Traumatic Optic Nerve Injury Elevates Plasma Biomarkers of Traumatic Brain Injury in a Porcine Model

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Abstract

A diagnosis of traumatic brain injury (TBI) is typically based on patient medical history, a clinical examination, and imaging tests. Elevated plasma levels of glial fibrillary acidic protein (GFAP), ubiquitin c-terminal hydrolase L1 (UCH-L1), and neurofilament light chain (NFL) have been observed in numerous studies of TBI patients. It is reasonable to view traumatic optic neuropathy (TON) as a focal form of TBI. The purpose of this study was to assess if circulating GFAP, UCH-L1, and NFL are also elevated in a porcine model of TON. Serum levels of GFAP, UCH-L1, and NFL were measured immediately before optic nerve crush and 1 h post-injury in 10 Yucatan minipigs. Severity of optic nerve crush was confirmed by visual inspection of the optic nerve at time of injury, loss of visual function as measured by flash visual evoked potential (fVEP) at 7 and 14 days, and histological analysis of axonal transport of cholera toxin-b (CT-b) within the optic nerve. Post-crush concentrations of GFAP, UCH-L1, and NFL were all significantly elevated compared with pre-crush concentrations (p < 0.01, p = 0.01, and p < 0.01, respectively). The largest increase was observed for GFAP with the post-injury median concentration increasing nearly sevenfold. The use of these TBI biomarkers for diagnosing and managing TON may be helpful for non-ophthalmologists in particular in diagnosing this condition. In addition, the potential utility of these biomarkers for diagnosing other optic nerve and/or retinal pathologies should be evaluated.

Keywords: optic nerve; porcine model; traumatic optic neuropathy

Introduction

A diagnosis of traumatic brain injury (TBI) is based on patient medical history, a clinical examination, and imaging tests.1–5 The ability to diagnose and classify TBI as well as to decide when imaging is indicated has advanced with the identification of circulating biomarkers often elevated in TBI patients. Specifically, the combined measurement of glial fibrillary acidic protein (GFAP, an abundant intermediate filament contained within astrocytes) and ubiquitin c-terminal hydrolase L1 (UCH-L1, a neuronal cystosolic enzyme) was recently approved by the United States Food and Drug Administration (FDA) as part of the Banyan Trauma Indicator (BTI) to aid in the evaluation of patients 18 years of age with a suspected TBI (Glasgow Coma Scale [GCS] score 13–15) when collected within 12 h of the injury. ALERT-TBI, the landmark multi-center observational study, key in obtaining FDA approval for these biomarkers, collected data from 1959 patients with suspected non-penetrating TBI and a Glasgow Coma Scale score of 9–15. The authors found that the combined use of GFAP and UCH-L1 had a sensitivity of 97.6% and negative predictive value (NPV) of 99.6% for the detection of acute intracranial lesions on computed tomography (CT).6 Other biomarkers, including neurofilament light chain (NFL, a neuron specific intermediate filament) are also elevated in TBI but are not FDA approved for use in TBI evaluation.6–20

It is reasonable to consider isolated traumatic optic neuropathy (TON) a focal form of TBI. The optic nerve is a specialized structure of the central nervous system (CNS) that serves the visual system. Although it is the second member of the 12 cranial “nerves,” the optic nerve is actually a white matter tract of the brain. It is (1) embryologically derived from the diencephalon; (2) myelinated by oligodendrocytes rather than Schwann cells; (3) surrounded by layers of dura, arachnoid, and pia as elsewhere in the brain; and (4) has a blood–brain barrier.21–30 Its axons are derived from retinal ganglion cells (RGCs), which are themselves specialized CNS neurons.27,31 Like other CNS neurons, damage to these axons results in Wallerian degeneration and apoptosis of their RGC cell bodies.32,33

As the name would suggest, TON is an injury caused by trauma to the optic nerve. TON injuries are usually divided into either
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Direct (e.g., penetrating injury) or indirect mechanisms of injury (e.g., concussive force transmitted to the optic nerve with globe or head trauma). Visual loss is usually immediate, but can be delayed in at least 10% of cases, presumably via secondary mechanisms that have not yet been fully elucidated. A prospective surveillance study conducted in the United Kingdom found that TON affects 1 in 1,000,000 of the general population. TON is present in 0.7–2.5% of cases of blunt and penetrating head and facial injury. Younger adult males predominate and male prevalence and other characteristics of TON are similar in pediatric and adult populations.

In a combat environment, the incidence of eye injuries among all combat-related injuries has been reported as being 13%, with TON making up 20% of eye-related injuries. Interestingly, seemingly trivial injuries such as trauma to the forehead with no alteration of awareness or loss of consciousness can cause TON. However, the trauma associated with TON is more often severe. Prospective and retrospective studies show that 34–75% of TON cases are associated with a loss of consciousness. There have not been any well-designed prospective studies to delineate optimum clinical management of patients with TON. Current mainstay management options are steroids, surgical management, and/or observation.

As with TBI, a TON diagnosis is based on patient medical history and clinical examination. In TON, imaging and other tests serve an adjunct role. Obtaining a medical history and testing for a relative afferent pupillary defect (RAPD, the “swinging flashlight” examination), visual acuity, color plate testing, the dilated fundoscopic examination (DFE) and static optic perimetry are mainstays of establishing a TON clinical diagnosis. Optical coherence tomography (OCT), magnetic resonance imaging (MRI), visual evoked potentials (VEP), CT, and ultrasound are all additional adjunctive tools.

Although all of these methods can be very useful, they have some inherent limitations. For example, testing for an RAPD is often limited by the examiner’s experience. This is problematic, as testing for an RAPD is considered by neuro-opthalmologists to be the most important clinical test in diagnosing TON. Non-ophthalmologists in particular often fail to properly perform and interpret this test. Imaging and ancillary tests such as OCT will also frequently be unrevealing, especially in the acute stages of disease, as it can take days to even weeks for some of these tests to show evidence of injury, if they do at all. Currently, no laboratory test exists to specifically assess the structural integrity or function of the optic nerve in a traumatic setting.

The purpose of this study was to assess if the biomarkers elevated after TBI were also elevated after development of isolated TON in a porcine model. This was a small nested study, conducted as part of a larger experiment investigating potential optic nerve regeneration, the details of which will be described in future publications. The Quanterix Simoa Human Neurology 4-plex A (N4PA) kit, which targets GFAP, UCH-L1, NFL, and total tau (t-tau, a group of alternatively spliced, axon-resident, microtubule-associated phosphoproteins), was used to measure these biomarkers in plasma. This assay is not FDA approved for clinical use, but is currently being used across the world for research purposes.

In efforts to elucidate the conditions conducive to the regeneration of RGCs and their axons, a well-established murine optic nerve crush model of TON has been developed, characterized, and extensively used. Using this model as a paradigm, we developed a porcine optic nerve crush model of TON to research neuroprotective and neuro-regenerative optic nerve treatments that have had some success in the murine model. Our porcine TON model demonstrates loss of visual function as proven by pathophysiological depression of flash visual evoked potential (fVEP) in addition to loss of normal physiological function of the optic nerve as demonstrated by abolished anterograde, axonal transport of cholera toxin-b subunit (CT-b). Within this investigation of optic nerve regeneration, we performed this pilot study to determine if the TBI biomarkers mentioned are also elevated in isolated TON.

Methods

Ten female Yucatan minipigs each weighing ~9–13 kg were independently evaluated and screened for pre-existing disease 3–7 days prior to the surgical procedure. Baseline flash fVEP were obtained at this time with a RETeval handheld electrophysiology device (LK C Technologies; Gaithersburg, MD) to ensure objective evidence of visual function. Access to the orbital portion of the optic nerve was achieved by lateral orbitotomy. Once generous exposure of the optic nerve was obtained (mean post-initial anesthesia time ± standard deviation (SD): 5 h 21 min ±49 min), blood was drawn from the right saphenous vein using standard phlebotomy techniques. A 23 gauge butterfly needle and lavender top tubes were used to collect two mL of whole blood immediately before and then 1 h after traumatic optic nerve injury. Injury was performed by crushing the optic nerve with a neurosurgical aneurysm clip (Aesculap Inc., Center Valley, PA, CAT No. FT820T) and reinforcing it by the addition of a titanium reinforcing clip (Aesculap Inc., Center Valley, PA, CAT No. FT900T), inducing ~12 N of crush force at the application site (measured using FlexiForce ELF System [Tekscan; South Boston, MA]). Blood was centrifuged at 2013g for 20 min at 4°C, and plasma volumes were aliquoted into 1.5 mL tubes and stored at -80°C. Plasma samples were shipped overnight to Quanterix (Billerica, MA) on dry ice. The Simoa Human Neurology 4-plex A (N4PA) assay was performed by Quanterix personnel under blinded conditions. Samples were run in duplicate at a dilution factor of four. Appropriate plate controls to establish limits of detection (LOD) and limits of quantification (LOQ) were used.

To confirm the validity and reproducibility of the crush lesions and that loss of visual function and loss of normal physiological measures had occurred, optic nerve crush was clearly visualized in each animal by the surgeon. Weekly serial fVEPs of the right and left eyes were obtained with the animals under general anesthesia. Axonal transport within the optic nerves using CT-b was also assessed. Two days before study end-point, 200 μg of CT-b in 150 μL of sterile saline was injected into the vitreous cavity of the left (uncrushed), and right (crushed) eyes of each pig. Eyes were fixed by transcardial perfusion fixation with 10% neutral buffered formalin with the animals under general anesthesia time

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Antibodies in the N4PA assay are known to cross-react with three of the four biomarkers from porcine samples. However, the monoclonal antibody used in the assay to measure the t-tau biomarker in human plasma does not cross-react with porcine t-tau (see Quanterix application note “Porcine Cross Reactivity in Human Simoa Assays”). Therefore, the t-tau biomarker data were not included in the statistical analysis. Samples with biomarker concentrations below the LOQ were assumed to have a concentration equal to the biomarker-specific LOQ (n = 4).

Statistical analysis

Differences between pre- and post-injury concentrations were examined using the Wilcoxon signed rank test. All analyses were conducted using SAS version 9.4 (Cary, NC), and statistical significance was evaluated using an α of 0.05.
Results

The distributions of pre- and post-injury biomarker concentrations for GFAP, UCH-L1, and NFL are illustrated in Figure 1. All three were significantly elevated post-crush. The largest increase was observed for GFAP, with a pre-injury median (interquartile range [IQR]) concentration of 7.76 pg/mL (5.48–15.3) and a post-injury median (IQR) concentration of 52.3 pg/mL (32.3–53.4) (p < 0.01). Increases in UCH-L1 (pre-injury median [IQR]: 63.6 pg/mL [32.8–121.0] vs. post-injury median [IQR]: 77.6 pg/mL [40.1–141.0]) and NFL (pre-injury median [IQR]: 11.8 pg/mL [8.89–14.8] vs. post-injury median [IQR]: 13.9 pg/mL [11.0–17.5]) were much smaller, but still statistically significant (p = 0.01 and p < 0.01, respectively).

There was strong evidence the optic nerves were severely and traumatically injured. Severe crush was visually confirmed at the

FIG. 1. Distribution of plasma concentrations of traumatic brain injury (TBI) biomarkers before and after optic nerve (ON) crush. Post-injury concentrations of all biomarkers were significantly elevated compared with pre-injury concentrations. Whiskers represent the 10th and 90th percentiles. (A) Glial fibrillary acidic protein (GFAP): pre-injury median (interquartile range [IQR]) concentration is 7.76 pg/mL (5.48–15.3) and post-injury median (IQR) concentration is 52.3 pg/mL (32.3–53.4) (p < 0.01). (B) Ubiquitin c-terminal hydrolase L1 (UCH-L1): pre-injury median (IQR) concentration is 63.6 pg/mL (32.8–121.0) and post-injury median (IQR) concentration is 77.6 pg/mL (40.1–141.0) (p = 0.01). (C) Neurofilament light chain (NFL): pre-injury median (IQR) concentration is 11.8 pg/mL (8.89–14.8) and post-injury median (IQR) concentration is 13.9 pg/mL (11.0–17.5) (p < 0.01).

FIG. 2. Assessment of porcine optic nerve form and function before and after optic nerve crush. (A) Baseline flash visual evoked potential (fVEP) recordings were obtained from 15 Yucatan minipigs 3–7 days prior to optic nerve crush. For all fVEP recordings, replicates were performed for oculus dextrus (OD) followed by oculus sinister (OS) (n = 30 waveforms). fVEP recording 7 days (B) and 14 days (C) after optic nerve crush from saline-only injected minipigs (n = 4 pigs, 8 waveforms per eye per time point). (D) Axonal transport of cholera toxin-β subunit (CT-β) in uncrushed and crushed optic nerves. Two days before euthanasia and harvest of eyes and optic nerves, 500 µg of CT-β was dissolved in 250 µL of injectable saline and 100 µL of CT-β solution was injected into the vitreous of both eyes. Two days after injection, the pigs were euthanized via transcardial perfusion fixation with 10% formalin, and the eyes and optic nerves were harvested. The optic nerves and peripapillary retina were then prepared for frozen sectioning. The right (crushed) optic nerve (D1) shows fluorescent signal from the CT-β up to, but not beyond, the crush site. In the immunofluorescence pattern obtained from the left eye (D2), the CT-β signal can be seen through the entire length of the nerve, demonstrating intact, unimpaired axonal transport throughout its entire length.
time of crush by the surgeon. Normal fVEPs were evident in uncrushed control eyes compared with a remarkably flat fVEP (after N1 waveform) in crushed eyes (Fig. 2, A–C). The flat fVEP response, indicating loss of visual function, fully appeared 7 days after optic nerve crush. This response was maintained through 14 days post-injury (Fig. 2 B and C). Additionally, axonal transport of CT-β was almost completely abolished posterior to the crush site (Fig. 2 D).

Discussion

The diagnosis and management of many diseases has benefited from the revolution in molecular diagnostics. However, this largely has not been the case in the diagnosis of optic nerve pathology. TON is a significant cause of trauma-related vision loss. Using a plasma-based enzyme-linked immunosorbent assay to measure biomarker concentrations in a porcine model of TON, we observed significant increases in the TBI biomarkers GFAP, UCH-L1, and NFL, with the post-injury median concentration of GFAP increasing nearly sevenfold. This is the first documented instance of TON being studied or detected using a laboratory assay of plasma biomarkers.

The research presented here is subject to a number of limitations. A relatively small number of animals (n = 10) was used. Post-hoc power calculations revealed that analyses of both the GFAP and NFL data were sufficiently powered (1-β > 80%). Statistical analysis of UCH-L1 data was underpowered, likely as a result of the large variability in the post-injury concentrations. Nonetheless, differences in pre- and post-injury concentrations were statistically and clinically significant for all three biomarkers. Another limitation of the current study was that only female pigs were used. Although we were unable to evaluate whether biomarker concentrations varied by sex, exclusion of male pigs was also a strength of the study as it prevented any potential confounding because of this factor. Finally, the severity of the optic nerve crush in this animal model is a weakness in that the nature of the injury is much more severe than many clinical cases of TON. Indeed, there was no doubt that each animal’s optic nerve was traumatically crushed. However, the severity of the injury prevented us from evaluating whether these biomarkers would be elevated in milder cases of TON.

As described previously, the broader objective of the primary experiment was to investigate porcine optic nerve regeneration. To measure TBI biomarkers as part of a sub-experiment, plasma samples were collected immediately before the optic nerve was crushed and 1 h post-crush. The 1 h time point post-crush was chosen for two reasons. First, previous work using a swine model of CT-β showed a linear increase in circulating GFAP and NFL concentrations 2 h after injury (see Quanterix application note “Porcine Cross Reactivity in Human Simoa Assays”), indicating that biomarker concentrations would also likely increase at 1 h post-injury. Second, because treatments to promote optic nerve regeneration as part of the primary study needed to be administered at 1 h post-crush, blood for the biomarker assay had to be drawn prior to treating the animals to prevent the treatments from obscuring the true effects of the optic nerve crush on biomarker concentrations. Although our findings demonstrate that GFAP, UCH-L1, and NFL concentrations are significantly elevated within a short time after injury, additional research is necessary to determine the kinetics.

These biomarkers could potentially be elevated in other clinical situations in which there is (1) interruption of structural integrity of the optic nerve or (2) presence of other pathologies affecting the optic nerve or possibly the retina (given the axons of the optic nerve originate from RGCs). Conceivably, increases in these plasma biomarkers that are a result of other optic nerve and/or retinal pathologies could have important clinical implications and need to be evaluated.

As discussed previously, the optic nerve is a white matter tract of the brain. The elevation of TBI plasma biomarkers in our model of TON supports the contention that TON is a focal form of TBI. A certain quandary exists, however, in that when there is concomitant damage to other portions of the brain beyond the optic nerve (a relatively common occurrence in TON), the elevation of plasma biomarkers cannot distinguish the regional source of the CNS damage. Nonetheless, well-designed human studies as well as additional animal studies may help determine if this assay is useful in both the diagnosis and management of TON in certain instances.

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Author Disclosure Statement

No competing financial interests exist. The views expressed in this article are those of the authors and do not reflect the official policy or position of the United States Army Medical Department, Department of the Army, Department of Defense, or the United States Government. Research was conducted in compliance with the Animal Welfare Act, the implementing Animal Welfare Regulations, and the principles of the Guide for the Care and Use of Laboratory Animals, National Research Council. The facility’s Institutional Animal Care and Use Committee approved all research conducted on this study. The facility where this research was conducted is fully accredited by the American Association for Accreditation of Laboratory Animal Care (AAALAC).

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