior edge of ON infarct OD</td>
<td>2/17/2011</td>
<td>Homicide at trial</td>
<td>OU; all layers Fe neg.</td>
<td>Yes OD Fe neg.</td>
<td>Yes</td>
<td>Yes</td>
<td>OU Fe neg</td>
<td>Yes</td>
<td>Yes</td>
</tr>
<tr>
<td>3</td>
<td>4 mo F</td>
<td>Contusion ON OD bracketed by +APP-A4</td>
<td>10/27/2015</td>
<td>AHT</td>
<td>Suspect</td>
<td>OU; all layers Mild VH Fe neg.</td>
<td>No</td>
<td>Yes</td>
<td>Yes</td>
<td>Fe neg</td>
<td>Yes</td>
</tr>
<tr>
<td>4</td>
<td>3 mo F</td>
<td>Bilateral APP-A4 macular input &amp; EBS in lamina</td>
<td>1/18/2013</td>
<td>Homicide at trial</td>
<td>OU; all layers Fe neg.</td>
<td>No</td>
<td>No</td>
<td>No</td>
<td>No</td>
<td>Yes</td>
<td>Yes</td>
</tr>
</tbody>
</table>

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### Results

We obtained heavy APP-A4 staining in RGCs from the 40-year-old Case #1 demonstrating potential for remarkable stability and durability of the protein components in formalin-fixed paraffinized tissues (Figure 1). Obviously one such demonstration does not establish that any similarly aged or older paraffin material will reliably work with this marker: However this one example is a good reason to further explore the use of this APP-A4 marker in long-stored paraffin embedded human ocular tissue if results might be pertinent to ongoing human autopsy studies. Our larger series of cases will include several in which paraffin embedding dates back to 2010, allowing better estimation of the marker’s durability in general over at least 10 years. Case #2 demonstrated an optic nerve infarct beginning approximately 6mm posterior to the globe, expanding to involve nearly the whole diameter of the nerve in deeper sections from the most posterior portion, some 29mm behind the globe likely into the chiasm since the orbital and optic canal portions of the nerve together average approximately 29mm in length (Figure 2). In this example, APP-A4 intra-axonal accumulations were found clearly outlining the anterior end of the infarct in agreement with routine Hematoxylin and Eosin (H&E) and Periodic Acid-Schiff (PAS) stains (Figure 2A-D). While not surprising this does illustrate how blocked orthograde axonal transport is sharply interrupted by an adjacent infarction. Case #3 illustrates for the first time both orthograde and retrograde axonal transport block corresponding to anterior and posterior sides of a hemorrhagic optic nerve contusion (Figure 3A-C & Figure 4). Case #4, to our knowledge is the best demonstration ever illustrated demonstrating bilaterally symmetrical APP-A4 macular input and classic end-bulb swellings in the lamina cribrosa OU, in our view reinforcing no need for non-AHT controls (Figure 5) [8,9]. The symmetry of APP-A4 staining of RGCs and optic nerve heads with laminar end-bulb swellings in axons across companion eyes in this case well documented expected vigorous bilateral pre-mortem orthograde axonal transport physiology at least to the lamina cribrosa.

### Discussion

The infarction of the optic nerve in Case #2 we believe expanded anteriorly from the CNS along its course to within 10mm of the globe where it tapered anteriorly. Since the ON measures about 1mm in the globe and 24mm in the orbit the ON segment of...
approximately 29mm would have extended into the chiasm. As additional evidence of chiasmal extension the posterior ON segment displayed dwindling pialseptae, characteristic of the chiasm. We suspect this injury was an extension of the global ischemic hypoxic injury (GHII) found during the neuropathology examination at autopsy. Normally, over the approximately 24mm course from eye to chiasm, both internal and external carotid arterial input supply blood to the optic nerve. Compromise of the ophthalmic artery’s orbital branches blood flows presumably occurs in GHII. To our knowledge, this case documents the first histological evidence of extension far anteriorly of GHII into the optic nerve. We otherwise found only suggestions and indirect evidence of such in experimental and human studies in current literature. Lamina cribrosa accumulation of APP-A4 after even transient IOP elevation would correlate well with past acute ocular hypertension experimental studies in foveated monkeys revealing lamina cribrosa blockade of both orthograde and retrograde axonal transport precisely in that location [15].

Conclusion

Three of these four cases demonstrate previously unreported new findings related to optic nerve injury in AHT. Axonal end bulb swellings in the lamina cribrosa, as in Case #4, have been previously reported using APP-A4 IHC without speculation as to the pathophysiology of their occurrence [8]. We suspect that IOP is at least transiently elevated in fatal AHT cases probably due to several factors including high ICP and related migration of blood into the nerve’s sheath spaces posterior to the globe, venous congestion of choroidal and retinal vessels, and vitreous hemorrhage in some with secondary hemorrhagic glaucoma. Ongoing studies of our whole series of AHT cases, to be reported in a separate manuscript, will explore correlations relevant to this concept. Controversy persists among forensic pathologists and defense lawyers as to the validity of AHT diagnoses. Even in instances where most of the typical findings of AHT are present, alternative diagnoses are often suggested by the defense and none of the typical findings are acknowledged as specific for AHT [3]. Much research is yet needed to develop specific markers of AHT assuming they exist; nevertheless APP-A4 can contribute valuable information to the diagnoses in these unfortunate and complex cases [7].

Acknowledgements

Funding/support: Supported in part by an unrestricted grant from Research to Prevent Blindness.
Conflict of Interest Disclosures

Authors in this paper have no financial or proprietary interest in a product, method, or material published in this manuscript. All authors have completed and submitted the ICMJE Form for Disclosure of Potential Conflicts of Interest. All information and materials in the manuscript are original.

IRB Approval

University of California Irvine IRB approval not required for small case series with clinical-pathological correlation.

References


