ABSTRACT

Objectives To estimate the prevalence of post-traumatic stress disorder (PTSD) in a large civilian population with traumatic brain injury (TBI), and to assess whether brain injury severity is correlated with PTSD symptoms.

Design Observational, cross-sectional study.

Setting and participants Outpatient clinic in a major UK trauma centre and secondary care hospital. Estimates of PTSD prevalence are based on 171 sampled individuals attending TBI clinic within an 18-month period. Analysis of the relationship between TBI severity and PTSD was performed on the subset of 127 patients for whom injury severity data were also available.

Methods Civilian TBI clinic attendees completed validated self-report questionnaires assessing PTSD (PTSD Checklist Civilian Version (PCL-C)) and other psychiatric symptoms. From this, the prevalence of PTSD was estimated in our cohort. Postresuscitation Glasgow Coma Score and Marshall grade on CT brain scan were recorded as indicators of brain injury severity. A hierarchical regression explored whether TBI severity may predict PTSD scores.

Results A high prevalence of PTSD was estimated (21% with PCL-C score >50). Higher Marshall grading displayed a slight negative correlation with PTSD symptoms. This statistically significant relationship persisted after confounding factors such as depression and postconcussion symptoms were controlled for.

Conclusions PTSD and TBI frequently coexist, share antecedents and overlap in their resultant symptoms. This complexity has given rise to conflicting hypotheses about relationships between the two. This research reveals that PTSD is common in civilians with TBI (adding to evidence drawn from military populations). The analysis indicated that more severe brain injury may exert a slight protective influence against the development of PTSD—potentially by disrupting implicit access to traumatic memories, or via overlapping neuropsychiatric symptoms that impede diagnosis. The association suggests that further research is warranted to explore the reuse of routine clinical and neuroimaging data—investigating its potential to predict risk of psychiatric morbidity.

INTRODUCTION

The complex relationship between post-traumatic stress disorder (PTSD) and traumatic brain injury (TBI) presents opportunities to further the understanding of both conditions individually as well as their interplay. PTSD is a common mental health condition with an estimated lifetime prevalence of 7.8%. Risk is increased by severely distressing experiences such as sexual assault, life-threatening injury or emotional trauma during military service. Its psychosocial impact is significant, with a high risk of suicidal behaviour in patients with PTSD, impairments in social and occupational functioning, as well as increased...
utilisation of health services. The purported aetiology of PTSD involves an antecedent psychologically traumatic event which is deemed severely threatening. The presence of such a stressor is common to diagnostic criteria in the Diagnostic and Statistical Manual 5 (DSM 5) and the International Classification of Diseases 10 (ICD-10).

A psychologically traumatic event involving physical brain injury can potentially complicate the development of PTSD. Pre-event and postevent amnesia is often a feature of brain injury and concussion and yet the integrity of traumatic memories may also play an important role in the development of the disease. Where memory of an antecedent event is impaired due to traumatic amnesia, it has been proposed that this memory loss may have a potentially protective or even preventative role in PTSD development. The lack of intact recollection of a traumatic event may be associated with a failure to develop intrusive, distressing memories which are a hallmark of PTSD.

A varied range of cognitive deficits can result from TBI. This may in fact render patients more vulnerable to the development of PTSD as better premorbid function, with increased cognitive reserve, has been found to be a protective factor. Further research suggested that both mild TBI (mTBI) and severe TBI may predispose to PTSD even in the presence of amnesia and other cognitive abnormalities. There are diverse mechanisms by which brain injury may produce cognitive deficits, for example, diffuse axonal fragmentation can disrupt connections between key networks of cortical grey matter. However, the extent to which neuroimaging and gross structural changes can be linked to the development of PTSD in this patient group is poorly understood. As a result, uncertainties remain about the neuropathological mechanisms by which TBI and PTSD may be linked, particularly outside of the military/blast injury context.

Large studies exploring the relationships between TBI and PTSD often involve military populations, typically those involved in wars in Afghanistan and Iraq. Though pragmatic, an approach based on military cohorts is complicated by the potential exposure to multiple psychological stressors aside from the event responsible for TBI. Furthermore research in this population is largely focused on damage attributable to blast injuries or other mechanisms quite specific to military combat.

In contrast, civilian TBI is most commonly due to falls, vehicle crashes and assaults (as well as a varied range of other mechanisms). These mechanisms may result in qualitatively differing patterns of brain damage and psychological trauma, limiting the extent to which findings from specific military studies can be generalised to the civilian populous.

The relationship between the severity of brain injury and the development of PTSD remains controversial, with mixed findings in patients with mild versus severe injury. As a result, some studies have focused on mTBI in order to explore the effect on PTSD. A systematic review of such studies has highlighted marked heterogeneity of study design which obscures the relationship between the conditions. Drawing a distinction between mTBI and more severe injury may introduce an artificial dichotomy onto the spectrum of brain injury, potentially limiting our understanding of the relationship with PTSD.

This current study aimed to explore the relationship between brain injury factors and PTSD symptoms in a large civilian, outpatient population—while controlling for confounding variables. This has been conducted with the objectives of estimating the prevalence of PTSD, and assessing whether indicators of TBI severity predict PTSD symptom levels.

**METHODS**

Data were collected prospectively between December 2013 and June 2015 from patients attending an outpatient TBI clinic at the Queen Elizabeth Hospital in Birmingham—a major trauma centre in the UK. This included processes to ensure participants provided informed consent for their clinical data to be stored in database form, and for anonymised information to be used for the purposes of research.

Admission records were interrogated to record demographic details and best postresuscitative Glasgow Coma Score. Patients completed a battery of self-report questionnaires, including: health-related quality of life (Quality of Life after Brain Injury (QOLIBRI)), post-concussion symptoms (PCS) (Rivermead PCS Questionnaire), depression (Patient Health Questionnaire (PHQ9)) and PTSD severity (PTSD Checklist Civilian Version (PCL-C)). Exclusion criteria were: attendance due to non-traumatic pathology, chronic subdural haematoma or declining to provide informed consent. Additionally, participants could not be included if required data for the analysis were not available (as reported in the Results section).

PCL-C is a measure of PTSD symptoms adapted from the military questionnaire use in a civilian population and scores can range from 17 (minimal symptoms) to 85. Two cut-off levels are established to estimate PTSD prevalence using PCL-C scores: scores≤50 have been regarded as a suitable diagnostic estimate in the mTBI population, and scores≤44 have been validated based on studies in

| Variable | Range/Score
|---------|-------------
| Age     | 16–82 years; median 38; IQR 32 |
| Sex     | F: 37 (22%), M: 134 (78%) |
| Ethnicity | White 131 (77%), African Caribbean 6 (4%), Asian 18 (11%), Mixed 8 (5%), other 8 (5%) |
| PTSD (PCL-C score/84) | Range 5–84; mean 34.46; SD 18.12 |
Table 2 Proportion of patients with traumatic brain injury of differing severities based on the best postresuscitation Glasgow Coma Scale (GCS) and Marshall grade n=127

| Variable          | Range | Median | IQR  |
|-------------------|-------|--------|------|
| Quality of Life (QOLIBRI—%) | 33–100 | 59     | 29   |
| Concussion symptoms (Rivermead PCS) | 0–60  | 24     | 26.25|
| Depression symptoms (PHQ9) | 0–26  | 7      | 15.5 |
| PTSD symptoms (PCL-C) | 17–85 | 25     | 28   |

Table 3 Descriptive statistics of the group included in hierarchical regression n=127

Table 4 Proportion of patients with traumatic brain injury of differing severities based on the best postresuscitation Glasgow Coma Scale (GCS) and Marshall grade n=127

| Variable          | Range | Median | IQR  |
|-------------------|-------|--------|------|
| Quality of Life (QOLIBRI—%) | 33–100 | 59     | 29   |
| Concussion symptoms (Rivermead PCS) | 0–60  | 24     | 26.25|
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1. Patients and public were not directly involved in the development of this study.

RESULTS

To produce estimates of PTSD prevalence in this cohort, 171 participants were included (as their full PCL-C scores were available), 79% were men and the median age was 38. See table 1 for sample description.

Using PCL-C cut-off score >50, the prevalence of PTSD was 20.6%; using the lower threshold (score >44), prevalence was 31.6%.

Brain injury severity data were not available for all of the participants described above, due to incomplete records. Hierarchical multiple regression was performed based on those 127 participants who completed questionnaires, had CT head scan results available and their admission GCS recorded (see table 2). The 44 participants excluded (due to missing data) did not differ significantly in demographic or injury characteristics, nor in PTSD, depression or PCS scores.

Uncorrected exploratory correlations conducted within the regression group (n=127) suggested that PCSs (r=0.70) and depression (r=0.76) were moderately positively correlated with PTSD severity (p<0.01).

A two-level hierarchical regression analysis was performed (see table 3), with PTSD severity (PCL-C score) as the dependent variable. Model assumptions were tested and met. The first level of the regression consisted of potential predictors of PTSD score which
may be confounding factors, specifically age, sex, depression scores (PHQ9), PCSs (Rivermead) and quality of life (QOLIBRI). The second level contained GCS and Marshall grade (see table 4).

At the first level, depression and other potential confounders contribute significantly to the model, \( F(5,121)=35.59, p<0.01 \), accounting for 57.9% of the variance in PTSD severity, with depression the only individual significant factor. The second level including Marshall grade added a modest but statistically significant contribution to PTSD severity—\( F(7,119)=28.06, p<0.05 \). In contrast, GCS was not a statistically significant predictor of PTSD severity when other potential confounders were controlled for.

**DISCUSSION**

These findings reveal a high level of PTSD symptoms in the civilian TBI clinic population. Dependant on diagnostic threshold used, estimated prevalence of PTSD is between 20.6% and 31.6%, which is in keeping with previous studies of smaller cohorts.\(^9\) Furthermore, this is in keeping with findings from military populations showing associations between even mTBI and PTSD.\(^8\) This PCS and PTSD potentially both include features such as irritability, and both conditions can be associated depressed mood.\(^10\) Such potential for overlap at the symptom level introduces the possibility that the co-occurrence of PTSD, depression and PCS may be overestimated. Furthermore, GCS was not a significant correlate of PTSD severity. This reflects the possibility that a conventional distinction between mild, moderate and severe brain injury based purely on postresuscitative GCS, may not reflect the particular factors that predispose toward psychiatric morbidity. Further study is required to explore whether there is a relationship between altered GCS and PTSD in other settings.

The current results suggest that PTSD is common in this large cohort and that routinely collected radiological data may be of use in identifying those at greatest risk of severe PTSD symptoms. The strengths of this study include a pragmatic emphasis on tools which can be employed in routine clinical practice—review of CT scans, GCS levels and self-report questionnaires. Different criteria for PTSD have been used in previous research (using the ICD-10, DSM 4 and DSM 5) each with subtly different emphases. The PCL-C is based on established diagnostic features and can be reliably administered in the clinical setting, thereby enabling comparisons in the wider literature.\(^28\)

**Table 4** Hierarchical regression models

| Potential predictor          | Coefficient beta | B (95% CI)   | B SE |
|-----------------------------|------------------|--------------|------|
| Model 1                     |                   |              |      |
| Sex                         | −0.10            | −0.47 (−6.22 to 5.28) | 2.90 |
| Age                         | −0.03            | −0.03 (−0.16 to 0.10) | 0.06 |
| QoL                         | 0.13             | 0.15 (−0.07 to 0.36) | 0.11 |
| Concussion symptoms         | 0.23             | 0.28 (−0.06 to 0.62) | 0.17 |
| Depression symptoms         | 0.65             | 1.57 (0.97 to 2.17)* | 0.30 |
| Model 2                     |                   |              |      |
| Sex                         | −0.01            | −0.51 (−6.13 to 5.11) | 2.84 |
| Age                         | −0.03            | −0.03 (−0.16 to 0.09) | 0.06 |
| QoL                         | 0.13             | −0.15 (−0.07 to 0.36) | 0.11 |
| Concussion symptoms         | 0.18             | 0.22 (−0.12 to 0.56) | 0.17 |
| Depression symptoms         | 0.69             | 1.67 (1.08 to 2.26)* | 0.30 |
| GCS† (mild/moderate/severe) | −0.08            | −1.86 (−4.67 to 0.96) | 1.42 |
| Marshall grade†             | −0.12            | −1.73 (−3.45 to 0.01)* | 0.87 |

*Statistically significant p<0.05.
†Second level of hierarchical regression (routinely recorded brain injury factors).
GCS, Glasgow Coma Scale; QoL, quality of life.
some mild cognitive deficits are associated with PCS which may increase vulnerability to PTSD as previously discussed. In this study, controlling for PCSs within the regression analysis served to partially mitigate against this potential source of confusion.

Inevitably, certain limitations apply to the approach presented. A brief survey inevitably produces less precise estimates of prevalence of PTSD than a full psychiatric assessment. However, prior research suggests a high rate of psychiatric symptoms and that PTSD may be underdiagnosed in this group, so measuring symptom severity may highlight those for whom psychiatric review would be beneficial and could lead to diagnosis. Broader concerns about self-report measures may apply—whether this manifests as patients denying the severity of their symptoms, or overstating them in the hope of receiving more support. In spite of this, the PCL-C has been found to be reliable across comparable populations.

The outpatient sample taking part in this study represents a large and well-categorised civilian group in a real-world hospital setting; however, some factors may limit its generalisability to the wider TBI population. There is potential for selection bias in favour of patients with more persistent symptoms, as those attending the clinic are more likely to have enduring neuropsychiatric symptoms which justify their attendance. Conversely, patients with more severe injuries may have cognitive deficits that render them unable to complete the necessary questionnaires for inclusion, or they may be in inpatient settings that make clinic attendance less likely. The potential also exists for the severity of TBI to be underestimated through the use of best GCS score after resuscitation, rather than the use of initial GCS on admission. However, this compromise improves the ability of GCS to predict long-term outcomes. In spite of the majority of the cohort consisting of mTBI, the sample contains a wide range of injury severity levels, which partially serves to mitigate against a systematic bias of this type. The use of routinely collected clinical brain injury data (GCS, CT scan findings) is advantageous in that it is readily available and quite objective in nature, but the fact that such data are/may be recorded by different clinical teams (without specific training for the purpose of this study) has the potential to reduce inter-rater reliability.

This study sought to characterise this particular civilian TBI cohort, as data from similar large populations are relatively limited. However, the absence of a control group does limit the extent to which one can meaningfully speculate about the neural mechanisms by which TBI and PTSD may be linked. To elucidate this in future, studies including a control group of participants with extracranial trauma may be valuable to isolate the effect of brain damage from other aspects of psychological trauma associated with injury and hospitalisation. Finally, the majority male sample may be typical of TBI sufferers; however, this may be less representative of the wider civilian PTSD cohort. This study included TBI both with and without structural changes identified on CT. A full understanding of the links between acquired brain injury and psychiatric symptoms will require the location of any overt injury to be taken into account, although this was beyond the scope of the analysis presented.

The high prevalence of PTSD found in this study provides an important epidemiological estimate within the UK civilian population. These prevalence findings are in accordance with research in populations who have suffered general trauma (including extracranial injury) such as those involved in motor vehicle accidents and assaults. The novel finding of an independent negative correlation identified between Marshall grade and PTSD invites speculation that more severe structural brain damage may exert a modest protective effect against PTSD symptoms. This is borne out in the previous literature suggesting that severe TBI may prevent the development of PTSD in some cases. For example, it has been proposed that the prolonged periods of unconsciousness may exert a protective influence. This may be attributable to amnesia interfering with the process by which traumatic memories are formed. While intuitively plausible, the picture is complicated by findings in patients with mTBI, in which a longer duration of post-traumatic amnesia was found to be protective against certain PTSD symptoms in spite of the absence of overt structural brain injury. Some have extended this line of reasoning further to suggest that mTBI and PTSD are mutually exclusive regardless of amnesia.

In order to reconcile the findings of these potentially conflicting studies, three main mechanisms have been proposed to link TBI with PTSD via memory systems: unimpaired traumatic memories, traumatic amnesia with spared implicit memory of trauma and ‘islands of memory’ within post-traumatic amnesia. The findings of this study can be recognised within this framework, as more severe brain injury findings on CT are a significant predictor of milder PTSD symptoms. In more severe TBI, structural damage (and resultant neuronal loss) may produce functional impairment of implicit memory systems. Deficits in implicit memory are not easily recognised in routine clinical assessment of post-traumatic amnesia (which essentially test declarative, but not implicit, memory). Future research into these mechanisms may benefit from avoiding a potentially arbitrary dichotomy between mild and more severe TBI. Quantifying or systematically classifying brain injury severity on a continuous basis using more sophisticated imaging may enable measurable brain injury factors to be linked to different symptoms within PTSD. A refined method based on this principle may enable information from more detailed radiological modalities (such as MRI) to predict psychiatric symptoms at a level of precision that could become clinically meaningful. In future, this may require the specific use of appropriate imaging (such as MRI) to search for relevant markers of poor long-term outcome, rather than repurposing existing scans in an opportunistic manner. In cases where a focal traumatic lesion is identified, future research may benefit from also...
exploring the effect of the anatomical location of TBI and its relationship to psychiatric symptoms. Such precision may in future enable a meaningful taxonomy of the specific psychiatric sequelae that may arise, depending on the nature of their brain injury,\textsuperscript{35} with interventions targeted accordingly.

In spite of the limitations inherent in observational study of an outpatient clinic cohort, this research illustrates that PTSD represents a common condition among people with TBI. Furthermore, routinely performed CT scans can be reviewed to identify features that relate to psychiatric morbidity in a real-world civilian population. Higher Marshall grades (eg, 5–6) are modestly associated with lower PCL-C scores. The presence of a relationship between more severe brain injury and milder PTSD symptoms represents a novel finding, given that depression and PCPs have been controlled for in this design. The implications of this extend from the theoretical to the practical—inviting further exploration using more sophisticated imaging, as well as pointing toward pragmatic approaches to screen those patients with TBI at the highest risk of PTSD.

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**REFERENCES**

1. Kessler RC, Sonnega A, Bromet E, et al. Posttraumatic stress disorder in the national comorbidity survey. *Arch Gen Psychiatry* 1995;52:1048–60.

2. Tarrier N, Gregg L. Suicide risk in civilian PTSD patients. *Soc Psychiatry Psychiatr Epidemiol* 2004;39:655–61.

3. Hidalgo RB, Davidson JR. Posttraumatic stress disorder: epidemiology and health-related considerations. *J Clin Psychiatry* 2000;61 Suppl 7:5–13.

4. American Psychiatric Association. *Diagnostic and Statistical Manual of Mental Disorders (5th edition)* (DSM-5); APA, 2013.

5. World Health Organization. The ICD-10 classification of mental and behavioural disorders: clinical descriptions and diagnostic guidelines. Geneva 1992.

6. Mayou R, Bryant B, Duthie R. Psychiatric consequences of road traffic accidents. *BMJ* 1993;307:647–51.

7. Sbordone RJ, Lifer JC. Mild traumatic brain injury does not produce post-traumatic stress disorder. *Brain Inj* 1995;9:405–12.

8. Stein MB, McAllister TW. Exploring the convergence of posttraumatic stress disorder and mild traumatic brain injury. *Am J Psychiatry* 2009;166:769–76.

9. Bryant RA, Marossekje Y, Crooks J, et al. Posttraumatic stress disorder after severe traumatic brain injury. *Am J Psychiatry* 2000;157:629–31.

10. Bryant RA, Harvey AG. Postconcussive symptoms and posttraumatic stress disorder after mild traumatic brain injury. *J Neurol Ment Dis* 1999;187:302–5.

11. Levin HS, Wilde E, Troyanskaya M, et al. Diffusion tensor imaging of mild to moderate blast-related traumatic brain injury and its sequelae. *J Neurotrauma* 2010;27:683–94.

12. Bigler ED, Maxwell WL. Neuropathology of mild traumatic brain injury: relationship to neuroimaging findings. *Brain Imaging Behav* 2012;6:108–36.

13. Lew HL, Vanderploeg RD, Moore DF, et al. Overlap of mild TBI and mental health conditions in returning OIF/OEF service members and veterans. *J Rehabil Res Dev* 2008;45:11.

14. Schneiderman AI, Braver ER, Kang HK. Understanding sequelae of injury mechanisms and mild traumatic brain injury incurred during the conflicts in Iraq and Afghanistan: persistent postconcussive symptoms and posttraumatic stress disorder. *Am J Epidemiol* 2008;167:1446–52.

15. Langlois JA, Rutland-Brown W, Wald MM. The epidemiology and impact of traumatic brain injury: a brief overview. *J Head Trauma Rehabil* 2006;21:375–8.

16. Carlson KF, Kehrle SM, Meis LA, et al. Prevalence, assessment, and treatment of mild traumatic brain injury and posttraumatic stress disorder: a systematic review of the evidence. *J Head Trauma Rehabil* 2011;26:103–15.

17. Marshall LF, Marshall SB, Klauber MR, et al. A new classification of head injury based on computerized tomography. *J Neurosurg* 1991;75:514–S20.

18. von Steinbüchel N, Petersen C, Bullinger M. QOLIBRI Group. Assessment of health-related quality of life in persons after traumatic brain injury--development of the Qolibri, a specific measure. *Acta Neurochir Suppl* 2005;93:43.

19. King NS, Crawford S, Wendon FJ, et al. The rivermead post concussion symptoms questionnaire: A measure of symptoms commonly experienced after head injury and its reliability. *J Neurol* 1995;242:587–92.

20. Kroenke K, Spitzer RL. The PHQ-9: A new depression diagnostic and severity measure. *Psychiatr Ann* 2002;32:590–15.

21. Wilkins KC, Lang AJ, Norman SB. Synthesis of the psychometric properties of the PTSD checklist (PCL) military, civilian, and specific versions. *Depress Anxiety* 2011;28:596–606.

22. Weathers F, Litz B, Herman D, et al. The PTSD Checklist: Reliability, validity, & diagnostic utility. *Annual Meeting of the Internation Society of Traumatic Stress Studies*. San Antonio, Texas, 1993.

23. Hoge CW, McGurk D, Thomas JL, et al. Mild traumatic brain injury in U.S. Soldiers returning from Iraq. *N Engl J Med* 2008;358:453–63.

24. Terhakopian A, Sinai N, Engel CC, et al. Estimating population prevalence of posttraumatic stress disorder: An example using the PTSD checklist. *J Trauma Stress* 2008;21:290–300.

25. Udekwu P, Kromhout-Schiro S, Vasilis S, et al. Glasgow Coma Scale score, mortality, and functional outcome in head-injured patients. *J Trauma* 2004;56:1084–9.

26. Cifu DX, Keyser-Marcus L, Lopez E, et al. Acute predictors of successful return to work 1 year after traumatic brain injury: A multicenter analysis. *Arch Phys Med Rehabil* 1997;78:125–31.

27. Keane TM, Wolfe J. Comorbidity in post-traumatic stress disorder an analysis of community and clinical studies. *J Appl Soc Psychol* 1990;20:1776–88.

28. Ruggiero KJ, Del Ben K, Scotti JR, et al. Psychometric properties of the PTSD Checklist-Civilian Version. *J Trauma Stress* 2003;16:495–502.

29. Qureshi AI, Malik AA, Adil MM, et al. Hematoma enlargement among patients with traumatic brain injury: Analysis of a prospective multicenter clinical trial. *J Vasc Interv Neurol* 2015;8:42.
30. Holeva V, Tarrier N, Wells A. Prevalence and predictors of acute stress disorder and PTSD following road traffic accidents: Thought control strategies and social support. *Behav Ther* 2001;32:65–83.

31. Kilpatrick DG, Ruggiero KJ, Acierno R, et al. Violence and risk of PTSD, major depression, substance abuse/dependence, and comorbidity: Results from the National Survey of Adolescents. *J Consult Clin Psychol* 2003;71:692–700.

32. Glaesser J, Neuner F, Lütgehetmann R, et al. Posttraumatic stress disorder in patients with traumatic brain injury. *BMC Psychiatry* 2004;4:1.

33. Bryant RA, Creamer M, O’Donnell M, et al. Post-traumatic amnesia and the nature of post-traumatic stress disorder after mild traumatic brain injury. *J Int Neuropsychol Soc* 2009;15:862–7.

34. King NS. Post-traumatic stress disorder and head injury as a dual diagnosis: “islands” of memory as a mechanism. *Journal of Neurology, Neurosurgery & Psychiatry* 1997;62:82–4.

35. Finset A, Andersson S. Coping strategies in patients with acquired brain injury: relationships between coping, apathy, depression and lesion location. *Brain Inj* 2000;14:987–905.