JAMA Neurology | Original Investigation

Neuropathologic and Clinical Findings in Young Contact Sport Athletes Exposed to Repetitive Head Impacts

Ann C. McKee, MD; Jesse Mez, MD, MS; Bobak Abdolmohammadi, BA; Morgane Butler, BSc; Bertrand Russell Huber, MD, PhD; Madeline Uretsky, MS; Katharine Babcock, PhD; Jonathan D. Cherry, PhD; Victor E. Alvarez, MD; Brett Martin, MS; Yorghos Tripodis, PhD; Joseph N. Palmisano, MS; Kerry A. Cormier, BA; Caroline A. Kubilus, BA; Raymond Nicks, MS; Daniel Kirsch, BA; Ian Mahar, PhD; Lisa McHale, EdS; Christopher Nowinski, PhD; Robert C. Cantu, MD; Robert A. Stern, PhD; Daniel Daneshvar, MD, PhD; Lee E. Goldstein, MD, PhD; Douglas I. Katz, MD; Neil W. Kowall, MD; Brigid Dwyer, MD; Thor D. Stein, MD, PhD; Michael L. Alosco, PhD

IMPORTANCE Young contact sport athletes may be at risk for long-term neuropathologic disorders, including chronic traumatic encephalopathy (CTE).

OBJECTIVE To characterize the neuropathologic and clinical symptoms of young brain donors who were contact sport athletes.

DESIGN, SETTING, AND PARTICIPANTS This case series analyzes findings from 152 of 156 brain donors younger than 30 years identified through the Understanding Neurologic Injury and Traumatic Encephalopathy (UNITE) Brain Bank who donated their brains from February 1, 2008, to September 31, 2022. Neuropathologic evaluations, retrospective telephone clinical assessments, and online questionnaires with informants were performed blinded. Data analysis was conducted between August 2021 and June 2023.

EXPOSURES Repetitive head impacts from contact sports.

MAIN OUTCOMES AND MEASURES Gross and microscopic neuropathologic assessment, including diagnosis of CTE, based on defined diagnostic criteria; and informant-reported athletic history and informant-completed scales that assess cognitive symptoms, mood disturbances, and neurobehavioral dysregulation.

RESULTS Among the 152 deceased contact sports participants (mean [SD] age, 22.97 [4.31] years; 141 [92.8%] male) included in the study, CTE was diagnosed in 63 (41.4%; median [IQR] age, 26 [24-27] years). Of the 63 brain donors diagnosed with CTE, 60 (95.2%) were diagnosed with mild CTE (stages I or II). Brain donors who had CTE were more likely to be older (mean difference, 3.92 years; 95% CI, 2.74-5.10 years) Of the 63 athletes with CTE, 45 (71.4%) were men who played amateur sports, including American football, ice hockey, soccer, rugby, and wrestling; 1 woman with CTE played collegiate soccer. For those who played football, duration of playing career was significantly longer in those with vs without CTE (mean difference, 2.81 years; 95% CI, 1.15-4.48 years). Athletes with CTE had more ventricular dilatation, cavum septum pellucidum, thalamic notching, and perivascular pigment-laden macrophages in the frontal white matter than those without CTE. Cognitive and neurobehavioral symptoms were frequent among all brain donors. Suicide was the most common cause of death, followed by unintentional overdose; there were no differences in cause of death or clinical symptoms based on CTE status.

CONCLUSIONS AND RELEVANCE This case series found that young brain donors exposed to repetitive head impacts were highly symptomatic regardless of CTE status, and the causes of symptoms in this sample are likely multifactorial. Future studies that include young brain donors unexposed to repetitive head impacts are needed to clarify the association among exposure, white matter and microvascular pathologic findings, CTE, and clinical symptoms.

JAMA Neurol. doi:10.1001/jamaneurol.2023.2907 Published online August 28, 2023. Multimedia
Supplemental content

Author Affiliations: Author affiliations are listed at the end of this article.

Corresponding Author: Ann C. McKee, MD, Veterans Affairs Boston Healthcare System, US Department of Veteran Affairs, 150 S Huntington Ave, Boston, MA 02130 (amckee@bu.edu).

cross the world, millions of people are exposed to repetitive head impacts (RHIs) through participation in contact and collision sports, military service, physical violence, and many other activities.¹⁻⁶ Repetitive head impacts can result in symptomatic concussions and the much more frequent, nonconcussive injuries that are asymptomatic.⁷ Sustained exposure to RHIs can produce persistent cognitive and neuropsychiatric symptoms⁸⁻¹¹ and a progressive, tau-based neurodegenerative disease, chronic traumatic encephalopathy (CTE).¹²⁻²¹ Multiple studies13,15,22 link a longer duration of RHI exposure in US football players with increased odds for the presence of CTE and increased severity of CTE. In older American football players with pathologically diagnosed CTE, RHI exposure is also associated with white matter rarefaction,²³⁻²⁵ loss of myelin associated proteins,²⁶ and oligodendrocyte loss.²⁷ Emerging data show structural white matter alterations on magnetic resonance imaging (MRI) in young, active, and recently retired contact sport players exposed to RHI,^{1,2,28-30} although the pathologic condition underlying these changes is unclear.

A definitive diagnosis of CTE requires neuropathologic evidence of perivascular hyperphosphorylated tau (p-tau) aggregates in neurons, with or without astrocytes, typically at the depths of the sulci in the cerebral cortex.^{31,32} The clinical syndrome associated with CTE is known as traumatic encephalopathy syndrome (TES).^{8,33} On the basis of the National Institute of Neurological Disorders and Stroke (NINDS) consensus diagnostic criteria for TES,⁸ the core clinical features of TES include cognitive impairment, especially episodic memory and executive dysfunction, and neurobehavioral dysregulation, such as impulsivity, explosivity, and emotional dysregulation.⁸ Supportive features include delayed onset (ie, core clinical features starting years after RHI exposure ends), parkinsonism, other motor signs (including amyotrophic lateral sclerosis), depression, anxiety, apathy, and paranoia.

The Understanding Neurologic Injury and Traumatic Encephalopathy (UNITE) Brain Bank¹ has harvested brains from more than 1350 donors exposed to RHIs who are considered at risk for CTE. Brain donors in the UNITE bank vary widely in age at death and include teenagers and young adults. Chronic traumatic encephalopathy has been reported in individuals as young as 17 years,¹³ yet, to date, there have been no largescale neuropathologic and clinical studies of young individuals exposed to RHIs. Attention to this age group has several important implications. The study of young athletes allows insight into the earliest features of RHI-induced neuropathologic injury and CTE. Furthermore, it allows analysis in the absence of common age-associated comorbidities. Moreover, most of these young contact sport athletes played only at amateur levels, as part of teams affiliated with educational institutions; consequently, the study of young athletes adds to our understanding of the long-term consequences of amateur contact sports participation. In this report, we describe the neuropathologic and clinical features of 152 brain donors from the UNITE brain bank who were younger than 30 years at the time of death.

Key Points

Question What are the neuropathologic and clinical findings in a convenience sample of young, deceased, symptomatic contact sport athletes?

Findings In this case series of 152 contact sport athletes younger than 30 years at the time of death, chronic traumatic encephalopathy (CTE) was found in 63 (41.4%), with nearly all having mild CTE (stages I and II). Neuropathologic abnormalities associated with CTE included ventricular enlargement, cavum septum pellucidum, thalamic notching, and perivascular pigment-laden macrophage deposition in the frontal white matter.

Meaning These findings confirm that CTE and other brain pathologies can be found in young, symptomatic contact sport athletes, but the clinical correlates of these pathologic conditions are uncertain.

Methods

Brain Donors and Study Design

The initial sample for this case series included 156 deceased individuals with a history of exposure to RHIs from contact sports participation who donated their brain to the UNITE Brain Bank from February 1, 2008, to September 31, 2022, and were younger than 30 years at the time of death. Race was determined by next-of-kin report and was included to understand the representativeness of the sample and the associated generalizability of the results. Procedures of brain donation have been previously described.14,34 The inclusion criterion was based on the presence of a history of exposure to RHIs without regard to symptom status. The restriction to brain donors younger than 30 years was selected to minimize any contribution from age-related conditions. Donors were excluded for poor tissue quality. Four donors were excluded because of incomplete brain fragments or prolonged premortem hypoxia, resulting in a final sample size of 152. Institutional review board approval for brain donation, postmortem clinical record review, interviews with informants, and neuropathologic evaluation was obtained through the Boston University Medical Campus and the Veterans Affairs Bedford Institutional Review Board. The next of kin or legally authorized representative of each brain donor provided written informed consent. The methods for this report followed the appropriate use and reporting of uncontrolled case series in the medical literature reporting guidelines.³⁵

Neuropathologic Evaluation

Neuropathologic evaluation occurred blinded to the clinical evaluation by neuropathologists (A.C.M., B.R.H., V.E.A., and T.D.S.). Pathologic processing and evaluation were conducted using previously published methods and as described in the eAppendix in Supplement 1.^{12-14,34,36} Neuropathologic diagnoses were made using NINDS National Institute of Biomedical Imaging and Bioengineering criteria for CTE^{31,32} and well-established criteria for other neurodegenerative diseases.³⁷⁻⁴⁰ The CTE p-tau pathologic findings were classified into 4 stages using the McKee staging scheme for CTE.^{13,41}

Brain tissue was also evaluated for the presence of vascular pathologic findings, including gross infarcts, microinfarcts, atherosclerosis, and arteriolosclerosis, as well as white matter rarefaction and the presence of perivascular pigmentladen macrophages within the deep cerebral white matter. For pathologic findings rated on a none, mild, or moderatesevere scale (cerebral amyloid angiopathy, white matter rarefaction, atherosclerosis, and arteriolosclerosis), scores were dichotomized as present vs absent. Neuropathologic diagnoses were reviewed by the 4 neuropathologists (A.C.M., B.R.H., V.E.A., and T.D.S.); any discrepancies were resolved by discussion and consensus of the group.

Clinical Evaluation

Retrospective clinical evaluations with next of kin were performed on 143 brain donors using online surveys and/or structured and semistructured postmortem telephone interviews, as described previously.34 Established scales modified for retrospective administration were given to informants of brain donors to assess cognitive symptoms, mood disturbances, and neurobehavioral dysregulation. Scales that assessed cognitive and functional symptoms included BRIEF-A Metacognition Index, Cognitive Difficulties Scale, and the Functional Activities Questionnaire. Scales that assessed neurobehavioral dysregulation included BRIEF-A Behavioral Regulation Index and Barratt Impulsiveness Scale 11. The Apathy Evaluation Scale and Geriatric Depression Scale, 15-item version assess apathy and depression symptoms, respectively. These scales were used as primary outcomes in this study. Further details on clinical protocols are outlined in the eAppendix in Supplement 1. Eighteen brain donors with missing clinical scale data were excluded from clinical analyses.

Statistical Analysis

Statistical analysis was conducted between August 2021 and June 2023. Qualitative assessments of the neuropathologic and clinical features were performed, and descriptive statistics were generated from SPSS software, version 20 (IBM Inc). Brain donors with and without CTE were compared on demographic, athletic, medical, and sport characteristics using independent-sample *t* tests (for continuous outcomes) and the χ^2 test or Fisher exact test (for binary outcomes). Neuropathologic and clinical features were compared between brain donors with and without CTE using analysis of covariance for continuous outcomes and binary logistic regression for binary outcomes. The analyses were controlled for age at death. Data were collected and stored using REDCap, version 8.5.1 (Vanderbilt University). A 2-sided *P* < .05 was considered statistically significant.

Results

Among the 152 brain donors, age at death ranged from 13 to 29 years (mean [SD] age, 22.97 [4.31] years) (**Table 1**). Eleven brain donors (7.2%) were female and 141 (92.8%) were male; 1 (0.7%) was American Indian or Alaska Native, 27 (17.8%)

jamaneurology.com

were Black, 111 (73.0%) were White, and 13 (8.6%) had missing or other race, including multiracial (White-African American, White-Indigenous American, White-Asian Indian, or White-Filipino), Tongan, and unspecified. Race information was not provided on 5 donors. Chronic traumatic encephalopathy was neuropathologically diagnosed in 63 brain donors (41.4%), 1 of whom was a woman (1.6%). Brain donors who had CTE were more likely to be older (mean difference, 3.92 years; 95% CI, 2.74-5.10 years; P < .001) and have a reported Black racial identity (16 [25.4%]; P = .047). Black brain donors were older than donors of other races (mean difference, 1.61 years; 95% CI, 0.01-3.23 years; P = .051) and had more years of football play (mean difference, 3.93 years; 95% CI, 2.07-5.78 years; *P* < .001). Brain donors with CTE had a higher level of education (28 [44.4%] vs 17 [19.1%] donors with a college degree or higher; P < .001). Suicide was the most common cause of death, followed by unintentional overdose; there were no differences in cause of death based on CTE status.

Of the 63 brain donors diagnosed with CTE, 60 (95.2%) were diagnosed with mild (stages I or II) CTE, including 39 (61.9%) with stage I and 21 (33.3%) with stage II. Three (4.8%) were diagnosed with stage III (**Table 2**, **Figure 1**, **Figure 2**, and **Figure 3**; eTable 1 in **Supplement 1**). Those with stage III CTE included 1 former National Football League (NFL) player, 1 college football player, and 1 professional rugby player. No brain donors were diagnosed with stage IV CTE.

Football Players

Of the 152 brain donors, 92 (60.5%) played US football as their primary sport, and 48 donors (76.2%) with CTE played football compared with 44 (49.4%) without CTE (P < .001). Although significantly more donors with CTE played football at the professional level (11 [22.9%]) compared with those without CTE (1 [2.3%]) (P < .001), 37 (58.7%) of those with CTE played football as their primary sport at the amateur level, with 21 (33.3%) playing in college and 16 (25.4%) never playing after high school. Two individuals (3.2%) who played only youth-level football were diagnosed with CTE; however, both had substantial RHI exposure through nonfootball activities, including military service with blast injury (n = 1) and motocross (n = 1). Eleven of 12 professional football players (91.7%) were diagnosed with CTE, including 11 of 11 NFL players (100%). For those who played football, duration of playing career was significantly longer in those with CTE compared with those without CTE (mean difference, 2.81 years; 95% CI, 1.15-4.48 years; P < .001). Playing position did not significantly differ between the groups.

Ice Hockey Players

Of the 16 primary ice hockey players, 6 (37.5%) were diagnosed with CTE. Four amateur ice hockey players were diagnosed with stage I or II CTE; 1 played hockey at the youth level, 2 reached the junior level, and 1 played in college. One of the high school hockey players also played 2 years of college football. One (of 1) National Hockey League (NHL) player had stage II CTE. One (of 1) non-NHL professional hockey player had stage I CTE.

Soccer and Rugby Players and Wrestlers

Four of 23 athletes (17.4%) who played soccer were diagnosed with CTE. Two high school soccer players were diagnosed with stage I CTE, 1 female collegiate player was diagnosed with stage I CTE, and 1 semiprofessional player was diagnosed with stage II CTE and amyotrophic lateral sclerosis. Two of 9 amateur wrestlers (22.2%) were diagnosed with CTE, 1 who wrestled in high school and 1 in college. There were 4 young rugby players in the sample, including a 17-year-old girl who died of second impact syndrome. Of the 4 players, 2 (50.0%) had CTE, a professional rugby player and a high school rugby

player who also played high school football. One (of 1) professional wrestler was diagnosed with stage II CTE.

Female Brain Donors

There were 11 female brain donors (7.2%). Overall, the female donors were younger (mean [SD] age at death, 20.82 [4.49] years; age range, 16-28 years) and had fewer years of contact sport exposure than the male donors. Their primary sources of exposure to RHIs included college soccer (n = 2), high school soccer (n = 4), high school rugby (n = 1), high school ice hockey (n = 1), professional cycling (n = 1), high school softball (n = 1),

	Total sample (N = 152)	No CTE	CTE	- P value	
Characteristic		(n = 89)	(n = 63)		
Age at death, mean (SD) [range], y	22.97 (4.31) [13-29]	21.35 (4.39) [13-29]	25.27 (2.97) [17-29]	<.001	
Sex					
Male	141 (92.8)	79 (88.8)	62 (98.4)	.03	
Female	11 (7.2)	10 (11.2)	1 (1.6)		
Racial identity					
American Indian or Alaska Native	1 (0.7)	1 (1.1)	0		
Black	27 (17.8)	11 (12.4)	16 (25.4)	.047 ^b	
White	111 (73.0)	67 (75.3)	44 (69.8)		
Other or missing ^c	13 (8.6)	10 (11.2)	3 (4.8)		
Educational level					
No or some high school	29 (19.1)	25 (28.1)	4 (6.3)		
High school or GED	20 (13.2)	15 (16.9)	5 (7.9)		
Some college, no degree	52 (34.2)	29 (32.6)	23 (36.5)	-	
College degree	42 (27.6)	15 (16.9)	27 (42.9)	- <.001 ^d	
More than college or graduate degree	3 (2)	2 (2.2)	1 (1.6)		
Unknown	6 (4)	3 (3.4)	3 (4.8)		
Primary sport played					
US football	92 (60.5)	44 (49.4)	48 (76.2)		
Ice hockey	16 (10.5)	10 (11.2)	6 (9.5)		
Soccer	23 (15.1)	19 (21.4)	4 (6.3)		
Amateur wrestling	9 (5.9)	7 (7.9)	2 (3.2)	<.001 ^e	
Karate	1 (0.7)	1 (1.1)	0		
Professional wrestling	1 (0.7)	0	1 (1.6)		
Rugby	4 (2.6)	2 (2.2)	2 (3.2)		
Lacrosse	2 (1.3)	2 (2.2)	0		
Equestrian	1 (0.7)	1 (1.1)	0		
Softball	1 (0.7)	1 (1.1)	0		
Ultimate frisbee	1 (0.7)	1 (1.1)	0		
Cycling	1 (0.7)	1 (1.1)	0		
Duration of American football play, mean (SD), y	10.29 (4.19)	8.83 (3.95)	11.64 (3.98)	<.001	
Age at first exposure to US football play, mean (SD), y	9.25 (2.71)	9.27 (2.58)	9.23 (2.87)	.94	
Highest level of US football played					
Youth	7 (7.6)	5 (11.4)	2 (4.2)		
High school	45 (48.9)	31 (7.5)	14 (29.2)	.004 ^f	
College	26 (23.3)	5 (11.4)	21 (43.8)		
Semiprofessional	2 (2.2)	2 (4.5)	0		
Professional	12 (13)	1 (2.3)	11 (22.9)		

(continued)

Table 1. Demographic and Athletic Characteristics of the Brain Donors^a (continued)

Characteristic	Total sample (N = 152)	No CTE (n = 89)	CTE (n = 63)	- P value
American football position played				
Offensive lineman	11 (12)	7 (15.9)	4 (8.3)	
Tight end	2 (2.2)	1 (2.3)	1 (2.1)	
Quarterback	2 (2.2)	1 (2.3)	1 (2.1)	
Running back	5 (5.4)	1 (2.3)	4 (8.3)	
Wide receiver	4 (4.3)	2 (4.5)	2 (4.2)	
Defensive lineman	8 (8.7)	5 (11.4)	3 (6.3)	
Linebacker	19 (20.7)	5 (11.4)	14 (29.2)	.20 ^g
Defensive back	12 (13)	0	12 (25)	
Punter	0	0	0	
Kicker	0	0	0	
Other	1 (1.1)	0	1 (2.1)	
Multiple	26 (28.3)	20 (45.5)	6 (12.5)	
Unknown	2 (2.2)	2 (4.5)	0	
Traumatic brain injury history				
Concussion count				
Postdefinition, median (range)	11 (0-1000)	10 (1-1000)	15 (0-1000)	.09
Traumatic brain injury with LOC				
Yes	62 (40.8)	38 (42.7)	24 (38.1)	.92
Military history				
Yes	9 (5.9)	6 (6.7)	3 (4.8)	.74
Combat	5 (3.3)	3 (3.4)	2 (3.2)	>.99
Primary cause of death				
Suicide	87 (57.2)	54 (60.7)	33 (52.4)	
Unintentional overdose	22 (14.5)	13 (14.6)	9 (14.3)	
Injury	16 (10.5)	7 (7.9)	9 (14.3)	
Cardiovascular disease	4 (2.6)	3 (3.4)	1 (1.6)	
Motor neuron disease	1 (0.7)	0	1 (1.6)	.25 ^h
Neurodegenerative	0	0	0	
Neoplasm	0	0	0	
Other	15 (9.9)	7 (7.9)	8 (12.7)	
Unknown	7 (4.6)	5 (6)	2 (3.2)	

Abbreviations: CTE, chronic traumatic encephalopathy; GED, general educational development; LOC, loss of consciousness; NA, not available.

^a Data are presented as number (percentage) of brain donors unless otherwise indicated. Donors with and without CTE were compared using independent-sample *t* tests for continuous measures and the χ^2 test or Fisher exact test for binary measures. Primary sport play was determined by the sport a donor played for the most years.

- ^b Compared Black vs other.
- ^c Other races include multiracial (White-African American, White-Indigenous American, White-Asian Indian, or White-Filipino), Tongan, and unspecified. Race information was not provided on 5 donors.
- ^d Compared college degree or greater vs others.
- ^e Compared US football vs others.
- ^f Compared professional vs others.

^g Compared linemen (offensive and defensive) vs others.

^h Compared suicide vs other.

and horseback riding (n = 1). The professional cyclist additionally played soccer for 10 years as a winger through high school, and the softball player also played field hockey as a forward through high school. The 1 female player diagnosed with CTE was 28 years old at death and played soccer as a forward for 18 years, beginning at age 3 years and playing through 3 years of Division I collegiate soccer. She was diagnosed with attention-deficit/hyperactivity disorder in college and prescribed stimulants. In addition to 2 concussions without loss of consciousness playing soccer, at age 24 years, she experienced a syncopal episode and a traumatic brain injury with loss of consciousness for 3 minutes. Computed tomographic findings were unremarkable. Four years later, she developed paranoia and suicidal thoughts. At 28 years of age, she died by suicide. Postmortem examination revealed stage I CTE, mild arteriolosclerosis, moderate white matter rarefaction, and marked perivascular pigment-laden macrophages in the frontal white matter (Figure 3).

jamaneurology.com

Amateur Athletes

Overall, amateur (ie, not semiprofessional or professional) athletes made up 128 of the 152 brain donors (84.2%). Of the 128 amateurs, 45 (35.2%) had CTE; of the 63 donors with CTE, 45 (71.4%) were amateurs. Amateur athletes included football players (78 [60.9%]), soccer players (22 [17.2%]), hockey players (10 [7.8%]), and wrestlers (9 [7.0%]).

Neuropathologic Features

Among those with available data, cavum septum pellucidum (26 [63.4%] vs 19 [30.2%]; P < .001), enlargement of the frontal horns of the lateral ventricles (17 [41.4%] vs 13 [20.4%]; P = .02), and a thalamic notch (5 [11.9%] vs 1 [1.5%]; P = .03) were significantly more frequent in those with vs without CTE, respectively (Figure 1 and Table 2). In cases of CTE, cavum septum pellucidum was often accompanied by thinning and atrophy of the fornices; 1 case had large septal fenestrations (Figure 1).

demonstrate of the state of	Table 1 (1) 453			. ·
Neuropathologic diagnosis	Iotal sample (N = 152)	No CTE (n = 89)	CTE (n = 63)	P value
	90 (EQ C)	80 (100)	0	
	09 (30.0)	03(100)	0	NA
Stage I	39 (25.7)	0	39 (61.9)	
Stage II	21 (13.8)	0	21 (33.3)	
Stage III	3 (2.0)	0	3 (4.8)	
Stage IV	0	0	0	
ross features				
Brain weight, mean (SD), g	1440.3 (229.71)	1435.61 (221.75)	1447 (242.45)	.77
Cavum septum pellucidum	45 (43.3)	19 (30.2)	26 (63.4)	.001
Septal fenestrations	1 (0.7)	0	1 (2.4)	NA
Frontal atrophy				
None	120 (93)	71 (94.7)	49 (90.7)	.49
Mild	7 (5.4)	3 (4.0)	4 (7.4)	
Moderate-severe	2 (1.6)	1 (1.3)	1 (1.9)	
Temporal atrophy				
None	123 (96.1)	73 (97.3)	50 (94.3)	
Mild	5 (3.9)	2 (2.7)	3 (5.7)	.65
Moderate-severe	0	0	0	
Parietal or occipital atrophy				
None	127 (99.2)	75 (100)	52 (98.1)	.41
Mild	1 (0.8)	0	1 (1.9)	
Moderate-severe	0	0	0	
Hippocampal atrophy				
None	127 (92.7)	77 (93.8)	50 (90.9)	
Mild	10 (7.3)	5 (6.2)	5 (9.1)	.74
Moderate-severe	0	0	0	
Ventricular dilation				
None	75 (71.4)	51 (79.7)	24 (58.5)	
Mild	26 (24.8)	12 (18.8)	14 (34.1)	.02
Moderate-severe	4 (3.9)	1 (1.6)	3 (7.3)	
halamic notch	6 (5.5)	1 (1.5)	5 (11.9)	.03
licroscopic features				
White matter rarefaction				
None	78 (52.7)	51 (59.3)	27 (43.5)	
Mild	55 (37 2)	28 (32 6)	27 (43 5)	058
Moderate-severe	15 (10.2)	7 (8.1)	8 (12.9)	.050
Frontal perivascular macrophages	\/	(/	- ()	
None	28 (20 1)	25 (29 8)	3 (5 5)	
Mild	34 (24 5)	22 (26 2)	12 (21.8)	< 001
Moderate-severe	77 (55 1)	37 (44 1)	40 (72 7)	<.001
Cribriform state	,, (33.1)	57 (17.1)	10 (12.1)	
Nono	114 (77)	69 (80.2)	45 (72 6)	
Mild	18 (12 2)	7 (8 1)	11 (17 7)	20
Modorato covoro	16 (12.2)	10(11.6)	£ (0,7)	.28
Athereesleresis	10(10.0)	10(11.0)	0(9.7)	
Ameroscierosis	127 (00 2)	77 (00 7)	50 (100)	
NOR	127 (99.2)	1 (1 2)	50 (100)	-
MILO	1 (0.8)	1 (1.3)	0	NA
Moderate-severe	U	U	0	
Arteriosclerosis				
None	109 (72.2)	69 (77.5)	40 (64.5)	.08
Mild	29 (19.2)	14 (15.7)	15 (24.2)	
Moderate-severe	13 (8.6)	6 (6.7)	7 (11.3)	

(continued)

E6 JAMA Neurology Published online August 28, 2023

Neuropathologic and Clinical Findings in Athletes With Repetitive Head Impacts

Table 2. Neuropathologic Characteristics of the Brain Donors^a (continued)

Total sample (N = 152)	No CTE (n = 89)	CTE (n = 63)	P value
0	0	0	NA
1 (0.7)	0	1 (1.6)	NA
5 (3.3)	3 (3.4)	2 (3.2)	>.99
0	0	0	NA
0	0	0	NA
1 (0.7)	0	1 (1.7)	NA
1 (0.7)	0	1 (1.7)	NA
1 (0.7)	0	1 (1.7)	NA
1 (0.7)	0	1 (1.7)	NA
	Total sample (N = 152) 0 1 (0.7) 5 (3.3) 0 0 1 (0.7) 1 (0.7) 1 (0.7) 1 (0.7) 1 (0.7)	Total sample (N = 152) No CTE (n = 89) 0 0 1 (0.7) 0 5 (3.3) 3 (3.4) 0 0 1 (0.7) 0 1 (0.7) 0 1 (0.7) 0 1 (0.7) 0 1 (0.7) 0 1 (0.7) 0	Total sample (N = 152) No CTE (n = 89) CTE (n = 63) 0 0 0 1 (0.7) 0 1 (1.6) 5 (3.3) 3 (3.4) 2 (3.2) 0 0 0 1 (0.7) 0 0 1 (0.7) 0 1 (1.7) 1 (0.7) 0 1 (1.7) 1 (0.7) 0 1 (1.7) 1 (0.7) 0 1 (1.7) 1 (0.7) 0 1 (1.7)

Abbreviations: CTE, chronic traumatic encephalopathy; NA, not available; TDP-43, TAR DNA-binding protein 43.

^a Data are presented as number (percentage by category) of brain donors unless otherwise indicated. Some characteristics were not assessed in all participants.

Figure 1. Gross Neuropathologic Features Associated With Chronic Traumatic Encephalopathy (CTE) in Young Athletes

A Control aged 27 y



B Young athletes with CTE



A, A 27-year-old control. Coronal brain sections at the level of the caudate, accumbens, and putamen (left); anterior thalamus and mammillary bodies (center); and midthalamus (right). B, Young athletes with CTE. Examples of macroscopic brain abnormalities in CTE. Cavum septum pellucidum (top left; arrowhead), thalamic notch (top center; arrowhead), degeneration of fornix (top right; arrowhead), enlargement of the frontal horns of the lateral ventricles and septal fenestrations (bottom left; asterisk), enlargement of the frontal horns of the lateral ventricles and cavum septum pellucidum (2 bottom center images; arrowheads), and thalamic notch (bottom right; asterisk).

jamaneurology.com

Figure 2. Stages of Chronic Traumatic Encephalopathy (CTE) Severity in Contact Sport Athletes Younger Than 30 Years

A Stage I CTE with isolated lesions in the depths of the superior frontal sulci



B Stage II CTE with 5 lesions in the superior and lateral frontal sulci



C Stage II CTE with multiple large lesions in the frontal, parietal, and temporal cortex



D Stage III CTE with multiple large lesions in the superior frontal lobe, with p-tau in the amygdala and transentorhinal cortex (arrowheads)



Hemispheric 50-µm tissue sections immunostained with CP-13, directed against phosphoserine 202 of tau (courtesy of Peter Davies, PhD, Feinstein Institute for Medical Research; 1:200); positive hyperphosphorylated tau (p-tau) immunostaining appears dark brown, showing representative images of CTE in the young athlete brain donors using the McKee staging scheme (I-IV).^{1.2}

Microscopic Features

In the 63 brain donors diagnosed with CTE, pathognomonic lesions of CTE consisting of p-tau immunoreactive neurofibrillary tangles (NFTs), pretangles, and dotlike neurites surrounding a central blood vessel were found most frequently at the depths of the sulci in the dorsolateral frontal, superior frontal, and superior temporal cortices, followed by the inferior parietal, inferior frontal, and Rolandic cortices (Table 2 and Figures 2 and 3; eTable 1 in Supplement 1). Neurofibrillary tangles were also common in entorhinal cortex, amygdala, and locus coeruleus in those with stage II or III CTE. Multiplex immunofluorescent labeling of the pathognomonic CTE lesions indicated that the NFT-containing cells colocalized with MAP2, a marker of neurons, and that the predominant p-tau isoform was 4 repeat (4R) p-tau (eFigure in Supplement 1). No p-tau-containing astrocytes were found in the pathognomonic CTE lesions or the surrounding neuropil, and no p-tau thorn-shaped astrocytes were found in the subpial region of any brain donor. In all brain regions sampled except the mammillary bodies and calcarine cortex, there were significantly more p-tau NFTs in brain donors with CTE compared with those without (eTable 1 in Supplement 1).

Non-p-Tau Pathologic Alterations

White matter rarefaction, evident as reduced density of myelinated nerve fibers, conspicuous reactive astrocytes, and macrophages in the white matter, was found more often in those with CTE than those without CTE, although this finding did not reach significance (Figure 3). Perivascular pigmentladen macrophages in the frontal subcortical white matter were found significantly more often in those with CTE (52 [94.5%] vs 59 [70.3%]; P < .001) compared with those without CTE (Figure 3 and Table 2).

Among the 92 football players in the sample, years of football play was significantly associated with CTE status (odds ratio, 1.20; 95% CI, 1.07-1.34; P = .002) and perivascular macrophages in the frontal white matter (odds ratio, 1.26; 95% CI, 1.08-1.47; P = .003). Years of play was not associated with ventricular enlargement, cavum septum pellucidum, thalamic notch, or white matter rarefaction. One 28-year-old brain donor with stage II CTE showed mild cerebral amyloid angiopathy in the frontal leptomeninges, as reported previously.⁴² Neither diffuse nor neuritic plaques were found throughout the sample. One 27-year-old brain donor with CTE stage II had sparse Lewy bodies in the medulla. A semiprofessional soccer player was diagnosed with comorbid stage II CTE and amyotrophic lateral sclerosis.⁴³

Clinical Features

Across the sample of symptomatic brain donors exposed to RHIs, with and without CTE, cognitive, behavioral, and mood symptoms, as reported by informants using standardized scales, were highly frequent (eTable 2 in Supplement 1). There were no statistically significant differences between donors with a CTE diagnosis compared with those without CTE for any clinical symptom (eTable 2 in Supplement 2). Mood and neurobehavioral dysregulation symptoms were particularly frequent. Clinically meaningful symptoms of neurobehavioral

Figure 3. Microscopic Pathologic Features Associated With Chronic Traumatic Encephalopathy (CTE) in Young Athletes

A Pathognomonic CTE lesions

B AT8-positive NFTs and dotlike neurites



C Macrophages surrounding arterioles in frontal subcortical white matter

D White matter refraction with myelin pallor, astrocytosis, and macrophage infiltration



E Macrophages surrounding arterioles in frontal subcortical white matter in female soccer player



G Activated microglia clusters



H Macrophages surrounding arteriole in frontal white matter

F Conspicuous macrophages and reactive astrocytes in white matter





I Subpial and interface astrogliosis in frontal cortex



J Astrogliosis at interface between gray and white matter



A, AT8-immunostained sections of 10-µm formalin-fixed, paraffin-embedded sections shows pathognomonic CTE lesions consisting of hyperphosphorylated tau (p-tau) neurofibrillary tangles (NFTs), pretangles, and dotlike neurites around a central vessel (original magnification ×200). B, Dorsolateral frontal cortex of a 28-year-old female soccer player shows a cluster of AT8-positive NFTs and dotlike neurites at the depth of the sulcus surrounding several small vessels (original magnification ×400). The p-tau lesion measured 1.3 × 1.3 × 1.4 mm in total dimension. C, Dense pigment-laden and clear macrophages surround arterioles in the frontal subcortical white matter (Luxol fast blue, hematoxylin-eosin stain; original magnification ×200). D, White matter rarefaction with myelin pallor, astrocytosis, and macrophage infiltration (Luxol fast blue, hematoxylin-eosin stain; original magnification ×200). E, Dense pigment-laden and clear macrophages surround arterioles in the frontal subcortical white matter in the young female soccer player with CTE (Luxol fast blue, hematoxylin-eosin stain; original magnification ×200). E, Dense pigment-laden and clear macrophages surround arterioles in the frontal subcortical white matter in the young female soccer player with CTE (Luxol fast blue, hematoxylin-eosin stain; original magnification ×400). F, Conspicuous macrophages (left, arrowheads) and reactive astrocytes (right, arrowheads) in the white matter in the young female soccer player with CTE (Luxol fast blue, hematoxylin-eosin stain; original magnification ×600). G, Clusters of activated microglia in the frontal white matter of the young female soccer player with CTE (LBA1 immunostaining; original magnification ×600). H, Macrophages surrounding arteriole in the frontal cortex of the young female soccer player with CTE (glial fibrillary acidic protein immunostaining; original magnification ×200). J, Astrogliosis at the interface between the gray and white matter in the frontal cortex of the young female soccer player with CTE (g

dysregulation, based on the BRIEF-A Behavioral Regulation Index, were present in 50 of 88 (56.8%). Mean scores on the Barratt Impulsiveness Scale 11 were also high (mean [SD], 74.25 [16.40]). Clinically meaningful symptoms of apathy (Apathy Evaluation Scale) were present in 72 of 101 donors (71.3%), and clinically meaningful symptoms of depression (Geriatric Depression Scale, 15-item version) were present in 77 of 110 donors (70.0%).

Clinically meaningful symptoms of executive dysfunction, based on the BRIEF-A Metacognition Index, were present in 48 of 88 donors (54.5%). Attention and memory symptoms were less frequent. Functional difficulties were not common, as evidenced by the low Functional Activities Questionnaire scores.

Thirty-four of 119 donors (28.6%) sought substance use treatment during life. Alcohol abuse was present in 60 of 140 (42.9%), drug abuse was present in 54 of 141 (38.3%), and steroid use was present in 7 of 141 (5.0%). Stimulant use was present in 22 of 127 (17.3%). There were no differences in substance, alcohol, steroid, or stimulant use based on CTE status.

Discussion

Among 152 brain donors exposed to RHIs who donated their brain to the UNITE Brain Bank and were younger than 30 years at the time of death, 63 (41.4%) had autopsy-confirmed CTE, including a female collegiate soccer player (stage I CTE). Chronic traumatic encephalopathy has been previously diagnosed in women who experienced interpersonal violence and frequent head-banging⁴⁴⁻⁴⁶ and, most recently, in an exprofessional Australian rules football player. This is, to our knowledge, the first report of CTE in a woman who was an amateur soccer player.⁴⁷ Nearly all (60 [95.2%]) of the 63 young brain donors with CTE were diagnosed with mild CTE (stages I or II), and 39 (61.9%) were diagnosed with stage I CTE. Young athletes with neuropathologically confirmed CTE (n = 63) included American football players, ice hockey players, soccer and rugby players, amateur wrestlers, military veterans, and a professional wrestler. The study highlights that CTE can affect amateur as well as professional contact sports athletes, with 45 (71.4%) of the athletes diagnosed with CTE playing only as high as the high school or college level. Professional players also develop CTE at a young age. In this cohort of players younger than 30 years, 11 of 11 NFL players and 1 of 1 NHL players were diagnosed with CTE.

Not all individuals exposed to RHIs will develop CTE, and 89 donors (58.6%) in this sample did not have CTE. Despite the narrow age range of the sample, brain donors with CTE were older, were more likely to play American football, had longer duration of football play, and were more likely to play at an elite level, in line with previous studies^{15,41} of older players. These findings emphasize the dual roles of age and duration of exposure to RHIs in the development of CTE, even among younger individuals.

In young brain donors, CTE was often accompanied by other pathologic abnormalities. Cavum septum pellucidum, often with degeneration and thinning of the fornices, was present significantly more often in donors with CTE. In addition, there was more enlargement of the frontal horns of the lateral ventricles and notching of the medial thalamus.

Chronic traumatic encephalopathy was evident in young athletes as neuronal p-tau aggregates, including NFTs and dotlike neurites, oriented around a small vessel most often in the superior frontal, dorsolateral frontal, and superior temporal cortices. The predominant p-tau isoform in the CTE lesions was 4R p-tau, consistent with a previous study⁴⁸ showing early 4R p-tau predominance and increasing 3-repeat (3R):4R p-tau ratio with increasing severity of CTE. No p-tau astrocytes were found in the parenchyma or subpial region. The lack of p-tau astrocytes was surprising because subpial p-tau thorn-shaped astrocytes are a frequent finding in older individuals with CTE.^{31,32,49,50} Their absence indicates that p-tau astrocytes are not an early or essential feature of CTE. The lack of p-tau astrocytes in young donors with CTE also supports the second NINDS consensus conference conclusion that the pathognomonic CTE lesion requires p-tau aggregates in neurons.32

Perivascular pigment-laden macrophages in the frontal white matter were significantly greater in CTE and associated with duration of exposure to RHIs. This finding suggests that disruption of the blood-brain barrier is increased after RHIs and might play a critical role in CTE pathogenesis. Previous autopsy studies^{51,52} in older individuals with CTE also have shown blood-brain barrier dysfunction and microvascular alterations. The vascular injury-associated markers intercellular adhesion molecule 1, vascular cell adhesion molecule 1, and C-reactive protein were increased in individuals with CTE compared with RHI-exposed and RHI-naive controls.⁵³ In addition, intercellular adhesion molecule 1 and C-reactive protein levels increased with RHI exposure duration and were associated with increased microglial inflammation and p-tau pathologic findings.⁵³ Microglial inflammation correlates with RHI duration and CTE severity,⁴⁹ and reactive astrocytosis at the gray-white matter interface has been reported after RHI and in CTE.^{54,55} Moreover, these findings are in line with dynamic contrast-enhanced MRI studies⁵⁶ of living American football players showing persistent blood-brain barrier dysfunction and white matter alterations on diffusion tensor imaging.

White matter rarefaction was increased in the deceased young athletes with CTE, albeit of marginal significance. White matter changes are also common in older individuals with autopsy-confirmed CTE.^{23,26,27,57} White matter rarefaction in older brain donors with CTE is directly associated with RHI exposure, CTE status, and dementia.²³ In addition, a single nuclear RNA sequencing study²⁷ found that the number of oligodendrocytes was reduced and altered in relative subtype proportions in the dorsolateral frontal white matter of older athletes with CTE compared with controls. A recent autopsy study²⁶ of 205 older male brain donors with and without CTE found decreased myelin-associated proteins in frontal white matter that corresponded to years of exposure and age at first exposure to American football. Using high spatial resolution ex vivo diffusion tensor imaging, researchers have reported alterations in fractional anisotropy in white matter underlying sulci with CTE lesions and microscopic evidence of axonal disruption.⁵⁷ In addition, in vivo MRI studies show alterations of the corpus callosum in former NFL players on diffusion imaging scans²⁵ and greater volumes of white matter hyperintensities on antemortem fluid-attenuated inversion recovery MRI²⁴ and T1 scans.⁵⁸ These observations suggest that white matter rarefaction, perivascular pigmentladen macrophages, neuroinflammation, and interface astrocytosis might represent key components of RHI-induced brain injury and CTE.^{1-6,28,30}

Clinical symptoms, as reported retrospectively by next of kin, were common in young donors exposed to RHI with and without CTE. Across the sample, based on modified standardized scales, approximately 50% had clinically meaningful symptoms of executive dysfunction. However, difficulties with instrumental activities of daily living were infrequent. Regarding neuropsychiatric symptoms, nearly 60% had symptoms of behavioral dysregulation as measured by the BRIEF-A Behavioral Regulation Index, and impulse control difficulties were also frequently endorsed. Approximately 70% reported meaningful symptoms of depression and apathy. The fact that there were no significant differences in clinical symptoms between those with CTE and those without may be attributable to selection bias (ie, those who donate are more likely to have symptoms). It may also indicate that retrospective review of symptoms with family members might not be granular enough to detect subtle clinical differences. It further implies that the symptoms are not specific to low-stage CTE. Despite all brain donors being symptomatic, 58.6% of the sample did not have CTE, emphasizing that not all contact sport athletes with symptoms have CTE. Nontau pathologic findings related to RHIs (eg, white matter rarefaction, perivascular macrophage infiltration, neuroinflammation, and astrocytosis) and unrelated to RHIs (eg, environmental stressors, medical history, genetic factors, and mental health issues) likely contribute to the clinical symptoms. Future studies that compare RHI-naive individuals with those with varying severities of RHI exposure, with and without CTE, are required to determine whether p-tau pathologic findings, microvascular alterations, or white matter loss are associated with cognitive, behavioral, or mood alterations that occur after RHIs and in CTE in young people.

There is a clear need for improved clinical characterization of living athletes across the age spectrum who have been exposed to RHIs through prospective objective assessments. The importance of prospective research studies on young athletes is underscored by the difficulty of assessing and managing symptoms after RHI in living athletes and the substantial contribution of uncertainty to their psychological stress. A recent study⁵⁹ reported that one-third of former NFL players are "extremely concerned" about cognitive difficulties and "having CTE." There is also a critical need for symptomatic individuals exposed to RHIs to be medically evaluated and followed up, because their symptoms can often be successfully treated in a clinical care setting.

Limitations

This study has some limitations. We did not evaluate the incidence or prevalence of CTE in the general population or in young contact sport athletes and other individuals exposed to RHIs, and no estimates of incidence or prevalence can be implied or concluded from this study. This study is constrained to brain donors whose families desired a neuropathologic examination after their loved one's death, primarily White male football players. There are limitations of ascertainment bias in studies associated with participation in a brain donation program, and those with symptoms during life, regardless of RHI or CTE status, are more likely to donate, potentially explaining the high rates of symptoms in the overall sample. The study is also limited by the lack of a comparison group that is representative of all young individuals exposed to RHIs and a control group of age- and sex-matched individuals not exposed to RHIs. Notably, such resources (ie, autopsies of young age individuals unexposed to RHIs) are extremely limited, underscoring the novelty and importance of this sample. Future studies comparing RHI-naive with RHI-exposed young brain donors will help isolate the clinical and neuropathologic effects of RHIs independent of CTE.

Conclusion

In a convenience brain bank sample of 152 young athletes exposed to RHI who were younger than 30 years at the time of death, 63 (41.4%) had neuropathologic evidence of CTE, including 1 female athlete. Most young athletes with CTE played at the high school and college levels, and sports included amateur football, ice hockey, rugby and soccer, and wrestling. Young athletes with CTE had significantly more ventricular dilatation, cavum septum pellucidum, thalamic notching, p-tau pathologic findings, and perivascular pigment-laden macrophages in the frontal white matter than those without CTE. Young donors exposed to RHIs were highly symptomatic regardless of CTE status, and the causes of symptoms in this sample, as reported by informants, are likely multifactorial and include RHI- and non-RHI-related causes. Furthermore, despite all donors being symptomatic, 58.6% did not have pathologic evidence of CTE. Future studies that include young brain donors unexposed to RHIs are needed to clarify the association among RHI exposure, white matter and microvascular pathologic findings, CTE, and clinical symptoms.

ARTICLE INFORMATION

Accepted for Publication: July 6, 2023. Published Online: August 28, 2023.

doi:10.1001/jamaneurol.2023.2907 **Open Access:** This is an open access article distributed under the terms of the CC-BY License.

© 2023 McKee AC et al. JAMA Neurology.

jamaneurology.com

Author Affiliations: Veterans Affairs (VA) Boston Healthcare System, US Department of Veteran Affairs, Boston, Massachusetts (McKee, Huber, Cherry, Alvarez, Cormier, Nicks, Stein); Alzheimer's Disease Research Center and Chronic Traumatic Encephalopathy Center, Chobanian and Avedisian School of Medicine, Boston University, Boston, Massachusetts (McKee, Mez, Abdolmohammadi, Butler, Huber, Uretsky, Babcock, Cherry, Alvarez, Martin, Tripodis, Palmisano, Cormier, Kubilus, Nicks, Kirsch, Mahar, McHale, Nowinski, Cantu, Stern, Daneshvar, Goldstein, Kowall, Stein, Alosco); Department of Neurology, Chobanian and Avedisian School of Medicine, Boston University, Boston, Massachusetts (McKee, Mez, Huber, Alvarez, Nicks, Stern, Katz, Kowall, Dwyer, Alosco); Department of Pathology and Laboratory Medicine, Chobanian and Avedisian School of Medicine, Boston University, Boston, Massachusetts (McKee, Cherry, Kirsch, Stein); VA Bedford Healthcare System, US Department of Veteran Affairs, Bedford, Massachusetts (McKee, Huber, Alvarez, Cormier, Kubilus, Nicks, Stein); National Center for PTSD VA Boston Healthcare Boston Massachusetts (McKee, Huber); Department of Biostatistics, Boston University School of Public Health. Boston. Massachusetts (McKee. Tripodis): Department of Anatomy and Neurobiology, Chobanian and Avedisian School of Medicine. Boston University, Boston, Massachusetts (Butler, Babcock); Biostatistics and Epidemiology Data Analytics Center, Boston University School of Public Health, Boston, Massachusetts (Martin, Tripodis, Palmisano); Concussion Legacy Foundation, Boston, Massachusetts (McHale, Nowinski, Cantu); Department of Neurosurgery, Emerson Hospital, Concord, Massachusetts (Cantu); Department of Neurosurgery, Chobanian and Avedisian School of Medicine, Boston University, Boston, Massachusetts (Cantu, Stern); Department of Rehabilitation Medicine, Harvard Medical School, Boston, Massachusetts (Daneshvar); Department of Psychiatry, Chobanian and Avedisian School of Medicine, Boston University, Boston, Massachusetts (Goldstein); Department of Biomedical, Electrical, and Computer Engineering. Boston University College of Engineering, Boston, Massachusetts (Goldstein); Braintree Rehabilitation Hospital, Braintree, Massachusetts (Katz, Dwyer).

Author Contributions: Drs McKee and Alosco had full access to all of the data in the study and take responsibility for the integrity of the data and the accuracy of the data analysis. Concept and design: McKee, Abdolmohammadi, Kirsch, Mahar, Daneshvar, Stein, Alosco. Acauisition, analysis, or interpretation of data: McKee, Mez, Abdolmohammadi, Butler, Huber, Uretsky, Babcock, Cherry, Alvarez, Martin, Tripodis, Palmisano, Cormier, Kubilus, Nicks, Kirsch, McHale, Nowinski, Cantu, Stern, Daneshvar, Goldstein, Katz, Kowall, Dwver, Stein, Alosco, Drafting of the manuscript: McKee, Abdolmohammadi, Butler, Martin, Alosco. Critical review of the manuscript for important intellectual content: McKee, Mez, Abdolmohammadi, Butler, Huber, Uretsky, Babcock, Cherry, Alvarez, Tripodis, Palmisano, Cormier, Kubilus, Nicks, Kirsch, Mahar, McHale, Nowinski, Cantu, Stern, Daneshvar, Goldstein, Katz, Kowall, Dwyer, Stein, Alosco. Statistical analysis: Abdolmohammadi, Cherry, Tripodis. Obtained funding: McKee, Mez, Kirsch, Nowinski, Stein. Administrative, technical, or material support: McKee, Butler, Huber, Uretsky, Babcock, Cherry, Palmisano, Cormier, Kubilus, Kirsch, Mahar, McHale, Goldstein, Kowall, Stein, Supervision: McKee, Alvarez, Palmisano, Cantu, Daneshvar, Alosco.

Conflict of Interest Disclosures: Dr McKee is a member of the Mackey-White Health and Safety Committee of the National Football League Players Association and reported receiving grants from the National Institutes of Health and Department of Veteran Affairs and other funding from the Buoniconti Foundation and MacParkman Foundation during the conduct of the study. Dr Mez reported receiving grants from the National Institutes of Health, Department of Defense, Alzheimer's Association, and Concussion Legacy Foundation during the conduct of the study. Dr Nowinski is the cofounder and chief executive officer of the Concussion Legacy Foundation, and serves as an adviser for Oxeia Biopharmaceuticals. Dr Cantu is a paid consultant to the National Football League Head Neck and Spine Committee, a vice president and chair of the scientific advisory committee of the National Operating Committee on Standards for Athletic Equipment, and a consultant to the Concussion Legacy Foundation: he also receives royalties from Houghton Mifflin Harcourt and compensation for expert legal opinion to the National Collegiate Athletic Association and National Hockey League and is a member of the Mackey-White Health and Safety Committee of the National Football League Players Association. Dr Stern is a paid consultant to Biogen, is a member of the Mackey-White Health and Safety Committee of the National Football League Players Association, receives royalties for published neuropsychological tests from Psychological Assessment Resources Inc, is a member of the board of directors of King-Devick Technologies, and reported grants from the National Institutes of Health during the conduct of the study. Dr Goldstein is a paid consultant to Johnson & Johnson, Janssen Research & Development LLC, and Rebiscan Inc and has received funding from the World Wrestling Entertainment (WWE) and Ivivi Health Sciences. Dr Katz reported receiving grants from Boston University School of Medicine Department of Neurology during the conduct of the study, receives royalties from Springer/Demos Publishing for a textbook on brain injury, serves as an expert witness in legal cases involving brain injury and concussion, receives a stipend from Encompass Health as program medical director for brain injury and chair of the annual neurorehabilitation conference, and has received honoraria for a keynote address for the HealthSouth annual medical directors meeting. Dr Alosco reported receiving grants from National Institutes of Health/ National Institute on Neurological Disorders and Stroke during the conduct of the study. No other disclosures were reported.

Funding/Support: This work was supported by grants from the Andlinger Foundation, National Football League, MacParkman Foundation, National Operating Committee on Standards for Athletic Equipment, and the Nick and Lynn Buoniconti Foundation (Dr McKee); the Alzheimer's Association (NIRG-15-362697 and NIRG305779 to Dr Cherry); the Concussion Legacy Foundation; World Wrestling Entertainment (Dr Goldstein); the National Center for Advancing Translational Medicine (UL1TROO1430 to Dr Mez); the National Institute of Neurological Disorders and Stroke (U01NS086659, U01NS093334, U54NS115266, R01NS078337, and R56NS078337 to all authors and K23NS102399 to Dr Alosco): the National Institute on Aging (AG057902, AG06348 and supplement 0572063345 to Dr McKee; and AG61028, K23AG046377, and R21HD089088 to Dr Mez; F32NSO96803 to Dr Alosco; P30AG13846 to Dr Kowall and AG1649 to Dr McKee); the US Department of Defense (W81XWH-13-2-0095 and W81XWH-13-20064 to Dr Mez); and the US Department of Veterans Affairs (CXOO135 to Dr McKee and CX001038 to Dr Stein).

Role of the Funder/Sponsor: The funders had no role in the design and conduct of the study; collection, management, analysis, and interpretation of the data; preparation, review, or approval of the manuscript; and decision to submit the manuscript for publication.

Data Sharing Statement: See Supplement 2.

REFERENCES

1. Rubin TG, Catenaccio E, Fleysher R, et al. MRI-defined white matter microstructural alteration associated with soccer heading is more extensive in women than men. *Radiology*. 2018; 289(2):478-486. doi:10.1148/radiol.2018180217

2. Lipton ML, Kim N, Zimmerman ME, et al. Soccer heading is associated with white matter microstructural and cognitive abnormalities. *Radiology*. 2013;268(3): 850-857. doi:10.1148/radiol.13130545

3. Smirl JD, Peacock D, Wright AD, et al. An acute bout of soccer heading subtly alters neurovascular coupling metrics. *Front Neurol*. 2020;11:738. doi:10. 3389/fneur.2020.00738

4. Smirl JD, Peacock D, Burma JS, et al. An acute bout of controlled subconcussive impacts can alter dynamic cerebral autoregulation indices: a preliminary investigation. *Eur J Appl Physiol*. 2022;122(4): 1059-1070. doi:10.1007/s00421-022-04908-4

5. Wright DK, O'Brien TJ, Shultz SR. Sub-acute changes on MRI measures of cerebral blood flow and venous oxygen saturation in concussed Australian rules footballers. *Sports Med Open*. 2022;8(1):45. doi:10.1186/s40798-022-00435-w

6. Bahrami N, Sharma D, Rosenthal S, et al. Subconcussive head impact exposure and white matter tract changes over a single season of youth football. *Radiology*. 2016;281(3):919-926. doi:10.1148/radiol.2016160564

7. Talavage TM, Nauman EA, Breedlove EL, et al. Functionally-detected cognitive impairment in high school football players without clinically-diagnosed concussion. J Neurotrauma. 2014;31(4):327-338. doi:10.1089/neu.2010.1512

8. Katz DI, Bernick C, Dodick DW, et al. National Institute of Neurological Disorders and Stroke consensus diagnostic criteria for traumatic encephalopathy syndrome. *Neurology*. 2021;96(18):848-863. doi:10.1212/WNL.000000000011850

9. Montenigro PH, Alosco ML, Martin BM, et al. Cumulative head impact exposure predicts later-life depression, apathy, executive dysfunction, and cognitive impairment in former high school and college football players. *J Neurotrauma*. 2017;34(2): 328-340. doi:10.1089/neu.2016.4413

10. Roberts AL, Pascual-Leone A, Speizer FE, et al. Exposure to American football and neuropsychiatric health in former National Football League players: findings from the Football Players Health Study. *Am J Sports Med.* 2019;47(12): 2871-2880. doi:10.1177/0363546519868989

11. Brett BL, Nader AM, Kerr ZY, et al. Disparate associations of years of football participation and a metric of head impact exposure with neurobehavioral outcomes in former collegiate football players. J Int Neuropsychol Soc. 2022;28(1): 22-34. doi:10.1017/S1355617721000047

12. McKee AC, Cantu RC, Nowinski CJ, et al. Chronic traumatic encephalopathy in athletes: progressive tauopathy after repetitive head injury. *J Neuropathol Exp Neurol*. 2009;68(7):709-735. doi:10.1097/NEN.0b013e3181a9d503

13. McKee AC, Stern RA, Nowinski CJ, et al. The spectrum of disease in chronic traumatic encephalopathy. *Brain.* 2013;136(pt 1):43-64. doi:10.1093/brain/aws307

14. Mez J, Daneshvar DH, Kiernan PT, et al. Clinicopathological evaluation of chronic traumatic encephalopathy in players of American football. *JAMA*. 2017;318(4):360-370. doi:10.1001/jama.2017.8334

15. Mez J, Daneshvar DH, Abdolmohammadi B, et al. Duration of American football play and chronic traumatic encephalopathy. *Ann Neurol.* 2020;87(1): 116-131. doi:10.1002/ana.25611

16. Bieniek KF, Ross OA, Cormier KA, et al. Chronic traumatic encephalopathy pathology in a neurodegenerative disorders brain bank. Acta Neuropathol. 2015;130(6):877-889. doi:10.1007/ s00401-015-1502-4

17. Suter CM, Affleck AJ, Lee M, Pearce AJ, Iles LE, Buckland ME. Chronic traumatic encephalopathy in Australia: the first three years of the Australian Sports Brain Bank. *Med J Aust*. 2022;216(10): 530-531. doi:10.5694/mja2.51420

 Omalu BI, DeKosky ST, Minster RL, Kamboh MI, Hamilton RL, Wecht CH. Chronic traumatic encephalopathy in a National Football League player. *Neurosurgery*. 2005;57(1):128-134. doi:10.1227/01.NEU.0000163407.92769.ED

19. Omalu BI, DeKosky ST, Hamilton RL, et al. Chronic traumatic encephalopathy in a national football league player: part II. *Neurosurgery*. 2006;59(5):1086-1092. doi:10.1227/01.NEU.0000245601.69451.27

20. Hazrati L-N, Tartaglia MC, Diamandis P, et al. Absence of chronic traumatic encephalopathy in retired football players with multiple concussions and neurological symptomatology. *Front Hum Neurosci.* 2013;7:222. doi:10.3389/fnhum.2013.00222

21. Stewart W, McNamara PH, Lawlor B, Hutchinson S, Farrell M. Chronic traumatic encephalopathy: a potential late and under recognized consequence of rugby union? *QJM*. 2016;109(1):11-15. doi:10.1093/qjmed/hcv070

22. LeClair J, Weuve J, Fox MP, et al. Selection bias analysis supports dose-response relationship between level of American football play and chronic traumatic encephalopathy diagnosis. *Am J Epidemiol.* 2022;191(8):1429-1443. doi:10.1093/aje/kwac075

23. Alosco ML, Stein TD, Tripodis Y, et al. Association of white matter rarefaction, arteriolosclerosis, and tau with dementia in chronic traumatic encephalopathy. *JAMA Neurol*. 2019;76 (11):1298-1308. doi:10.1001/jamaneurol.2019.2244

24. Uretsky M, Bouix S, Killiany RJ, et al. Association between antemortem FLAIR white matter hyperintensities and neuropathology in brain donors exposed to repetitive head impacts. *Neurology*. 2022;98(1):e27-e39. doi:10.1212/WNL. 000000000013012

25. Kochsiek J, O'Donnell LJ, Zhang F, et al. Exposure to repetitive head impacts is associated with corpus callosum microstructure and plasma total tau in former professional American football players. *J Magn Reson Imaging*. 2021;54(6): 1819-1829. doi:10.1002/jmri.27774

26. Alosco ML, Ly M, Mosaheb S, et al. Decreased myelin proteins in brain donors exposed to football-related repetitive head impacts. *Brain Commun.* 2023;5(2):fcadO19. doi:10.1093/ braincomms/fcadO19

27. Chancellor KB, Chancellor SE, Duke-Cohan JE, et al. Altered oligodendroglia and astroglia in chronic traumatic encephalopathy. *Acta Neuropathol*. 2021;142(2): 295-321. doi:10.1007/s00401-021-02322-2

28. Zimmerman KA, Laverse E, Samra R, et al. White matter abnormalities in active elite adult rugby players. *Brain Commun*. 2021;3(3):fcab133. doi:10.1093/braincomms/fcab133

29. Asken BM, DeKosky ST, Clugston JR, Jaffee MS, Bauer RM. Diffusion tensor imaging (DTI) findings in

adult civilian, military, and sport-related mild traumatic brain injury (mTBI): a systematic critical review. *Brain Imaging Behav*. 2018;12(2):585-612. doi:10.1007/s11682-017-9708-9

30. Weissberg I, Veksler R, Kamintsky L, et al. Imaging blood-brain barrier dysfunction in football players. *JAMA Neurol*. 2014;71(11):1453-1455. doi:10.1001/jamaneurol.2014.2682

31. McKee AC, Cairns NJ, Dickson DW, et al; TBI/CTE group. The first NINDS/NIBIB consensus meeting to define neuropathological criteria for the diagnosis of chronic traumatic encephalopathy. *Acta Neuropathol.* 2016;131(1):75-86. doi:10.1007/s00401-015-1515-z

32. Bieniek KF, Cairns NJ, Crary JF, et al; TBI/CTE Research Group. The Second NINDS/NIBIB Consensus Meeting to Define Neuropathological Criteria for the Diagnosis of Chronic Traumatic Encephalopathy. *J Neuropathol Exp Neurol*. 2021; 80(3):210-219. doi:10.1093/jnen/nlab001

33. Montenigro PH, Baugh CM, Daneshvar DH, et al. Clinical subtypes of chronic traumatic encephalopathy: literature review and proposed research diagnostic criteria for traumatic encephalopathy syndrome. *Alzheimers Res Ther.* 2014;6(5):68. doi:10.1186/s13195-014-0068-z

34. Mez J, Solomon TM, Daneshvar DH, et al. Assessing clinicopathological correlation in chronic traumatic encephalopathy: rationale and methods for the UNITE study. *Alzheimers Res Ther*. 2015;7(1): 62. doi:10.1186/s13195-015-0148-8

35. Kempen JH. Appropriate use and reporting of uncontrolled case series in the medical literature. *Am J Ophthalmol.* 2011;151(1):7-10.e1. doi:10.1016/j. ajo.2010.08.047

 McKee AC, Gavett BE, Stern RA, et al. TDP-43 proteinopathy and motor neuron disease in chronic traumatic encephalopathy. J Neuropathol Exp Neurol. 2010;69(9):918-929. doi:10.1097/NEN. 0b013e3181ee7d85

37. Beekly DL, Ramos EM, van Belle G, et al; NIA-Alzheimer's Disease Centers. The National Alzheimer's Coordinating Center (NACC) Database: an Alzheimer disease database. *Alzheimer Dis Assoc Disord*. 2004;18(4):270-277.

38. Brownell B, Oppenheimer DR, Hughes JT. The central nervous system in motor neurone disease. *J Neurol Neurosurg Psychiatry*. 1970;33(3): 338-357. doi:10.1136/jnnp.33.3.338

39. Mackenzie IRA, Neumann M, Bigio EH, et al. Nomenclature for neuropathologic subtypes of frontotemporal lobar degeneration: consensus recommendations. *Acta Neuropathol*. 2009;117(1): 15-18. doi:10.1007/s00401-008-0460-5

40. McKeith IG. Consensus guidelines for the clinical and pathologic diagnosis of dementia with Lewy bodies (DLB): report of the Consortium on DLB International Workshop. *J Alzheimers Dis.* 2006;9(3)(suppl): 417-423. doi:10.3233/JAD-2006-95347

41. Alosco ML, Cherry JD, Huber BR, et al. Characterizing tau deposition in chronic traumatic encephalopathy (CTE): utility of the McKee CTE staging scheme. *Acta Neuropathol*. 2020;140(4): 495-512. doi:10.1007/s00401-020-02197-9

42. Standring OJ, Friedberg J, Tripodis Y, et al. Contact sport participation and chronic traumatic encephalopathy are associated with altered severity and distribution of cerebral amyloid angiopathy. *Acta Neuropathol*. 2019;138(3):401-413. doi:10.1007/s00401-019-02031-x

43. McKee AC, Daneshvar DH, Alvarez VE, Stein TD. The neuropathology of sport. *Acta Neuropathol*. 2014; 127(1):29-51. doi:10.1007/s00401-013-1230-6 44. Danielsen T, Hauch C, Kelly L, White CL. Chronic traumatic encephalopathy (CTE)-type neuropathology in a young victim of domestic abuse. *J Neuropathol Exp Neurol*. 2021;80(6):624-627. doi:10.1093/jnen/nlab015

45. Roberts GW, Whitwell HL, Acland PR, Bruton CJ. Dementia in a punch-drunk wife. *Lancet*. 1990;335 (8694):918-919. doi:10.1016/0140-6736(90)90520-F

46. Hof PR, Knabe R, Bovier P, Bouras C. Neuropathological observations in a case of autism presenting with self-injury behavior. *Acta Neuropathol.* 1991;82(4):321-326. doi:10.1007/BF00308819

47. Suter CM, Affleck AJ, Pearce AJ, Junckerstorff R, Lee M, Buckland ME. Chronic traumatic encephalopathy in a female ex-professional Australian rules footballer. *Acta Neuropathol*. 2023. Published online June 30, 2023. doi:10.1007/s00401-023-02610-z

48. Cherry JD, Kim SH, Stein TD, et al. Evolution of neuronal and glial tau isoforms in chronic traumatic encephalopathy. *Brain Pathol.* 2020;30(5):913-925. doi:10.1111/bpa.12867

49. McKee AC, Stein TD, Huber BR, et al. Chronic traumatic encephalopathy (CTE): criteria for neuropathological diagnosis and relationship to repetitive head impacts. *Acta Neuropathol.* 2023; 145(4):371-394. doi:10.1007/s00401-023-02540-w

50. Butler MLMD, Dixon E, Stein TD, et al. Tau pathology in chronic traumatic encephalopathy is primarily neuronal. *J Neuropathol Exp Neurol*. 2022; 81(10):773-780. doi:10.1093/jnen/nlac065

51. Doherty CP, O'Keefe E, Wallace E, et al. Blood-brain barrier dysfunction as a hallmark pathology in chronic traumatic encephalopathy. *J Neuropathol Exp Neurol.* 2016;75(7):656-662. doi:10.1093/jnen/nlw036

52. Buée L, Hof PR, Bouras C, et al. Pathological alterations of the cerebral microvasculature in Alzheimer's disease and related dementing disorders. *Acta Neuropathol.* 1994;87(5):469-480. doi:10.1007/BF00294173

53. Kirsch D, Shah A, Dixon E, et al. Vascular injury is associated with repetitive head impacts and tau pathology in chronic traumatic encephalopathy. *J Neuropathol Exp Neurol*. 2023;82(2):127-139. doi:10.1093/jnen/nlac122

54. Cherry JD, Tripodis Y, Alvarez VE, et al. Microglial neuroinflammation contributes to tau accumulation in chronic traumatic encephalopathy. *Acta Neuropathol Commun.* 2016;4(1):112. doi:10.1186/s40478-016-0382-8

55. Babcock KJ, Abdolmohammadi B, Kiernan PT, et al. Interface astrogliosis in contact sport head impacts and military blast exposure. *Acta Neuropathol Commun*. 2022;10(1):52. doi:10.1186/s40478-022-01358-z

56. Veksler R, Vazana U, Serlin Y, et al. Slow blood-to-brain transport underlies enduring barrier dysfunction in American football players. *Brain*. 2020;143(6):1826-1842. doi:10.1093/brain/awaa140

57. Holleran L, Kim JH, Gangolli M, et al. Axonal disruption in white matter underlying cortical sulcus tau pathology in chronic traumatic encephalopathy. *Acta Neuropathol*. 2017;133(3): 367-380. doi:10.1007/s00401-017-1686-x

58. Alosco ML, Koerte IK, Tripodis Y, et al. White matter signal abnormalities in former National Football League players. *Alzheimers Dement (Amst)*. 2017;10:56-65. doi:10.1016/j.dadm.2017.10.003

59. Walton SR, Kerr ZY, Mannix R, et al. Subjective concerns regarding the effects of sport-related concussion on long-term brain health among former NFL players: an NFL-LONG study. *Sports Med*. 2022;52(5): 1189-1203. doi:10.1007/s40279-021-01589-5