







Accelerated Aging after Traumatic Brain Injury: An ENIGMA Multi-Cohort Mega-Analysis

Emily L Dennis, PhD ,^{1,2} Samantha Vervoordt, MS,³ Maheen M Adamson, PhD,^{4,5} Amiri Houshang, PhD,^{6,7} Erin D Bigler, PhD,^{1,8} Karen Caeyenberghs, PhD ,⁹ James H Cole, PhD ,^{10,11} Kristen Dams-O'Connor, PhD,^{12,13} Evelyn M Deutscher, PhD,⁹ Ekaterina Dobryakova, PhD,^{14,15} Helen M Genova, PhD,^{15,16} Jordan H Grafman, PhD,¹⁷ Asta K Håberg, MD, PhD,^{18,19} Torgeir Hellstrøm, MD, PhD,²⁰ Andrei Irimia, PhD,^{21,22,23} Vassilis E Koliatsos, MD, MBA,^{24,25} Hannah M Lindsey, PhD,¹ Abigail Livny, PhD,^{26,27,28} David K Menon, PhD,²⁹ Tricia L Merkley, PhD,^{1,8} Abdalla Z Mohamed, PhD,³⁰ Stefania Mondello, MD, MPH PhD ,³¹ Martin M Monti, PhD ,^{32,33} Virginia FJ Newcombe, MD, PhD ,²⁹ Mary R Newsome, PhD,^{1,2} Jennie Ponsford, PhD,^{34,35} Amanda Rabinowitz, PhD,^{36,37} Hanne Smevik, PhD,^{38,39} Gershon Spitz, PhD,^{34,35,40} Umesh M Venkatesan, PhD,^{36,37} Lars T Westlye, PhD,^{41,42} Ross Zafonte, DO,^{43,44} Paul M Thompson, PhD,^{45,46} Elisabeth A Wilde, PhD,^{1,2} Alexander Olsen, PhD,^{38,39,47†} and Frank G Hillary, PhD^{3†}

View this article online at [wileyonlinelibrary.com](https://www.wileyonlinelibrary.com). DOI: 10.1002/ana.26952

Received Oct 21, 2023, and in revised form Apr 10, 2024. Accepted for publication Apr 15, 2024.

Address correspondence to Prof Frank G. Hillary, Department of Psychology, Pennsylvania State University, State College, PA 16801, USA. E-mail: fhillary@psu.edu. Prof Alexander Olsen, Department of Psychology, Norwegian University of Science and Technology, Sem Sæløds vei 7, 7491 Trondheim, Norway. E-mail: alexander.olsen@ntnu.no

[†]These authors contributed equally to this manuscript.

From the ¹Department of Neurology, University of Utah, Salt Lake City, UT, USA; ²George E. Whalen Veterans Affairs Medical Center, Salt Lake City, UT, USA; ³Department of Psychology, Penn State University, State College, PA, USA; ⁴Women's Operational Military Exposure Network (WOMEN) & Rehabilitation, VA Palo Alto Healthcare System, Palo Alto, CA, USA; ⁵Neurosurgery, Stanford University School of Medicine, Stanford, CA, USA; ⁶Neuroscience Research Center, Institute of Neuropharmacology, Kerman University of Medical Sciences, Kerman, Iran; ⁷Department of Radiology & Nuclear Medicine, Amsterdam UMC, Amsterdam, The Netherlands; ⁸Department of Psychology and Neuroscience Center, Brigham Young University, Provo, UT, USA; ⁹Cognitive Neuroscience Unit, School of Psychology, Deakin University, Geelong, Australia; ¹⁰Centre for Medical Image Computing, Computer Science, University College London, London, UK; ¹¹Dementia Research Centre, Queen Square Institute of Neurology, University College London, London, UK; ¹²Department of Rehabilitation and Human Performance, Icahn School of Medicine at Mount Sinai, New York, NY, USA; ¹³Department of Neurology, Icahn School of Medicine at Mount Sinai, New York, NY, USA; ¹⁴Center for Traumatic Brain Injury, Kessler Foundation, East Hanover, NJ, USA; ¹⁵Rutgers New Jersey Medical School, Newark, NJ, USA; ¹⁶Center for Neuropsychology and Neuroscience Research, Kessler Foundation, East Hanover, NJ, USA; ¹⁷Shirley Ryan Ability Lab, Chicago, IL, USA; ¹⁸Department of Neuromedicine and Movement Science, Faculty of Medicine and Health Sciences, NTNU, Trondheim, Norway; ¹⁹Department of Radiology and Nuclear Medicine, St. Olavs Hospital, Trondheim University Hospital, Trondheim, Norway; ²⁰Department of Physical Medicine and Rehabilitation, Oslo University Hospital, Oslo, Norway; ²¹Ethel Percy Andrus Gerontology Center, Leonard Davis School of Gerontology, University of Southern California, Los Angeles, CA, USA; ²²Alfred E. Mann Department of Biomedical Engineering, Viterbi School of Engineering, University of Southern California, Los Angeles, CA, USA; ²³Department of Quantitative and Computational Biology, Dornsife College of Arts and Sciences, University of Southern California, Los Angeles, CA, USA; ²⁴Departments of Pathology (Neuropathology), Neurology, and Psychiatry and Behavioral Sciences, Johns Hopkins University School of Medicine, Baltimore, MD, USA; ²⁵Neuropsychiatry Program, Sheppard and Enoch Pratt Hospital, Baltimore, MD, USA; ²⁶Division of Diagnostic Imaging, Sheba Medical Center, Tel-Aviv,

Objective: The long-term consequences of traumatic brain injury (TBI) on brain structure remain uncertain. Given evidence that a single significant brain injury event increases the risk of dementia, brain-age estimation could provide a novel and efficient indexing of the long-term consequences of TBI. Brain-age procedures use predictive modeling to calculate brain-age scores for an individual using structural magnetic resonance imaging (MRI) data. Complicated mild, moderate, and severe TBI (cmsTBI) is associated with a higher predicted age difference (PAD), but the progression of PAD over time remains unclear. We sought to examine whether PAD increases as a function of time since injury (TSI) and if injury severity and sex interacted to influence this progression.

Methods: Through the ENIGMA Adult Moderate and Severe (AMS)-TBI working group, we examine the largest TBI sample to date ($n = 343$), along with controls, for a total sample size of $n = 540$, to replicate and extend prior findings in the study of TBI brain age. Cross-sectional T1w-MRI data were aggregated across 7 cohorts, and brain age was established using a similar brain age algorithm to prior work in TBI.

Results: Findings show that PAD widens with longer TSI, and there was evidence for differences between sexes in PAD, with men showing more advanced brain age. We did not find strong evidence supporting a link between PAD and cognitive performance.

Interpretation: This work provides evidence that changes in brain structure after cmsTBI are dynamic, with an initial period of change, followed by relative stability in brain morphometry, eventually leading to further changes in the decades after a single cmsTBI.

ANN NEUROL 2024;00:1–13

Complicated mild (mild traumatic brain injury [TBI] with trauma-related intracranial pathology on computed tomography/magnetic resonance imaging [CT/MRI]), moderate, and severe TBI (cmsTBI) results in downstream consequences for brain structure and physiology, altering the course of brain aging and increasing risk for neurodegeneration.^{1,2} Traumatic axonal injury (TAI) is a hallmark pathology in TBI, and a predictor of brain atrophy, observable as Wallerian-like degeneration in the early phase and more recently been linked to promotion of toxic proteinopathies in the chronic phase.^{3,4} The trajectories for brain atrophy in remote cmsTBI (>10 years) samples have not been studied extensively but some evidence suggests a pattern distinct from that in Alzheimer's disease.⁵ Modifiers of aging after cmsTBI include chronic neuroinflammation,⁶ blood-brain barrier and microvascular disruption, and proteinopathy primarily involving tau, beta-amyloid, and alpha-synuclein.^{7,8} Given the multiple interacting factors influencing outcome in chronic TBI, time since injury (TSI) may be an important indicator of the degree of exposure to the pathology of TBI and, therefore, a moderator of long-term outcome.⁹

Brain-age-gap has been developed as a potential biomarker for outcome in psychiatric and neurological disorders. Brain-age is established by comparing brain

characteristics of an individual to their chronological age, determined using data from healthy participants.^{10,11} Studies of the predicted age difference (PAD) have been applied to a range of clinical populations, including depression,¹² PTSD,¹³ and stroke.¹⁴ Brain age prediction modeling shows that, within the first several years after cmsTBI, there is increased atrophy in gray and white matter equivalent to about half a decade in chronological age in msTBI,^{15–18} and 1–3 years in mild TBI.^{2,19,20} The extent to which post-traumatic atrophy may evolve as individuals transition through various stages of life, continuing into senescence for years and even decades, remains to be established.

Greater PAD after cmsTBI has been associated with greater injury severity and poorer cognitive function. However, it remains unclear whether the *brain atrophy* – observed in the well-documented sulcal and ventricular enlargement occurring over the first few years secondary to lesion resolution and transsynaptic or Wallerian degeneration²¹ – has long-term effects and even *accelerates* over the lifespan. While there is some evidence of an association between TSI and PAD in mid-life, interpreted as *accelerated aging*,¹⁵ other work has failed to substantiate this effect.¹⁷

This study extends prior work^{15–17} by analyzing a larger sample with a wider age range (18–85) and TSI (mean = 5.3 years \pm 6.4, range 0.2–34), with severity

Israel; ²⁷Faculty of Medicine, Tel-Aviv University, Tel-Aviv, Israel; ²⁸Sagol Neuroscience School, Tel-Aviv University, Tel-Aviv, Israel; ²⁹Department of Medicine, University of Cambridge, Cambridge, UK; ³⁰Thompson Institute, University of the Sunshine Coast, Birtinya, Australia; ³¹Department of Biomedical and Dental Sciences and Morphofunctional Imaging, University of Messina, Messina, Italy; ³²Department of Psychology, UCLA, Los Angeles, CA, USA; ³³Brain Injury Research Center (BIRC), Department of Neurosurgery, UCLA, Los Angeles, CA, USA; ³⁴Monash-Epworth Rehabilitation Research Centre, School of Psychological Sciences, Monash University, Melbourne, Australia; ³⁵School of Psychological Sciences, Monash University, Melbourne, Australia; ³⁶Moss Rehabilitation Research Institute, Elkins Park, PA, USA; ³⁷Department of Rehabilitation Medicine, Sidney Kimmel Medical College, Thomas Jefferson University, Philadelphia, PA, USA; ³⁸Department of Psychology, NTNU, Trondheim, Norway; ³⁹NorHEAD – Norwegian Centre for Headache Research, NTNU, Trondheim, Norway; ⁴⁰Department of Neuroscience, Central Clinical School, Monash University, Melbourne, Australia; ⁴¹Department of Psychology, University of Oslo, Oslo, Norway; ⁴²NORMENT, Division of Mental Health and Addiction, Oslo University Hospital, Oslo, Norway; ⁴³Department of Physical Medicine and Rehabilitation, Massachusetts General Hospital/Brigham & Women's Hospital, Boston, MA, USA; ⁴⁴Spaulding Rehabilitation Hospital, Boston, MA, USA; ⁴⁵Imaging Genetics Center, Stevens Neuroimaging & Informatics Institute, Keck School of Medicine of USC, Marina del Rey, CA, USA; ⁴⁶Departments of Neurology, Pediatrics, Psychiatry, Radiology, Engineering, and Ophthalmology, USC, Los Angeles, CA, USA; and ⁴⁷Clinic of Rehabilitation, St. Olavs Hospital, Trondheim University Hospital, Trondheim, Norway

ranging between complicated mild and severe. We leveraged a team science approach through ENIGMA (Enhancing Neuro-Imaging Genetics through Meta-Analysis)²² and the ENIGMA Adult msTBI (AMS-TBI) working group.²³ We hypothesized that we would see both *advanced brain age* in individuals with cmsTBI, and *accelerated brain aging*, defined as increasing PAD as a function of TSI. We hypothesized that injury severity, lower educational attainment, and poorer cognitive function would be independent predictors of greater PAD. Additionally, we examined the influence of sex on brain age trajectory without making specific predictions, due to mixed findings in the literature.²⁴ Finally, given its role as a potential risk factor for poor outcome after TBI,²⁵ we examined the role of genetic risk (apolipoprotein E [APOE]) on PAD as it interacts with TSI, anticipating that the $\epsilon 4$ allele would confer risk for more accelerated brain aging.

Methods

Study Samples

Study samples consisted of 7 cohorts from parent studies originating in 3 countries (see Table). The final sample size (as detailed below and in Fig 1) was 540: 343 cmsTBI

(237 M/106F, mean age = 44.5 ± 16.2 years, range = 20–85) and 197 control (113 M/84F, mean age = 38.6 ± 16.2 years, range = 18–84). There was a significant difference in age between groups ($p < 0.001$), partially attributable to 2 cohorts with older participants not including a control sample. Participants were recruited from hospitals, outpatient rehabilitation clinics, and the surrounding community. Details on the inclusion and exclusion criteria for each cohort are included in Supplementary Table S1. Original studies were reviewed by the appropriate institutional review board for each respective institution. All participants provided written or verbal informed consent as part of involvement with the parent study.

For the analyses in this study, only participants 18 years of age or older at the time of enrollment were included and participants who sustained their injury before the age of 18 were excluded from most analyses (N = 61) in order to avoid confounding brain changes associated with neurodevelopment with morphological alterations attributed to injury. Level of education was measured using ISCED 2011 categories.²⁶

Patients had to have sustained a cmsTBI, defined by having a TBI and trauma-related intracranial pathology

TABLE. Cohort Demographics

Cohort	Total N	Subgroup N	M/F	Age (SD) Range	TSI (in years; average range)	
Kessler	63	cmsTBI	36	25/11	40.2 (11.2) 21–65	7.4 (5.8) 1.4–27
		Control	27	12/15	41.2 (10.8) 21–63	–
LETBI	53	cmsTBI	53	30/23	58.0 (10.3) 40–85	11.1 (9.2) 1.2–34.4
		Control	0	–	–	–
Monash	97	cmsTBI	74	60/14	38.3 (14.5) 20–78	2.1 (3.1) 0.2–21.1
		Control	23	10/13	30.4 (12.6) 18–69	–
NTNU	113	cmsTBI	45	33/12	32.3 (11.3) 20–65	2.8 (1.1) 1.4–5.4
		Control	68	50/18	35.2 (14.3) 19–64	–
Oslo	64	cmsTBI	64	43/21	42.9 (13.1) 20–67	1.2 (0.4) 0.6–1.8
		Control	0	–	–	–
PSU	112	cmsTBI	53	36/17	55.4 (16.9) 20–79	9.3 (6.7) 0.6–23.6
		Control	59	31/28	45.3 (20.2) 18–84	–
VA Palo Alto	38	cmsTBI	18	10/8	39.9 (13.6) 24–71	8.1 (5.5) 0.7–20.2
		Control	20	10/10	37.4 (10.2) 23–54	–
Total	540	cmsTBI	343	237/106	44.5 (16.2) 20–85	5.3 (6.4) 0.2–34.4
		Control	197	113/84	38.6 (16.2) 18–84	–

Note: The total sample size, number of cmsTBI and control participants, male and female participants, average age (and standard deviation), average time since injury (TSI; in years, and standard deviation), and range of TSI are shown for each cohort.

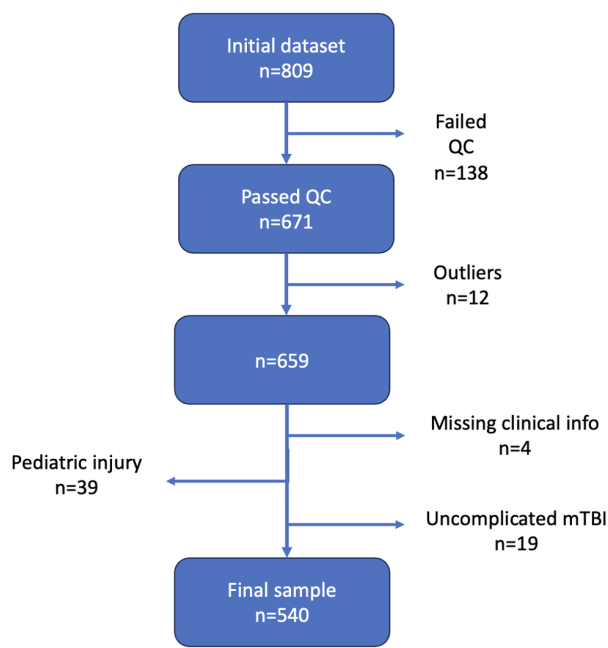


FIGURE 1: Participant exclusion flowchart. The number of participants included and various reasons for exclusion. This information is also detailed by site in Supplementary Table S1.

and/or significant loss-of-consciousness (for details see below). Complicated mild TBI has been shown to result in more severe and chronic cognitive deficits compared to mild TBI/concussion, constituting a distinct class of injury,²⁷ and was therefore included in the analysis. As injury severity was operationalized differently across the parent studies, patients were reclassified into complicated mild, moderate, or severe TBI based on GCS (Glasgow Coma Scale) score, where available: (1) GCS 13–15 and trauma-related intracranial pathology = complicated mild TBI, (2) GCS 9–12 = moderate TBI, and (3) GCS 3–8 = severe TBI. Where GCS scores were not available, injury severity was determined by other available study specific procedures (see Supplementary Table S1) or inferred based on inclusion/exclusion criteria. Severity was coded as 1 = complicated mild TBI, 2 = moderate TBI, 3 = severe TBI. Some studies excluded individuals with a history of more than 1 TBI, while others did not: see Supplementary Table S1.

Image Acquisition

All cohorts shared their raw T1-weighted MRI data with the central processing site (University of Utah, ED). The acquisition parameters for each cohort are shown in Supplementary Table S2.

Cognitive Data

Most sites collected a version of the Trail Making Test (TMT) and Digit Span from their cohorts; therefore, we examined whether TMT condition A and B (or D-KEFS conditions

3 and 4) performance or Digit Span scores (forward + backward) was associated with PAD in the cmsTBI group. For TMT, using raw or scaled scores together was not appropriate given differences in Halstead Trails and D-KEFS Trails test administration and norming procedures. For this reason, we normed the data based on our healthy control subjects to calculate *T*-scores, separately for each test. Cognitive data were collected at the same time point as the imaging data analyzed.

Brain Age Prediction

We implemented the Gaussian processes regression approach for brain age estimation.¹⁵ Raw T1-weighted MR images were processed with the brainageR v2.1 workflow (<https://github.com/james-cole/brainageR>), with a model similar to those described previously.^{13,15} The brainageR model was trained on brain MRIs from 3,377 individuals from 7 publicly available datasets. Overall, this training sample included individuals 18–92 years old from samples in the United States, United Kingdom, Australia, and China. Briefly, T1-weighted images were segmented into gray matter, white matter, and cerebrospinal fluid using SPM12 (<https://www.fil.ion.ucl.ac.uk/spm/software/spm12/>) and spatially normalized. The resulting images were vectorized and subjected to principal components analysis (using R *prcomp* <https://cran.r-project.org>), where components explaining the top 80% of variance were retained, resulting in 435 components. The brainageR model was trained using mean absolute error (MAE) of *y* (i.e., age) as the loss function. The kernel used was the radial basis function. All other GPR parameters were set as the default in kernlab (<https://cran.r-project.org/web/packages/kernlab/kernlab.pdf>), except for the initial noise variance, which was determined using nested cross-validation for hyperparameter tuning (comparing values of: 0.001, 0.01, 0.1, 1, 10), which gave 0.001 as the optimal value that was used for final training. Processing for the ENIGMA AMS-TBI data was the same, with raw T1-weighted images segmented, normalized, vectorized, and the rotation matrix from the training dataset applied to yield 435 components for each participant. The resulting components were used to predict brain age using *kernlab*,²⁸ and tissue segmentations were visually checked for quality. Of the 809 scans across 7 cohorts, 17% failed visual quality control due to poor tissue segmentation (QC, *N* = 138). The failure rate was similar between the cmsTBI and control groups. Outliers based on predicted age difference (PAD; $\pm 3SD$) were removed (*N* = 12). A flowchart of reasons for exclusions may be seen in Figure 1 and site-level information in Supplementary Table S3. Due to missing demographic or clinical information for some participants, the final sample size was 540 participants (343 cmsTBI and 197 controls).

The variable of interest was PAD, calculated by subtracting the chronological age from the predicted age. A negative PAD score indicates brain age values that are lower (younger) than expected given an individual's chronological age. A positive PAD indicates a brain that appears older than expected, and could imply either advanced and/or accelerated aging. Plots of the chronological age and predicted brain age across cohorts are shown in Supplementary Figure S1. For the purposes of this paper, *advanced* brain aging refers to a larger PAD in TBI, while *accelerated* brain aging refers to a PAD that increases with more advanced chronological age or more time post-injury.

APOE Analyses

A total of 166 participants (128 cmsTBI) across 3 cohorts had available APOE genotype. Of 166 participants with APOE genotype, 56 had at least 1 $\epsilon 4$ allele. Due to the limited APOE sample size, all findings related to APOE were considered exploratory and should be interpreted with caution.

Statistical Analyses

Statistical analyses were run as mixed effects models in R 3.1.3 with the *nlme* package, setting PAD as the dependent variable. Some cohorts consisted of participants from multiple studies or sites. Nested random effects (intercepts) were used to control for cohort and site/study. A flowchart for the statistical models tested may be found in Supplementary Figure S2. Normalized residuals, accounting for age, sex, and random effects of cohort and site, were calculated from regression analyses and used for charting.

Demographic Variables

As a first test of model accuracy, we examined the correlation between predicted brain age and chronological age in the healthy control sample. The model accurately predicted chronological age in healthy individuals ($r = 0.92$, $MAE = 1.4$). Across the whole sample, a significant correlation between age and PAD was found ($r = -0.2$, $p < 0.001$), meaning that PAD was higher for younger participants. The negative association between age and PAD has been shown in numerous papers and may result when there is insufficient information to estimate brain age while attempting to minimize residuals, resulting in regression to the mean/median.²⁹ There was not a significant sex difference in PAD ($t(538) = -0.4$, $p = 0.66$). Both age and sex were included as covariates in the models. Last, we examined the association between PAD and years of education, which was non-significant ($t(526) = -1.9$, $p = .06$).

Primary Group Comparison

We examined differences in PAD between the cmsTBI and control group, covarying for chronological age and sex. We also compared controls to TBI groups with patients broken into severity categories: cmTBI, modTBI, and sevTBI. All subsequent analyses were considered *post hoc* following a significant primary group effect.

Sensitivity Analyses

Several additional sensitivity analyses were run. First, we excluded cmsTBI participants with lesions visible on the T1-weighted image ($n = 152$) and those who were scanned < 1 year post-injury ($n = 51$). The rationale for the former is that lesions could lead to errors in the processing pipeline, and we wanted to ensure results were not due to such bias. The rationale for the latter is that this is a dynamic period during which most recovery occurs, and there may be diaschisis-related atrophy that is distinct from the more long-term interaction between aging and TSI.³⁰ We also ran analyses excluding individuals over 60 (age at scan, $n = 94$) to check a sample with better age matching between patients and controls. Additionally, we ran analyses excluding sites that *did not exclude* on the basis of prior TBI, thus limiting analyses to only participants with 1 recorded TBI.

Within-TBI-Group Analyses

We examined associations between PAD and a number of demographic, clinical, and cognitive variables within the cmsTBI group.

TSI. To examine a potential accelerated aging effect using cross-sectional data, we examined whether PAD remained consistent over TSI. We conceptualize accelerated aging as changing (advancing PAD) with increasing years since the time of injury. We tested linear and nonlinear association with PAD across all cmsTBI participants with TSI > 1 year, TSI > 5 years, and TSI > 10 years. To determine the shape of nonlinear associations, we performed spline interpolation with the *gam* function in the *splines* R library, testing 3, 5, and 7 degrees of freedom. We tested this association in these distinct windows of TSI as we would expect that detectable evidence of accelerated aging may not emerge for a few years after injury. *That is, if the pathological consequences of TBI (proteinopathy, inflammation) are active in chronic TBI, these interacting and perhaps cumulative effects should be more evident with longer windows of TSI.*

Age. Chronological age and TSI are related, making it difficult to tease apart the individual effects of each on PAD, especially given the known association between

chronological age and PAD. In addition, injury severity and age-at-injury are related (Supplementary Fig S3). To address this, we examined associations with TSI within separate age brackets, and associations with age within different TSI brackets.

Cognitive Performance. We examined associations with cognitive function, specifically performance on the TMT task (measured using either Delis-Kaplan Executive Function System [D-KEFS] or Halstead-Reitan Neuropsychological Battery [HRNB]; harmonization described above) and Digit Span. There were 175 participants in the cmsTBI group with Digit Span data (Digit Span Forward and Backward – DSF and DSB) and 203 with TMT (81 with D-KEFS and 122 with HRNB, normed separately using 95 controls with D-KEFS and 72 with HRNB). These analyses were adjusted for age and sex.

Interactions

We tested the following interactions: group \times age, group \times sex, and within the cmsTBI group, age \times TSI, TSI \times sex, and TSI \times education.

APOE Analyses

We compared PAD between participants negative for $\epsilon 4$ alleles and individuals either homo- or heterozygous for the allele across the whole sample and within the cmsTBI group only.

Defining Survivor Bias

Survivor bias is a pervasive methodological issue that faces any research efforts examining disease and mortality. This can result in research recruitment of an artificially healthy sample with respect to brain and behavioral health. Given the link between education and health, longevity, and mortality,^{31,32} we examined higher education in our older cohorts as a proxy for survivor bias.

Results

Primary Group Comparison

Across 540 individuals, the cmsTBI group had a significantly larger PAD than the control group ($b = 4.99$, $p < 0.001$, Fig 2A), indicating a substantial deviation (5 years) between brain age and chronological age with older appearing brains (*advanced aging*) in cmsTBI. There was a significant association between PAD and injury severity group, with more severe injury being associated with greater PAD ($b = 1.51$, $p = 0.019$, Fig 2B). Separated by severity, the cmsTBI group showed the smallest group difference (cmsTBI: $N = 288$, $b = 2.3$, $p = 0.055$; modTBI: $N = 248$, $b = 4.4$, $p \leq 0.001$; sevTBI: $N = 304$, $b = 5.8$, $p \leq 0.001$).

Sensitivity Analyses. The group difference in PAD remained after repeating the analysis while excluding participants with lesions visible on T1 scans ($N = 382$, $b = 4.55$, $p = 6.6 \times 10^{-8}$) and excluding participants < 1 year post-injury ($N = 476$, $b = 5.02$, $p = 3.7 \times 10^{-10}$). Group differences were larger when individuals > 60 years (age-at-scan) were excluded ($N = 458$, $b = 5.95$, $p = 8.5 \times 10^{-12}$) and slightly smaller when patients injured as children were included ($N = 515$, $b = 4.69$, $p = 1.3 \times 10^{-9}$). Additionally, we ran the group analysis covarying for time since injury and age at injury, separately, and results were consistent ($N = 527$, $b = 4.4$, $p < 0.001$; $N = 522$, $b = 4.5$, $p < 0.001$). Finally, when excluding sites that may have included individuals with multiple TBIs, results were consistent ($N = 337$, $b = 6.7$, $p < 0.001$).

Within-TBI Analyses

TSI. Across all cmsTBI participants more than a year post-injury, there was no linear association between TSI and PAD ($N = 276$, $b = 0.14$, $p = 0.10$), but there was a nonlinear association ($b = 0.02$, $p = 0.04$). There was however, a positive linear association between TSI and PAD across cmsTBI participants more than 5 years post-injury ($N = 109$, $b = 0.25$, $p = 0.02$) and a non-significant trend in participants > 10 years post-injury ($N = 58$, $b = 0.34$, $p = 0.057$), but no nonlinear associations in either of these age brackets (Fig 3). Fitting natural splines with 3, 5, or 7 degrees of freedom (DF), we found that both the 3 and 5 DF spline models were significant ($ps < .05$) with virtually identical model fit (based on Akaike Information Criterion). These curves exhibit an initial increase in brain age, followed by a decrease, followed by a slow and continuous increase (Fig 3).

Age. The positive association between TSI and PAD did not hold for the individuals who were older at age-of-scan, in fact it was reversed (Supplementary Fig S4). This inversion, starting around age 65, suggests that survivor bias may have influenced brain age results in the oldest individuals. There are many factors that can influence longevity, most of which were not accessible with the data available, with the exception of education. This possibility of survivor bias in our data is supported by the greater portion of older individuals (> 60 years at age-at-injury) who were highly educated (Bachelor's degree and up, ISCED level 6+) ($\chi^2 [6, N = 518] = 14.2$, $p = 0.028$; Supplementary Fig S7 and Table S4). Education is a well-known proxy for cognitive reserve that may mitigate some aging effects.

Cognitive Performance. There were no significant associations between PAD and any of the cognitive tests included

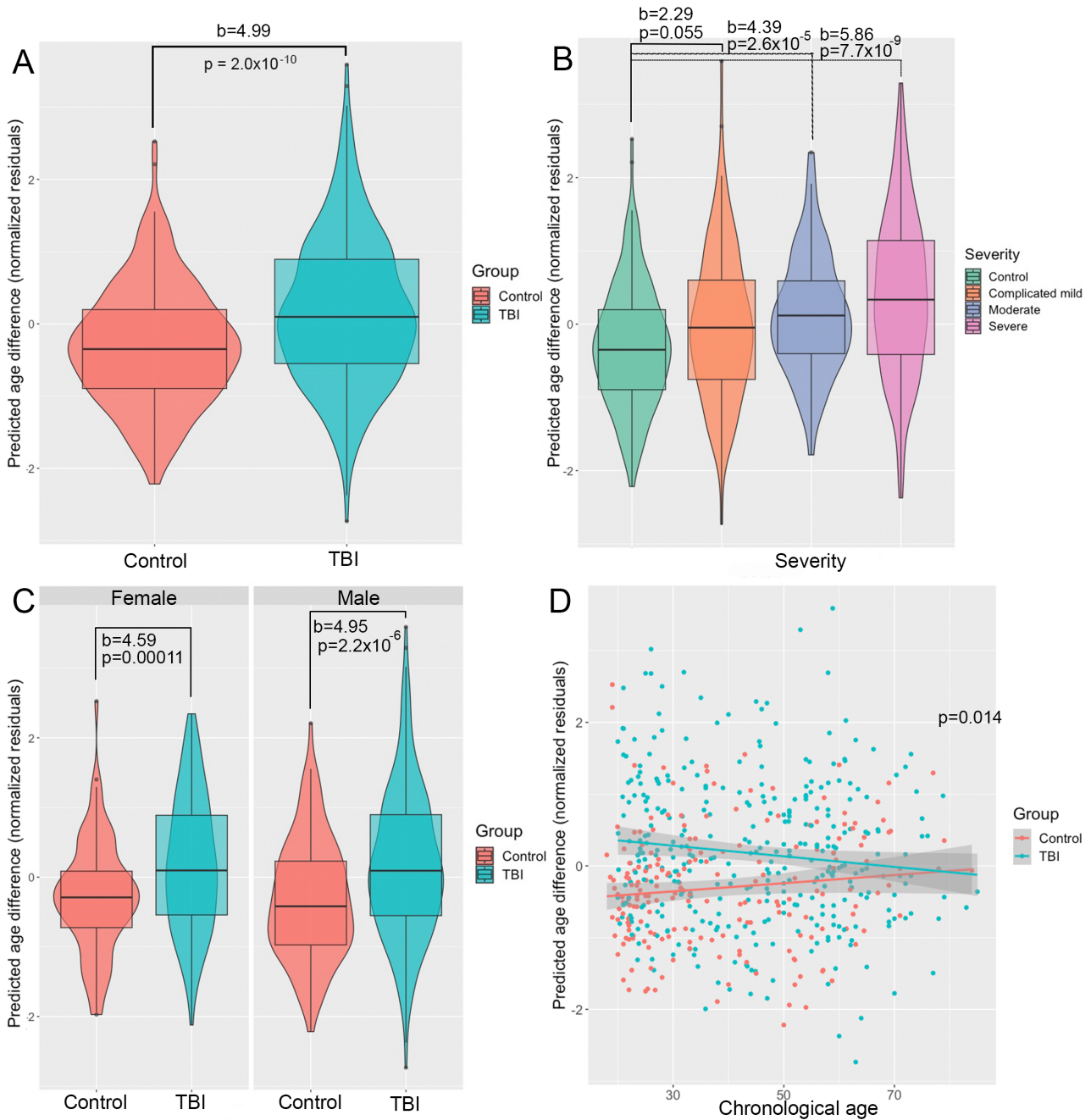


FIGURE 2: Group differences in PAD (A), associations with severity (B), group \times sex interaction (C), and group \times age interaction (D). Box/violin plots are shown for group differences in panels A–C, with control group in red and cmsTBI group in blue (in A, C, and D). Statistics are displayed for each panel. Trendlines in panel D are linear estimates with 95% confidence intervals in gray. The y axis displays normalized residuals – PAD after accounting for age, sex, and site effects.

across the cmsTBI group (DSF: $N = 175$, $p = 0.48$; DSB: $N = 175$, $p = 0.29$; TMT-A: $N = 173$, $p = 0.43$; TMT-B: $N = 173$, $p = 0.86$). These models were also non-significant when we covaried for education or tested associations across the full sample.

Interactions

There was a significant group \times sex interaction ($b = 1.63$, $p = 0.009$, Fig 2C), indicating that group

differences in PAD (cmsTBI vs Controls) were not the same between males and females. Group differences among females/women ($b = 4.59$, $N = 179$) were slightly smaller than among males/men ($b = 4.95$, $N = 297$). These remained significant after covarying for TSI or severity. There was also a significant group \times age interaction ($b = -0.11$, $p = 0.014$, Fig 2D), indicating that the group differences were not consistent over adulthood. In fact, group differences were smaller among older participants.

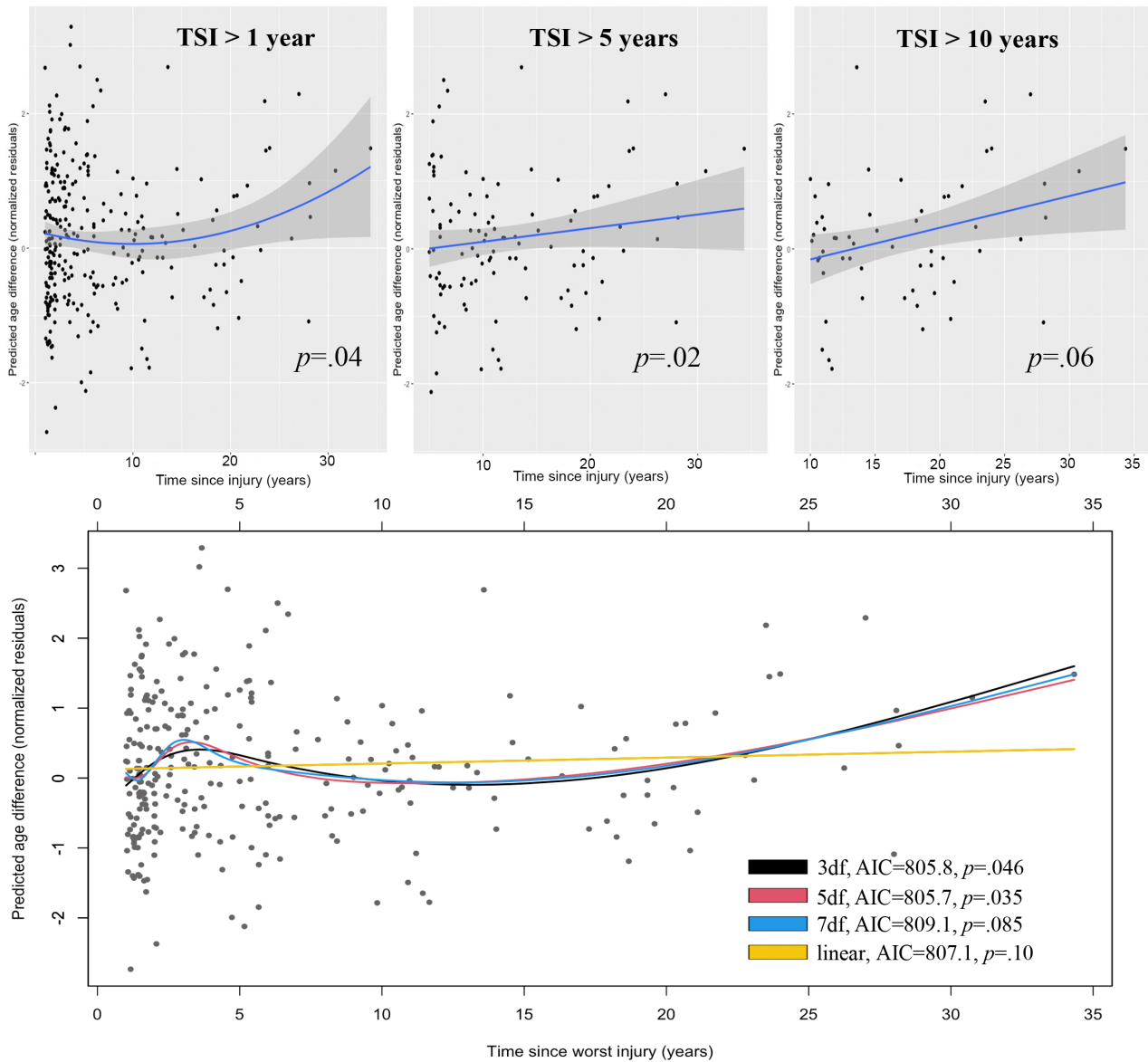


FIGURE 3: Association between PAD and TSI. The associations between time since injury in years and PAD (normalized residuals – accounting for age, sex, and random effects of cohort|site) are shown for 3 post-injury intervals (greater than 1 year, greater than 5 years, and greater than 10 years). Trendlines were plotted in R 4.2.2 with 95% confidence intervals in gray. There was a significant non-linear relationship with TSI across all cmsTBI participants more than 1 year post-injury and linear relationships with TSI beginning at 5 and 10 years post-injury. Given the significant nonlinear association, we also added a spline interpolation to better understand the shape. Using the *gam* function in the *splines* R library, we fit a natural spline with 3, 5, and 7 DF. The model with 5 DF provided the best fit. The linear fit is also displayed in yellow for comparison. AIC = Akaike Information Criterion.

There were no significant interactions between chronological age × TSI, TSI × sex, or TSI × education in the cmsTBI group.

APOE Analyses

We hypothesized that individuals with at least 1 APOE ε4 allele would have greater PAD. The analysis included 56 individuals hetero- or homozygous for the ε4 allele and 110 individuals without an ε4 allele. The hypothesis was not supported; there was no significant difference in PAD based on

APOE genotype in the whole sample or within the cmsTBI group ($p = 0.64$ and $.48$, respectively). There were also no significant interactions between APOE × group or APOE × age ($p = 0.97$ and $.40$, respectively). With a relatively small sample size, we considered this analysis to be exploratory.

Discussion

Our goal was to examine the effects of remote cmsTBI on brain aging. In a mega-analysis of 540 individuals

from 7 cohorts, we show evidence of accelerated brain aging after cmsTBI. Individuals with cmsTBI showed a PAD 5 years greater than the control group, consistent with other data in cmsTBI.^{15,17} These analyses extend prior work in 3 ways. First, the current sample included a large number of younger and older adults with a range of post-injury intervals. While prior work has focused on young to middle aged adults in the first few years after injury (e.g., mean 28 months post-injury¹⁵) or older adults in the late chronic timepoint (over 10 years post-injury),¹⁷ the median age for this study was 41.2 (range 18–85) and a mean TSI of 5.3 ± 6.4 years (range 0.2–34 years) with 198 participants at least 50 years of age (age-at-scan, mean TSI 7.5 ± 7.4 years). Second, there have been mixed findings with respect to the effects of injury severity and brain age, which are addressed below. Finally, in view of the established role of sex in TBI recovery, the current sample size allowed us to examine potential sex differences.

Brain Age and TSI

The positive association between TSI and PAD supports an accelerated aging effect, with the most dramatic effects observed after a decade post-injury (Fig 3). These data are consistent with prior work and extend those findings to a longer post-injury period. Spline interpolation revealed a pattern of initial injury response and an upward inflection in brain age (1–3 years in our data), followed by a slight decrease in brain age, perhaps reflecting injury accommodation and compensation, followed by a progressing increase in PAD around 7 years post-injury. While the year markers reported are biased by the TSI distribution in our sample (weighted toward TSI < 10 years), we present the general shape and progression, which mirrors a biphasic response to injury observed in other measurements of TBI pathology including serum protein concentration.³³

TAI is a strong predictor for brain atrophy, and more recently Wallerian degeneration and propagation of proteinopathies has been demonstrated to continue into the chronic phase in humans.⁴ There also remain a number of possible interacting variables including blood–brain barrier disruption and neuroinflammation³⁴, or functional network changes that facilitate functional recovery in the acute stage but may promote proteinopathy over the long-term.³⁵ Our data are consistent with the idea that brain volumetric changes post-injury cannot be accounted for by lesion resolution and transsynaptic or Wallerian-like events alone, and are likely attributable to more insidious physiological processes post-injury. While brain age may serve as a sensitive marker for brain health, future research efforts should clarify the clinical significance of this

variable, establish the critical windows when cortical atrophy is occurring, and define the clinical and demographic modifiers of post-traumatic increase in PAD as well as its underlying mechanisms.

Injury Severity and PAD

In contrast to others,^{2,19,20} Cole and colleagues found little evidence of PAD in a single uncomplicated mild TBI event. On the other hand, a pronounced effect has been seen in cmsTBI.¹⁵ With the benefit of a larger cmsTBI sample, the current results extend these findings, showing a stepwise increase in PAD with injury severity (Fig 3). This finding is intuitive and consistent with extensive literature supporting a direct relationship between injury severity and atrophy based on T1w-MRI volumetrics.³⁰

Sex and PAD

There is growing evidence that sex plays a critical role in TBI recovery and outcome.²⁴ The reasons for observed sex-based disparities in TBI outcome are a growing area of investigation, a welcome transition after decades of focus on male-only models for TBI.³⁶ While some evidence points to hormonal differences as a factor contributing,³⁷ mechanisms are still poorly understood. One aspect complicating sex comparisons is that the mechanism of injury may differ in ways that have implications for outcome.³⁸ With regard to sex and brain morphometry, recent work in mild TBI revealed no differences.³⁹ Our findings point to modestly greater PAD in men, although the effect was small (Fig 2C).

Brain Age and Genetics

Contrary to our hypotheses, APOE $\epsilon 4$ did not have a greater PAD compared to non-carriers. Of note, the APOE literature is mixed, with some findings revealing a clear effect of APOE on brain volumetrics,⁴⁰ while others show no relationship.⁴¹ In 1 recent examination of over 1,100 individuals, examiners showed a differential effect of APOE on brain volume with the $\epsilon 4$ allele based on age, i.e., a paradoxical “protective” effect in individuals <60 years of age⁴² that reversed in older age groups. The null finding in the present data could *not* be explained by greater injury severity in the non-carriers or sex differences between subgroups. This null finding should be interpreted cautiously due to limited statistical power due to the small sample size, but may also point to limitations in brain structure-based analysis in assessing functional changes; where brain organization, including functional connectivity, may better approximate neurological resilience.⁴³ However, this null finding is consistent with a large genetic association study in over 4,700 patients with

TBI, which did not replicate the effect of APOE $\epsilon 4$ carrier status on outcome.⁴⁴

Brain Age and Behavioral Outcome

We observed no significant relationship between the accelerating PAD in cmsTBI and cognitive functioning, i.e., psychomotor speed, executive functioning, and working memory. This finding is inconsistent with some prior work,^{2,15} but generally in line with a long history of incongruence between behavior and brain structure,⁴⁵ given the numerous factors responsible for change in brain volume and for behavioral decline, including the role of neural and cognitive reserves. Therefore, while we did not find cognitive consequences for advancing PAD, this negative result may be simply due to the fact that neural networks deteriorate prior to the appearance of clinical changes. In support of this, a recent study showed PAD to be predictive of later progression to dementia in a typical aging sample.⁴⁶

Addressing Challenges to External Validity

The study of cmsTBI over the lifespan poses natural challenges to study design and subject enrollment. Subject recruitment in cmsTBI has multiple challenges, including sample bias with regard to socioeconomic status, sex, and race.⁴⁷ Survivor bias can also impact studies of illness, aging, and mortality, and may amplify counterintuitive effects.⁴⁸

In our data, survivor bias may have contributed to the finding that PAD decreases with greater TSI in the oldest individuals (>65 years of age). Survivor bias is often difficult to track and demonstrate, but we used uncharacteristic differences in the sample as markers for potential biased sampling. Education is a predictor of longevity, and a proxy for cognitive reserve, which may serve to increase resilience post-cmsTBI.⁴⁹ In our data, there were more individuals with advanced degrees (Bachelor's and higher) beginning at age 60. One explanation for this upward inflection in sample education is age-related attrition in less educated individuals (e.g., due to mortality and dementia⁵⁰) and greater health and resilience factors associated with advanced education, which is a modifier of MRI brain volume.⁵¹ *Aging with TBI* is distinct from *TBI during aging*, partly because TBIs sustained in older individuals are more likely due to falls (lower impact) than motor vehicles (higher impact) producing different brain pathology. Moreover, individuals who sustain a *TBI during aging (older age-at-injury)*, will typically have completed their educational goals and potential, whereas those who are *aging with TBI (younger age-at-injury)* may have their educational and career trajectories negatively altered by TBI.

Cross-sectional research addressing questions that are by nature developmental or evolve over a lifespan have significant limitations, and is the most important limitation of our study. To ideally address the goals of the current study, large datasets with longitudinal data spanning decades are needed. We also recognize that a majority of the sample was from participants of white-European descent, and there is a great need to determine the role of health care access, race, and socioeconomic status on outcomes.⁴⁷ This is particularly true given that non-whites are less likely to be represented in the research literature,⁴⁷ have poorer clinical outcomes post-TBI,⁴⁷ and potentially carry a higher risk for Alzheimer's disease and related dementias.⁵² Our analyses are based on T1-weighted MRI data only, while other modalities have been shown to have greater sensitivity to the neuropathology of TBI.^{53,54} Future studies and algorithms could leverage information from multi-modal data to more fully assess brain alterations due to TBI.⁵⁵ Finally, while the methodology chosen in this study to determine brain age is well validated,^{15,16} there are multiple additional brain age algorithms that could also be used, some of which provide high-dimensional whole brain data allowing for more anatomical interpretation.¹⁰

Conclusions

The current mega-analysis of cross-sectional data reveals that a single cmsTBI is associated with a PAD of nearly 5 years along with *accelerated aging* years after injury. We show a non-linear time course for the initial effects of injury, followed by relative stability, and then faster progression of brain age in the late chronic phase. Injury severity showed a stepwise relationship with advancing brain age. There was also a mild influence of sex, with men showing relatively larger PAD. Longitudinal designs are needed to assess disease progression or mitigation. Brain age holds promise as a useful biomarker to track changes over time due to its dynamic nature and amenability to modification with preventive measures (e.g., lifestyle adjustments) and, possibly, treatment.

Acknowledgments

NIH/NINDS R61NS120249 to Dr. Dennis. Dr. Menon is an Emeritus Senior Investigator of the National Institute for Health Research, UK, and is supported by the CENTER-TBI grant from the European Union and the TBI-REPORTER grant from UKRI, NIHR, AR UK and UK Ministry of Defense. Israel Innovation Authority, "Magnetron" Grant to Dr Livny. Maryland Technology Development Corporation 1 R01 EY028039 Congressionally

Directed Medical Research, JH Discovery Award to Dr. Koliatsos. National Institute for Health and Care Research (NIHR) Advanced Fellowship to Dr. Newcombe. NHMRC Early Career Fellowship to Dr. Spitz. NIH grants R01NS100973, R01AG079957 and RF1AG082201, DoD contract W81XWH-18-1-0413, an anonymous donor family, and the Hanson-Thorell Research Scholarship Fund at the University of Southern California to Dr. Irimia. NIH/NIA R01AG061028, NIH/NINDS RF1NS115268, NIH/NINDS RF1NS128961 and NIH/NINDS/NICHD1U01NS086625-01 to Dr. Dams-O'Connor. NIH/NINDS R01-NS121107 to Dr. Dobryakova. PA Health Research Grant SAP #4100077082 to Dr. Hillary. The European Research Council (ERC) under the European Union's Horizon 2020 research and innovation programme (ERC Starting Grant, Grant agreement No. 802998) and the Research Council of Norway (249795) to Dr. Westlye. To Dr. Caeyenberghs: The Victorian Near-miss Award Pilot is administered by veski for the Victorian Health and Medical Research Workforce Project on behalf of the Victorian Government and the Association of Australian Medical Research Institutes. Funding for the Pilot has been provided by the Victorian Department of Jobs, Precincts and Regions. The Liaison Committee between the Central Norway Regional Health Authority (RHA) and the Norwegian University of Science and Technology (NTNU) (2020/39645) to Dr. Olsen. Tiny Blue Dot Foundation to Dr. Monti. Transport Accident Commission to Dr. Ponsford. NIH U54EB020403, R01MH116147, R56AG058854, P41EB015922, R01MH11671 to Dr. Thompson. VA R&D SPIRE AWARD.

Author Contributions

E.L.D., M.M.A., H.A., E.D.B., K.C., J.H.C., K.D.O.C., E.M.D., E.D., H.M.G., J.H.G., A.K.H., T.H., A.I., V.E.K., H.M.L., A.L., D.K.M., T.L.M., A.Z.M., S.M., M.M.M., V.F.J.N., M.R.N., J.P., H.S., G.S., L.T.W., R.Z., P.M.T., E.A.W., A.O., and F.G.H. contributed to the conception and design of the study. E.L.D., S.V., M.M.A., K.C., J.H.C., K.D.O.C., E.D., H.M.G., T.H., A.R., G.S., U.V., L.T.W., A.O., and F.G.H. contributed to the acquisition and analysis of data. E.L.D., A.O., and F.G.H. contributed to drafting the text of preparing the figures.

Potential Conflicts of Interest

Dr. Olsen is a co-founder and owner of Nordic Brain Tech AS. Dr. Zafonte received royalties from: (1) Oakstone for an educational CD- Physical Medicine and Rehabilitation a Comprehensive Review; (2) Demos

publishing for serving as co- editor of the text Brain Injury Medicine. Dr Zafonte serves on the Scientific Advisory Board of Myomo, Oxeia Biopharma, Biodirection and ELMINDA. He also evaluates patients in the MGH Brain and Body-TRUST Program which is funded by the NFL Players Association. Dr. Adamson is the CEO & Founder of Soof Solutions Inc. Dr. Cole is a scientific advisor and shareholder of Claritas HealthTech PTE and of BrainKey. Dr. Thompson received partial research support from Biogen, Inc., for research unrelated to this manuscript.

Data Availability

Data are available to researchers who join the working group and submit a secondary analysis proposal to the group for approval, which is granted on a cohort-by-cohort level. Interested researchers should contact the corresponding authors.

References

- Graham NS, Sharp DJ. Understanding neurodegeneration after traumatic brain injury: from mechanisms to clinical trials in dementia. *J Neurol Neurosurg Psychiatry* 2019;90:1221–1233.
- Amgalan A, Maher AS, Ghosh S, et al. Brain age estimation reveals older adults' accelerated senescence after traumatic brain injury. *Geroscience* 2022;44:2509–2525.
- Povlishock JT, Katz DI. Update of neuropathology and neurological recovery after traumatic brain injury. *J Head Trauma Rehabil* 2005; 20:76–94.
- Graham NSN, Jolly A, Zimmerman K, et al. Diffuse axonal injury predicts neurodegeneration after moderate-severe traumatic brain injury. *Brain* 2020;143:3685–3698.
- Graham NSN, Cole JH, Bourke NJ, et al. Distinct patterns of neurodegeneration after TBI and in Alzheimer's disease. *Alzheimers Dement* 2023;19:3065–3077.
- Lu Y, Jarrahi A, Moore N, et al. Inflammaging, cellular senescence, and cognitive aging after traumatic brain injury. *Neurobiol Dis* 2023; 180:106090.
- Huang X, You W, Zhu Y, et al. Microglia: a potential drug target for traumatic axonal injury. *Neural Plast* 2021;2021:5554824.
- Ayubcha C, Revheim M-E, Newberg A, et al. A critical review of radiotracers in the positron emission tomography imaging of traumatic brain injury: FDG, tau, and amyloid imaging in mild traumatic brain injury and chronic traumatic encephalopathy. *Eur J Nucl Med Mol Imaging* 2021;48:623–641.
- Moretti L, Cristofori I, Weaver SM, et al. Cognitive decline in older adults with a history of traumatic brain injury. *Lancet Neurol* 2012;11: 1103–1112.
- Yin C, Imms P, Cheng M, et al. Anatomically interpretable deep learning of brain age captures domain-specific cognitive impairment. *Proc Natl Acad Sci U S A* 2023;120:e2214634120.
- Franke K, Gaser C. Ten years of BrainAGE as a neuroimaging biomarker of brain aging: what insights have we gained? *Front Neurol* 2019;10:789.
- Han LKM, Dinga R, Leenings R, et al. A large-scale ENIGMA multisite replication study of brain age in depression. *NeuroImage* 2022;2: 100149.

13. Clausen AN, Fercho KA, Monsour M, et al. Assessment of brain age in posttraumatic stress disorder: findings from the ENIGMA PTSD and brain age working groups. *Brain Behav* 2022;12:e2413.
14. Liew S-L, Schweighofer N, Cole JH, et al. Association of Brain age, lesion volume, and functional outcome in patients with stroke. *Neurology* 2023;100:e2103–e2113. <https://doi.org/10.1212/WNL.000000000207219>.
15. Cole JH, Leech R, Sharp DJ. Alzheimer's disease neuroimaging initiative. Prediction of brain age suggests accelerated atrophy after traumatic brain injury. *Ann Neurol* 2015;77:571–581.
16. Cole JH, Jolly A, de Simoni S, et al. Spatial patterns of progressive brain volume loss after moderate-severe traumatic brain injury. *Brain* 2018;141:822–836.
17. Spitz G, Hicks AJ, Roberts C, et al. Brain age in chronic traumatic brain injury. *Neuroimage Clin* 2022;35:103039.
18. Newcombe VFJ, Ashton NJ, Posti JP, et al. Post-acute blood biomarkers and disease progression in traumatic brain injury. *Brain* 2022;145:2064–2076.
19. Dennis EL, Taylor BA, Newsome MR, et al. Advanced brain age in deployment-related traumatic brain injury: a LIMBIC-CENC neuroimaging study. *Brain Inj* 2022;36:662–672.
20. Gan S, Shi W, Wang S, et al. Accelerated brain aging in mild traumatic brain injury: longitudinal pattern recognition with White matter integrity. *J Neurotrauma* 2021;38:2549–2559.
21. Poudel GR, Dominguez DJF, Verhelst H, et al. Network diffusion modeling predicts neurodegeneration in traumatic brain injury. *Ann Clin Transl Neurol* 2020;7:270–279.
22. Thompson PM, Jahanshad N, Ching CRK, et al. ENIGMA and global neuroscience: a decade of large-scale studies of the brain in health and disease across more than 40 countries. *Transl Psychiatry* 2020;10:100.
23. Olsen A, Babikian T, Bigler E, et al. Toward a global and reproducible science for brain imaging in neurotrauma: the ENIGMA adult moderate/severe traumatic brain injury working group. *Br Imaging Behav* 2019;15:526–554. Under Review.
24. Gupte R, Brooks W, Vukas R, et al. Sex differences in traumatic brain injury: what we know and what we should know. *J Neurotrauma* 2019;36:3063–3091.
25. Maiti TK, Konar S, Bir S, et al. Role of apolipoprotein E polymorphism as a prognostic marker in traumatic brain injury and neurodegenerative disease: a critical review. *Neurosurg Focus* 2015;39:E3.
26. For Statistics UI. International standard classification of education: ISCED 2011. *Comp Soc Res* 2012;30. Available from: <https://www.voced.edu.au/content/ngv:54992>
27. Hacker D, Jones CA, Yasin E, et al. Cognitive outcome after complicated mild traumatic brain injury: a literature review and meta-analysis. *J Neurotrauma* 2023;40:1995–2014. <https://doi.org/10.1089/neu.2023.0020>.
28. Karatzoglou A, Smola A, Hornik K, Zeileis A. Kernlab – an S4 package for kernel methods in R. *J Stat Softw* 2004;11:1–20. [cited 2021 Mar 18].
29. de Lange A-MG, Anatürk M, Rokicki J, et al. Mind the gap: performance metric evaluation in brain-age prediction. *Hum Brain Mapp* 2022;43:3113–3129.
30. Bigler ED. Distinguished neuropsychologist Award lecture 1999. The lesion(s) in traumatic brain injury: implications for clinical neuropsychology. *Arch Clin Neuropsychol* 2001;16:95–131.
31. McHutchison CA, Backhouse EV, Cvoro V, et al. Education, socioeconomic status, and intelligence in childhood and stroke risk in later life: a meta-analysis. *Epidemiology* 2017;28:608–618.
32. Deary IJ, Taylor MD, Hart CL, et al. Intergenerational social mobility and mid-life status attainment: influences of childhood intelligence, childhood social factors, and education. *Dermatol Int* 2005;33:455–472.
33. Shahim P, Politis A, van der Merwe A, et al. Time course and diagnostic utility of NFL, tau, GFAP, and UCH-L1 in subacute and chronic TBI. *Neurology* 2020;95:e623–e636.
34. Visser K, Koggel M, Blaauw J, et al. Blood-based biomarkers of inflammation in mild traumatic brain injury: a systematic review. *Neurosci Biobehav Rev* 2022;132:154–168.
35. Hillary FG, Grafman JH. Injured brains and adaptive networks: the benefits and costs of Hyperconnectivity. *Trends Cogn Sci* 2017;21:385–401.
36. Shansky RM, Murphy AZ. Considering sex as a biological variable will require a global shift in science culture. *Nat Neurosci* 2021;24:457–464.
37. Franke K, Hagemann G, Schleussner E, Gaser C. Changes of individual BrainAGE during the course of the menstrual cycle. *NeuroImage* 2015;115:1–6.
38. Mikolić A, van Klaveren D, Groeniger JO, et al. Differences between men and women in treatment and outcome after traumatic brain injury. *J Neurotrauma* 2021;38:235–251.
39. Shida AF, Massett RJ, Imms P, et al. Significant acceleration of regional brain aging and atrophy after mild traumatic brain injury. *J Gerontol A Biol Sci Med Sci* 2023;78:1328–1338. <https://doi.org/10.1093/gerona/glad079>.
40. Rogojin A, Gorbet DJ, Hawkins KM, Sergio LE. Differences in structural MRI and diffusion tensor imaging underlie visuomotor performance declines in older adults with an increased risk for Alzheimer's disease. *Front Aging Neurosci* 2022;14:1054516.
41. Vervoordt SM, Arnett P, Engeland C, et al. Depression associated with APOE status and hippocampal volume but not cognitive decline in older adults aging with traumatic brain injury. *Neuropsychology* 2021;35:863–875.
42. Tang M, Su N, Zhang D, et al. The differential effects of Apoε4 on cerebral volumetric structures in different lifespan in community-dwelling populations. *J Alzheimers Dis* 2023;92:691–700. <https://doi.org/10.3233/JAD-220834>.
43. Stern Y, Varangis E, Habeck C. A framework for identification of a resting-bold connectome associated with cognitive reserve. *NeuroImage* 2021;232:117875.
44. Kals M, Kunzmann K, Parodi L, et al. A genome-wide association study of outcome from traumatic brain injury. *EBioMedicine* 2022;77:103933.
45. Marek S, Tervo-Clemmens B, Calabro FJ, et al. Reproducible brain-wide association studies require thousands of individuals. *Nature* 2022;603:654–660.
46. Biondo F, Jewell A, Pritchard M, et al. Brain-age is associated with progression to dementia in memory clinic patients. *Neuroimage Clin* 2022;36:103175.
47. Brenner EK, Grossner EC, Johnson BN, et al. Race and ethnicity considerations in traumatic brain injury research: incidence, reporting, and outcome. *Brain Inj* 2020;34:799–808.
48. Mitchell E, Chohan H, Bestwick JP, Noyce AJ. Alcohol and Parkinson's disease: a systematic review and meta-analysis. *J Parkinsons Dis* 2022;12:2369–2381.
49. Stern Y, Barnes CA, Grady C, et al. Brain reserve, cognitive reserve, compensation, and maintenance: operationalization, validity, and mechanisms of cognitive resilience. *Neurobiol Aging* 2019;83:124–129.
50. Fuller GW, Ransom J, Mandrekar J, Brown AW. Long-term survival following traumatic brain injury: a population-based parametric survival analysis. *Neuroepidemiology* 2016;47:1–10.
51. Steffener J. Education and age-related differences in cortical thickness and volume across the lifespan. *Neurobiol Aging* 2021;102:102–110.

52. Barnes LL. Alzheimer disease in African American individuals: increased incidence or not enough data? *Nat Rev Neurol* 2022;18:56–62.
53. Mckee AC, Daneshvar DH. The neuropathology of traumatic brain injury. *Handb Clin Neurol* 2015;127:45–66.
54. Bigler ED, Maxwell WL. Neuroimaging and neuropathology of TBI. *NeuroRehabilitation* 2011;28:63–74.
55. Niu X, Zhang F, Kounios J, Liang H. Improved prediction of brain age using multimodal neuroimaging data. *Hum Brain Mapp* 2020;41:1626–1643.