

HHS Public Access

Author manuscript *J Neurosurg*. Author manuscript; available in PMC 2017 February 01.

Published in final edited form as:

J Neurosurg. 2016 February ; 124(2): 511–526. doi:10.3171/2015.2.JNS14503.

Traumatic brain injury is associated with subsequent neurologic and psychiatric disease: a meta-analysis

David C. Perry, MD¹, Virginia E. Sturm, PhD¹, Matthew J. Peterson, PhD^{4,5}, Carl F. Pieper, DPH⁶, Thomas Bullock, BA³, Bradley F. Boeve, MD⁸, Bruce L. Miller, MD¹, Kevin M. Guskiewicz, PhD, ATC⁹, Mitchel S. Berger, MD², Joel H. Kramer, PsyD¹, and Kathleen A. Welsh-Bohmer, PhD⁷

¹Department of Neurology, University of California, San Francisco, San Francisco, CA, USA

²Department of Neurosurgery, University of California, San Francisco, San Francisco, CA, USA

³School of Medicine, University of California, San Francisco, San Francisco, CA, USA

⁴Geriatric Research, Education, and Clinical Center, Veterans Affairs Medical Center, Duke University, Durham, NC, USA

⁵Department of Medicine, Duke University, Durham, NC, USA

⁶Department of Biostatistics and Bioinformatics, Duke University, Durham, NC, USA

⁷Department of Psychiatry and Behavioral Sciences, Duke University, Durham, NC, USA

⁸Department of Neurology, Mayo Clinic, Rochester, MN, USA

⁹Department of Exercise and Sport Science, University of North Carolina, Chapel Hill, NC, USA

Abstract

Object—Mild traumatic brain injury (TBI) has been proposed as a risk factor for development of Alzheimer's disease, Parkinson's disease, depression, and other illnesses. This study's objective was to determine the association of prior mild TBI with subsequent diagnosis (i.e., at least one year post-injury) of neurologic or psychiatric disease.

Methods—All studies from 1995–2012 reporting TBI as a risk factor for diagnoses of interest were identified by searching PubMed, study references, and review articles. Reviewers abstracted the data and assessed study design and characteristics.

Results—57 studies met inclusion criteria. A random effects meta-analysis revealed a significant association of prior TBI with subsequent neurologic and psychiatric diagnosis. The pooled odds ratio (OR) for TBI on development of any illness was 1.67 (95% CI 1.44–1.93, p<.001). Prior TBI was independently associated with both neurologic [OR 1.55 (95% CI 1.31–1.83, p<.001)] and psychiatric [OR 2.00 (95% CI 1.50–2.66, p<.001)] outcomes. Analyses of individual diagnoses

Corresponding author: David C. Perry, MD, University of California, San Francisco, Memory and Aging Center MC: 1207, 675 Nelson Rising Lane, Suite 190, San Francisco, CA 94158, Telephone: (415) 476-8678, Fax: (415) 476-1816, david.perry2@ucsf.edu. **DISCLOSURES**

Dr. Perry reports no competing interests. Dr. Pieper reports no competing interests. Mr. Bullock reports no competing interests. Dr. Guskiewicz reports no competing interests. Dr. Berger reports no competing interests. Dr. Kramer reports no competing interests.

found higher odds of Alzheimer's disease, Parkinson's disease, mild cognitive impairment, depression, mixed affective disorders, and bipolar disorder in individuals with previous TBI compared to those without TBI. This association was present when examining only studies of mild TBI and when considering the influence of study design and characteristics. Analysis of a subset of studies found no evidence that multiple TBIs were associated with higher odds of disease than a single TBI.

Conclusions—History of TBI, including mild TBI, is associated with the development of neurologic and psychiatric illness. This indicates that either TBI is a risk factor for heterogeneous pathologic processes or that TBI may contribute to a common pathologic mechanism.

Keywords

dementia; psychiatry; head injury; meta-analysis

INTRODUCTION

Since the 1928 description of the "punch drunk" condition,⁴⁸ there has been speculation about a connection between traumatic brain injury (TBI) and late-life neurologic or psychiatric illness. Though this syndrome was later referred to as "dementia pugilistica" because it was thought to uniquely affect boxers,¹⁴ an accumulation of cases in recent years have suggested that repeated brain injury in other sports such as football, soccer, and wrestling might also predispose to neurodegenerative disease⁵² and that non-sports-related TBI, such as that sustained on the battlefield, can lead to this same illness.³⁰ It has recently been proposed that a history of even minor brain injury may predispose certain individuals to develop this pathologic process, now referred to as "chronic traumatic encephalopathy" or CTE⁵². The presentation of CTE is variable and may include neurologic and/or psychiatric manifestations. The current CTE literature suggests two common syndromes: a behavior and mood predominant illness, frequently accompanied by paranoia, which would be diagnosed as psychiatric illness and a predominantly cognitive disorder that is frequently diagnosed as Alzheimer's disease ⁸². A third syndrome, which was emphasized by the prior literature on boxers includes motor dysfunction with parkinsonism¹⁴. Some CTE cases have also been described with motor neuron disease^{53, 54}. Epidemiological study of CTE has been significantly limited since it is a pathological, rather than clinical diagnosis, and its presence can only be definitively confirmed after death. There is accumulating evidence, however, that CTE may be a pathologic process that unites seemingly disparate clinical syndromes and reflects a shared vulnerability to cognitive-behavioral-motor dysfunction. Recent studies have found support for a relationship between TBI and risk for later development of these individual neurologic and psychiatric syndromes. Since James Parkinson himself theorized a causative link to trauma in 1817, there has been continuing debate regarding the relationship between TBI and Parkinson's disease,¹⁹ with many^{17, 29, 87} but not all^{3, 43, 49} studies finding a positive association. Epidemiological studies investigating the risk of Alzheimer's disease after TBI have also shown mixed results. Meta-analyses of these studies in 199158 and in 2003²⁴ have shown an elevated risk. Prior TBI has also been associated with a significantly elevated risk of frontotemporal dementia⁷⁰ and although a prior meta-analysis of the risk of TBI on development of amyotrophic lateral sclerosis (ALS) showed a mildly elevated risk,¹¹

others have disputed the connection.⁹³ Although psychiatric symptoms (e.g., depression and anxiety) are common acutely after TBI,^{6, 35, 40} whether there are protracted psychiatric sequelae from earlier-life TBI remains poorly understood.⁹⁶

Our aim was to determine the association of mild TBI with the later development of those neurologic and psychiatric illnesses that have previously been linked to TBI and are potential manifestations of CTE. To investigate the wide range of disorders associated with prior TBI, we reviewed the literature examining TBI and subsequent neurologic or psychiatric diagnoses and performed a meta-analysis according to current guidelines^{56, 84}. Consistent with the notion that mild TBI may make certain individuals vulnerable to a number of neurologic and psychiatric conditions, we hypothesized that there would be an association between all diagnoses and a history of TBI, including mild TBI.

METHODS

Identification of the studies

Searches were conducted in Medline (1995 to February 2012) using a comprehensive search strategy. We used two components in each search: component A identified papers with keywords "craniocerebral trauma," "head injury," "brain injury," or "concussion." This was combined with component B or component C. Component B identified papers pertaining to the neurologic disorders of interest (i.e., "neurodegenerative diseases," "mild cognitive impairment," "Alzheimer," "Parkinson," "frontotemporal dementia," "amyotrophic lateral sclerosis," "vascular dementia," or "dementia"), and component C identified papers pertaining to the psychiatric illnesses of interest (i.e., "anxiety disorders," "mood disorders," or "schizophrenia and disorders with psychotic features"). We limited our search to papers in English and humans.

Three additional steps were taken to ensure search comprehensiveness: (1) references from included papers were reviewed, (2) to avoid any bias toward positive results inherent in the search strategy an additional search for "risk factors" for each diagnosis was performed to capture studies with weak or null findings that did not include our search terms in their title, abstract or keywords, (3) the citation lists in review papers were examined. For papers in which the required metrics were not easily identified the authors were contacted. A pair of reviewers (a neurologist and a neuropsychologist) discussed all papers at each stage of the process (Figure 1). Concordance between the reviewers for determining study inclusion was high; in cases of disagreement, studies were discussed until a consensus decision was reached. Ethics committee approval was not needed for this study as it included only analysis of previously published data.

Broad inclusion criteria

We first applied broad inclusion criteria (developed by a team of expert neurologists, neurosurgeons, and neuropsychologists) to select papers for further review.

- Original, peer-reviewed research articles (no case reports)
- Adult subjects over 18 years of age at the time of evaluation (not TBI)

- Presence of TBI without accompanying structural lesion (e.g., subdural hematoma or penetrating brain injury). Though our goal was to specifically examine the effect of mild TBI, in order to capture all pertinent studies, at this search stage we broadly included studies employing the various definitions and labels that are used to refer to minor head trauma (e.g., concussion).
- Presence of neurologic or psychiatric diagnosis
- TBI occurred before the diagnosis of the neurologic or psychiatric disorder (with at least 12 months between the TBI and outcome diagnosis, if specified)

Narrow inclusion criteria

Papers that met broad inclusion criteria were next reviewed in detail. In addition to ensuring adherence to broad criteria, we also confirmed that they met narrow inclusion criteria. If some subjects in a study were reported to have structural lesions, but they could be separated from those without lesions, we only included subjects with mild TBI.

- Presence of neurologic or psychiatric disorder. For neurologic disorders, studies must have utilized consensus diagnostic criteria or clinical evaluation. For psychiatric disorders, diagnoses were based on either criteria (e.g., DSM-IV) or scores from standardized measures (e.g., Beck Depression Inventory).
- **2.** *Inclusion of a control group.* Included studies were cross-sectional, cohort, or case-control studies in which all subjects underwent identical assessment and diagnostic procedures.
- 3. *TBI preceded neurologic or psychiatric diagnosis.* We excluded studies that reported that the diagnosis of the neurologic or psychiatric disorder had been made less than 12 months post-TBI. For studies in which the date of the TBI was not reported, we included studies of subjects with neurologic or psychiatric illness who were asked about TBI earlier in life.
- 4. No redundant subjects across studies. In cases where multiple papers used the same study cohort we included the most recent papers to capture the largest sample size. If multiple outcome diagnoses were reported in one paper, we included each odds ratio (OR) if the diagnoses were mutually exclusive. If the diagnoses were not mutually exclusive, in the analyses that examined the association of TBI with any neurologic or psychiatric outcome, we chose the broader diagnosis (e.g., dementia was preferred over Alzheimer's disease) or, if that distinction was not possible, we chose the diagnosis with the larger number of subjects.

Assessment of study characteristics

We recorded additional data regarding factors that could influence the relationship between TBI and outcome diagnoses. These included (1) the rigor with which each study employed a 12-month TBI-outcome diagnosis interval, (2) the TBI characteristics required in each study (e.g., whether subjects met accepted criteria for mild TBI or had any individual symptoms such as loss of consciousness), (3) whether the TBI diagnosis was based on patient or informant self-report as opposed to being made by a clinician, derived from medical records,

or based on diagnostic criteria, (4) the study design (cohort, case-control, or cross-sectional), and (5) whether information was provided regarding the number of TBIs sustained by each subject. These data were used in subgroup analyses geared towards assessing whether study characteristics influenced the meta-analysis results.

Statistical Analysis

Primary analyses—The effect of interest for this meta-analysis was the pooled OR. For the majority of the studies (51/57), unadjusted ORs were directly calculated from data extraction. Standard errors were calculated from the logarithm of the OR to allow for symmetry of the estimate on both sides of unity.²³ Where sample sizes were not available the published unadjusted ORs were used. We then applied standard meta-analytic techniques,³⁴ including weighted estimates of the pooled OR with a 95% confidence interval (CI). For those studies where the raw cell frequencies did not exist and only the standard error of the OR was available, to provide appropriate weighting of the study in the meta-analysis, the standard error of the OR was transformed to the standard error of the logarithm of the OR by linear interpolation. To determine whether there was significant variation among studies, tests of heterogeneity were performed.³⁴ All analyses were conducted using SAS v9.3 (Cary, NC).

Subgroup analyses—Since the overall analysis was inclusive of various TBI definitions and study characteristics, we next conducted seven additional subgroup analyses to examine whether our results differed when pooling studies with more uniformity of TBI assessment, TBI diagnostic criteria, and study design. When possible, we selected out only those subjects from the total study that met criteria for each subgroup analysis. The result is that for some studies a different number of subjects was included in the overall analysis compared to each subgroup analysis.

Subgroup 1: Effect of time interval between TBI and diagnosis

 Clearest interval - To ensure that studies with less stringent guidelines about timing of TBI were not significantly impacting our results, we excluded studies with the possibility that some subjects had a less than 12 month interval between TBI and diagnosis.

Subgroups 2–4: Effect of TBI features and severity

- Brief loss of consciousness This subgroup included only studies that required that loss of consciousness not exceed 30 minutes. This is the maximum duration established in the mild TBI criteria of the American Congress of Rehabilitation Medicine, Centers for Disease Control, and World Health Organization^{10, 42, 59}.
- **3.** Required loss of consciousness We included only studies that required brain injury with loss of consciousness. This subgroup considered the effect of TBI with a uniform minimum severity.
- **4.** Any mild TBI feature In order to exclude extremely mild or asymptomatic brain injury we performed an analysis including only those studies that required the brain injury be accompanied by any one (or more than one) common feature

of mild TBI, including loss of consciousness, post-traumatic amnesia, Glasgow Coma Scale (GCS) score 13, focal neurological deficit, altered mental status, brain injury requiring medical care, or symptoms of the postconcussive syndrome (e.g., headache, dizziness, nausea, photo- or phonophobia, fatigue, sleep difficulty, blurred vision).

Subgroups 5–6: Effect of study design

- 5. Excluding self-report In order to assess the impact of recall bias we conducted an additional analysis excluding studies with self-reported TBI.
- **6.** Cohort studies To eliminate recall bias we also performed an analysis including only cohort studies (rather than cross-sectional or case-control).

Subgroup 7: Effect of number of TBI

7. Repeated injury - Because we were also interested in whether there is a dose effect of TBI on development of later illness, we conducted an additional analysis in which we calculated the odds of neurologic or psychiatric diagnosis in subjects with more than one TBI compared to a single TBI using a subset of studies that provided this information.

Publication bias analysis—To assess for the effect of publication bias on our results we used the Egger method¹⁸ to examine whether the logarithm of the included ORs are predicted by the standard error, which reflects the sample size. We visually examined funnel plots of OR against sample size and the logarithm of the OR against the standard error of the logarithm of the OR and quantified the degree of bias by multiple regression. Using standard error rather than sample size in funnel plots may provide a more accurate visual depiction of whether bias is present⁸³.

RESULTS

57 papers met narrow inclusion criteria and were used in the meta-

analyses

Among the included papers, a sufficient number were found to apply meta-analytic methods for the diagnoses of dementia, Alzheimer's disease, Parkinson's disease, ALS, mild cognitive impairment, depression, psychotic disorders, bipolar disorder, and mixed affective disorder (a combined group of depression and anxiety). Insufficient numbers of studies were found to calculate a pooled OR for frontotemporal dementia, vascular dementia, or anxiety disorders. There was significant heterogeneity among studies (Q = 381.99, df = 58, p<.001), justifying the use of the random effects meta-analysis.

Prior TBI was associated with development of any of the neurologic and psychiatric illnesses of interest [OR 1.67 (95% CI 1.44–1.93), p<.0001]. This association was found for both neurologic [OR 1.55 (95% CI 1.31–1.83, p<.0001)] and psychiatric [OR 2.00 (95% CI 1.50–2.66, p<.0001)] disease in individuals with TBI, and was also found in the following diagnoses: Alzheimer's disease [OR 1.40 (95% CI 1.02–1.90, p<.05)], Parkinson's disease

[OR 1.45 (95% CI 1.18–1.78, *p*<.001)], mild cognitive impairment [OR 2.69 (95% CI 1.51–4.77, *p*<.001)], depression [OR 2.14 (95% CI 1.65–2.77, *p*<.0001)], bipolar disorder [OR 1.85 (95% CI 1.17–2.94, *p*<.01)], and mixed affective disorder [OR 1.84 (95% CI 1.50–2.66, *p*<.0001)]. See Table 1 and Figure 2.

Analyses of subgroups revealed a robust relationship between TBI and remote neurologic and psychiatric outcomes. The studies included in each subgroup analysis are specified in Table 1. Table 2 includes the features reported in each study regarding the time interval between TBI and diagnosis, the TBI features and severity, and the study design. Results of the subgroup analyses are reported in Table 3. Overall odds and the independent OR for neurologic, but not psychiatric disease remained significant when only including studies with the clearest greater than 12-month interval between TBI and diagnosis (Subgroup 1). The overall OR was significant among those studies that adhered to mild TBI criteria limiting duration of loss of consciousness to less than 30 minutes (Subgroup 2). The overall OR and OR for any of the studied neurologic and psychiatric diagnoses were also significant when only including studies that required loss of consciousness (Subgroup 3). When including studies that required the presence of at least one mild TBI symptom (Subgroup 4), the overall OR and OR for any of the neurologic and all psychiatric diagnoses of interest remained significant. After eliminating the studies with TBI diagnoses based on self-report (Subgroup 5), the overall OR and OR for neurologic disorders remained significant, though the OR for psychiatric outcomes no longer reached significance. When only cohort studies were included (Subgroup 6), the OR for neurologic outcomes was not significant though the overall OR and OR for psychiatric illness remained significant. The odds were not higher among those that reported more than one TBI compared to those with a single injury (Subgroup 7).

Publication bias analyses did not show evidence of bias in the included studies. Visual inspection of a funnel plot based on sample size showed that three studies with large samples strongly influenced the appearance (Figure 3A). When these studies are removed a more expected funnel shape is appreciated (Figure 3B). Regression indicates that the effects size (the logarithm of the OR) is not significantly predicted by the standard error when all studies are included (F(1,60)=3.08, p=.08) or when the three large sample studies are excluded (F(1,57)=1.11, p=.30, Figure 3C).

DISCUSSION

This meta-analysis supports an association between prior TBI and later diagnosis of the relevant neurologic or psychiatric diseases. This association was found independently for both neurologic and psychiatric outcomes. Alzheimer's disease, Parkinson's disease, mild cognitive impairment, depression, mixed affective disorders, and bipolar disorder showed a statistically significant association with prior TBI. The magnitude of effect is comparable across diagnoses, with mild cognitive impairment, depression, and bipolar disorder having the highest OR among those results that reached significance. The OR of Alzheimer's disease in this analysis is comparable to the findings of prior meta-analyses.^{24, 58} The OR of ALS was among the highest in the study, and there was some evidence of an association of TBI with dementia and psychotic disorders, but these did not reach statistical significance.

The overall combined OR for the selected neurologic and psychiatric illnesses and for neurologic illness independently in individuals with TBI remained significant when including only articles that explicitly specified a minimum 12-month interval between TBI and outcome diagnosis. The magnitude of association with psychiatric illness, however, did not remain significant. These results suggest that there may be a different time course in which psychiatric and neurologic symptoms manifest after TBI. While psychiatric symptoms are common in the acute phase after mild TBI^{6, 21, 35, 40} and some of these may be short-lived manifestations of the injury, others may reflect a more sustained susceptibility to mental illness. The results of this study suggest that TBI is a risk factor for both remote psychiatric and neurologic disease and are consistent with the possibility that both types of illness arise secondary to a common shared pathologic mechanism.

We conducted additional subgroup analyses to determine whether TBI characteristics or methodological factors would influence our findings. The overall OR of TBI remained significant when including only studies that required adherence to typical loss of consciousness criteria for mild TBI, the presence of any specific mild TBI symptom, or loss of consciousness. Though TBI definitions varied widely among studies, these additional analyses support an association of mild TBI with the studied neurologic and psychiatric outcomes. A significant OR for combined neurologic and psychiatric outcomes was also found when eliminating studies that used self-reported diagnosis of TBI and when including only cohort studies. Though statistical significance was lost when assessing the association with psychiatric outcomes when eliminating self-report and the odds of neurologic outcomes among cohort studies, the magnitudes of the ORs were largely consistent with the main analysis, and the change in significance is likely due to the small number of articles in these analyses and resulting loss of power rather than reflecting a weaker association due to recall bias, though this cannot be excluded. Our analyses also suggest that the finding of an association with TBI is unlikely to be due to publication bias, though low power may affect the publication bias test.

The results of this meta-analysis support an association of illness with a single TBI. A relevant associated question is whether this effect is compounded by multiple TBIs. In our analysis of multiple head traumas, the results do not show strong evidence for elevated odds of illness associated with having more than one head trauma compared to a single TBI. Given that only six studies were included in this analysis, lower power may have influenced these results. More research on the relationship between TBI exposure and diagnostic outcomes is needed.

The magnitude of the OR of TBI in this meta-analysis is relatively modest, but comparable to other strongly implicated risk factors. For example, for Alzheimer's disease, the Apolipoprotein E e4 allele is associated with an OR of 1.80–9.05,⁴¹ and obesity with an OR of 1.80.⁴ The OR for pesticide exposure and Parkinson's disease is 1.94.⁶⁶ Therefore, the presence of a risk factor in an individual does not indicate an inevitable development of disease. The ORs found in this study suggest that others factors modify an individual's susceptibility to develop a neuropsychiatric disorder after TBI. These factors are largely unknown and worthy of further investigation.

The fact that multiple neurodegenerative and psychiatric diagnoses are associated with the same exposure raises questions about possible mechanisms of shared vulnerability. Trauma could predispose the brain to different types of neurodegeneration through common mechanisms such as oxidative stress and microglial activation^{77, 99} or induction of plasma proteins associated with degeneration such as MCP-1.³⁶ Trauma might also activate molecular pathways leading to specific degenerative diseases, such as the finding that Alzheimer's disease-associated proteins including beta amyloid, beta secretase, presenilin-1, and caspase-3 accumulate in axons of brain injured animals.¹² Cleaved forms of the tau protein, which is associated with Alzheimer's disease and frontotemporal lobar degeneration, accumulate after trauma²⁶ and tau abnormalities after trauma have been found to be independent of beta amyloid effects.⁸⁹ The nature of the TBI could also influence the clinical presentation in an individual. For example, boxers may suffer from more torsional injury that could injury brainstem structures such as the substantia nigra, leading to parkinsonism⁸². Genetic variation could also help to explain the susceptibility of individuals to late-life effects of TBI. For example, apolipoprotein E, which is associated with risk of Alzheimer's disease, has shown a variable interaction with mild TBI.^{50, 63, 88}

An alternate explanation for the association across diagnostic groups is that the various clinical presentations could be different expressions of a common pathology^{28, 78}. Although CTE has been described as a distinct pathological process, the clinical characterization is not clearly established, and case reports suggest cognitive, motor, and psychiatric presentations. This phenotypic variability could lead to a diagnosis of dementia, Parkinson's disease, motor neuron disease, or primary psychiatric illness in different individuals. A study of causes of death among retired National Football League players found a three-fold higher rate of dying from neurodegenerative disease compared to the typical population frequency, with Alzheimer's disease and ALS being the most overrepresented⁴⁴, which would be consistent either a shared vulnerability hypothesis across neurodegenerative diseases or a common pathology. This meta-analysis examined clinical, not pathological, studies. Thus, it is unknown whether any of the subjects would have shown characteristic CTE pathology rather than (or in addition to) the more typical neuropathological features associated with their syndromes.

Among the articles that were reviewed, several addressed the association between TBI and clinical outcomes among athletes. These articles assessed the risk of Parkinson's disease among retired Thai traditional boxers,⁴⁶ depression and dementia among retired football players,^{32, 33} ALS or chronic encephalopathy among soccer players,^{13, 39} and chronic TBI in boxers.³⁸ Only two of the articles^{32, 33} that directly evaluated TBI in sports met strict inclusion criteria for this study. The ability of this meta-analysis to inform the questions surrounding the long-term consequences of sports-related mild TBI is therefore limited by this lack in the existing literature. Further longitudinal studies among athletes with appropriate control groups, characterization of head injuries (including severity, number, and exposure to repetitive subconcussive trauma), and assessment of late-life neurologic and psychiatric outcomes will be needed to address this question.

Several limitations of this meta-analysis warrant consideration. One is the possible bias of the included studies. We took several steps to mitigate this possibility. Our search strategy

included a variety of epidemiological studies that focused on many possible risk factors, not just TBI, thereby capturing negative studies that might otherwise have not been published. Our formal analyses also did not support publication bias. Although the strict inclusion criteria should reduce this possibility, variation in the studies themselves (e.g., different criteria for diagnosis of illness, or co-morbid environmental and genetic factors of the study population) limits the generalizability of the results. Variable study quality could also have resulted in heterogeneity, and it is possible that the presence of other confounding factors could have led to the observed association between TBI and later clinical outcomes. For example, patients who sustain a TBI as a result of a fall or motor vehicle accident may have other medical comorbidities (e.g., vascular disease or substance abuse) or differences in socioeconomic status that could predispose to neurologic or psychiatric illness. Another possibility is that the TBI itself could lead to injury of change in lifestyle that could modify risk of a mood disorder. Finally, ill patients who fall and suffer TBI may also undergo more medical testing and therefore be more likely to receive one of these neurologic or psychiatric diagnoses. Only English language studies were reviewed, which could have led to exclusion of some relevant studies. In spite of our criteria regarding an interval between TBI and illness onset, an alternative explanation for the observed association is that some head injuries may have been early manifestations of neurologic or psychiatric disease rather than an independent predisposing factor for illness. The authors of one of the studies concluded this reverse causality was responsible for their findings. They stratified the interval between TBI and diagnosis and found the association between TBI and Parkinson's was no longer present when only looking at TBI that occurred greater than 10 years prior to diagnosis⁷¹.

A major strength of this meta-analysis was the inclusion of a variety of different neurologic and psychiatric outcomes rather than a single diagnosis. By focusing on diagnoses rather than self-reported symptoms or performance on cognitive tests, this study assessed outcomes of sufficient magnitude to affect quality of life. The included studies also come from countries around the world, allowing for more generalizable results. The literature search was comprehensive, making this a rigorous examination of the topic.

CONCLUSIONS

This study supports an association of TBI, including mild TBI, on subsequent development of neurologic and psychiatric illness, including Alzheimer's disease, Parkinson's disease, mild cognitive impairment, depression, mixed affective disorders, and bipolar disorder. Due to limitations and heterogeneity in the existing studies, prospective studies with uniform assessment will be needed to confirm this result and determine the risk conferred by the number and severity of TBI in different settings, such as sports or the military.

Acknowledgments

Dr. Sturm is supported by National Institute on Aging 1K23AG040127. Dr. Peterson is supported by National Cancer Institute Award KM1CA156687. Dr. Boeve receives research support from the National Institute on Aging (P50 AG016574, U01 AG006786, R01 AG032306, and R01 AG041797) and the Mangurian Foundation. Dr. Miller is funded by NIH grants P50AG023501, P01AG019724, P50 AG1657303, and the state of California. Dr. Welsh-Bohmer received funding from the National Institute of Aging (P30 AG28377), from private donors to the Joseph & Kathleen Bryan Alzheimer's Disease Center at Duke University, and from Takeda and Zinfandel Pharmaceutical companies.

REFERENCES

- 1. AbdelMalik P, Husted J, Chow EW, Bassett AS. Childhood head injury and expression of schizophrenia in multiply affected families. Arch Gen Psychiatry. 2003; 60:231-236. [PubMed: 12622655]
- 2. Bachman DL, Green RC, Benke KS, Cupples LA, Farrer LA. MIRAGE Study Group. Comparison of alzheimer's disease risk factors in white and african american families. Neurology. 2003; 60:1372-1374. [PubMed: 12707449]
- 3. Baldereschi M, Di Carlo A, Vanni P, Ghetti A, Carbonin P, Amaducci L, et al. Lifestyle-related risk factors for parkinson's disease: A population-based study. Acta Neurol Scand. 2003; 108:239-244. [PubMed: 12956856]
- 4. Beydoun MA, Beydoun HA, Wang Y. Obesity and central obesity as risk factors for incident dementia and its subtypes: A systematic review and meta-analysis. Obes Rev. 2008; 9:204–218. [PubMed: 18331422]
- 5. Binazzi A, Belli S, Uccelli R, Desiato MT, Talamanca IF, Antonini G, et al. An exploratory casecontrol study on spinal and bulbar forms of amyotrophic lateral sclerosis in the province of rome. Amyotroph Lateral Scler. 2009; 10:361–369. [PubMed: 19922125]
- 6. Bombardier CH, Fann JR, Temkin NR, Esselman PC, Barber J, Dikmen SS. Rates of major depressive disorder and clinical outcomes following traumatic brain injury. JAMA. 2010; 303:1938–1945. [PubMed: 20483970]
- 7. Boston PF, Dennis MS, Jagger C. Factors associated with vascular dementia in an elderly community population. Int J Geriatr Psychiatry. 1999; 14:761–766. [PubMed: 10479748]
- 8. Bower JH, Maraganore DM, Peterson BJ, McDonnell SK, Ahlskog JE, Rocca WA. Head trauma preceding PD: A case-control study. Neurology. 2003; 60:1610–1615. [PubMed: 12771250]
- 9. Bryant RA, O'Donnell ML, Creamer M, McFarlane AC, Clark CR, Silove D. The psychiatric sequelae of traumatic injury. Am J Psychiatry. 2010; 167:312–320. [PubMed: 20048022]
- 10. Carroll L, Cassidy J, Holm L, Kraus J, Coronado V. Methodological issues and research recommendations for mild traumatic brain injury: The WHO collaborating centre task force on mild traumatic brain injury. J Rehabil Med. 2004; 43:113-125. [PubMed: 15083875]
- 11. Chen H, Richard M, Sandler DP, Umbach DM, Kamel F. Head injury and amyotrophic lateral sclerosis. Am J Epidemiol. 2007; 166:810-816. [PubMed: 17641152]
- 12. Chen XH, Siman R, Iwata A, Meaney DF, Trojanowski JQ, Smith DH. Long-term accumulation of amyloid-beta, beta-secretase, presenilin-1, and caspase-3 in damaged axons following brain trauma. Am J Pathol. 2004; 165:357–371. [PubMed: 15277212]
- 13. Chio A, Benzi G, Dossena M, Mutani R, Mora G. Severely increased risk of amyotrophic lateral sclerosis among italian professional football players. Brain. 2005; 128:472-476. [PubMed: 15634730]
- 14. Corsellis JA, Bruton CJ, Freeman-Browne D. The aftermath of boxing. Psychol Med. 1973; 3:270-303. [PubMed: 4729191]
- 15. De Michele G, Filla A, Volpe G, De Marco V, Gogliettino A, Ambrosio G, et al. Environmental and genetic risk factors in parkinson's disease: A case-control study in southern italy. Mov Disord. 1996; 11:17–23. [PubMed: 8771062]
- 16. DelBello MP, Soutullo CA, Zimmerman ME, Sax KW, Williams JR, McElroy SL, et al. Traumatic brain injury in individuals convicted of sexual offenses with and without bipolar disorder. Psychiatry Res. 1999; 89:281-286. [PubMed: 10708275]
- 17. Dick FD, De Palma G, Ahmadi A, Scott NW, Prescott GJ, Bennett J, et al. Environmental risk factors for parkinson's disease and parkinsonism: The geoparkinson study. Occup Environ Med. 2007; 64:666-672. [PubMed: 17332139]
- 18. Egger M, Davey Smith G, Schneider M, Minder C. Bias in meta-analysis detected by a simple, graphical test. BMJ. 1997; 315:629-634. [PubMed: 9310563]
- 19. Factor SA, Sanchez-Ramos J, Weiner WJ. Trauma as an etiology of parkinsonism: A historical review of the concept. Mov Disord. 1988; 3:30-36. [PubMed: 3050470]

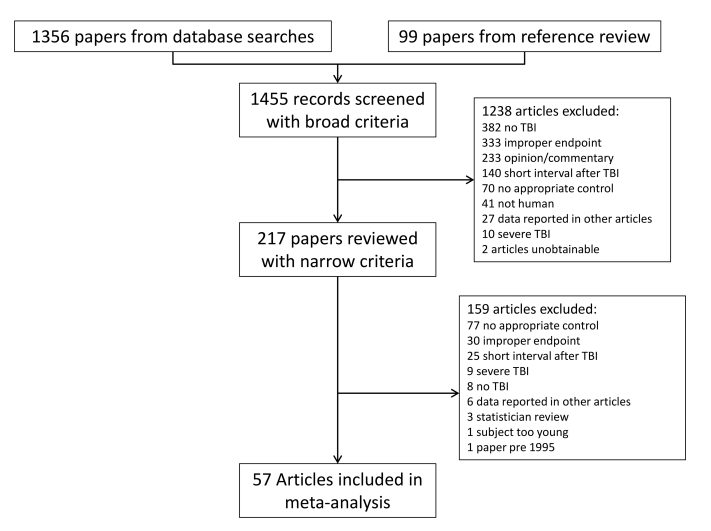
- Fann JR, Burington B, Leonetti A, Jaffe K, Katon WJ, Thompson RS. Psychiatric illness following traumatic brain injury in an adult health maintenance organization population. Arch Gen Psychiatry. 2004; 61:53–61. [PubMed: 14706944]
- Fedoroff JP, Starkstein SE, Forrester AW, Geisler FH, Jorge RE, Arndt SV, et al. Depression in patients with acute traumatic brain injury. Am J Psychiatry. 1992; 149:918–923. [PubMed: 1609872]
- 22. Fischer P, Zehetmayer S, Jungwirth S, Weissgram S, Krampla W, Hinterberger M, et al. Risk factors for alzheimer dementia in a community-based birth cohort at the age of 75 years. Dement Geriatr Cogn Disord. 2008; 25:501–507. [PubMed: 18446027]
- 23. Fleiss, JL.; Levin, BA.; Paik, MC. Statistical Methods for Rates and Proportions. New York City: Wiley; 2003.
- 24. Fleminger S, Oliver DL, Lovestone S, Rabe-Hesketh S, Giora A. Head injury as a risk factor for Alzheimer's disease: The evidence 10 years on; a partial replication. Journal of Neurology, Neurosurgery & Psychiatry. 2003; 74:857–862.
- Forster DP, Newens AJ, Kay DW, Edwardson JA. Risk factors in clinically diagnosed presenile dementia of the alzheimer type: A case-control study in northern england. J Epidemiol Community Health. 1995; 49:253–258. [PubMed: 7629459]
- 26. Gabbita SP, Scheff SW, Menard RM, Roberts K, Fugaccia I, Zemlan FP. Cleaved-tau: A biomarker of neuronal damage after traumatic brain injury. J Neurotrauma. 2005; 22:83–94. [PubMed: 15665604]
- 27. Gao S, Jin Y, Unverzagt FW, Liang C, Hall KS, Ma F, et al. Correlates of depressive symptoms in rural elderly chinese. Int J Geriatr Psychiatry. 2009; 24:1358–1366. [PubMed: 19347839]
- Gavett BE, Stern RA, Cantu RC, Nowinski CJ, McKee AC. Mild traumatic brain injury: A risk factor for neurodegeneration. Alzheimers Res Ther. 2010; 2:18. [PubMed: 20587081]
- Goldman SM, Tanner CM, Oakes D, Bhudhikanok GS, Gupta A, Langston JW. Head injury and parkinson's disease risk in twins. Ann Neurol. 2006; 60:65–72. [PubMed: 16718702]
- Goldstein LE, Fisher AM, Tagge CA, Zhang XL, Velisek L, Sullivan JA, et al. Chronic traumatic encephalopathy in blast-exposed military veterans and a blast neurotrauma mouse model. Sci Transl Med. 2012; 4:134ra60.
- Guo Z, Cupples LA, Kurz A, Auerbach SH, Volicer L, Chui H, et al. Head injury and the risk of AD in the MIRAGE study. Neurology. 2000; 54:1316–1323. [PubMed: 10746604]
- Guskiewicz KM, Marshall SW, Bailes J, McCrea M, Cantu RC, Randolph C, et al. Association between recurrent concussion and late-life cognitive impairment in retired professional football players. Neurosurgery. 2005; 57:719–726. discussion 719–26. [PubMed: 16239884]
- Guskiewicz KM, Marshall SW, Bailes J, McCrea M, Harding HP Jr, Matthews A, et al. Recurrent concussion and risk of depression in retired professional football players. Med Sci Sports Exerc. 2007; 39:903–909. [PubMed: 17545878]
- 34. Hedges, LV.; Olkin, I. Statistical Methods for Meta-Analysis. Orlando, FL: Academic Press; 1985.
- 35. Hibbard MR, Uysal S, Kepler K, Bogdany J, Silver J. Axis I psychopathology in individuals with traumatic brain injury. J Head Trauma Rehabil. 1998; 13:24–39. [PubMed: 9651237]
- 36. Ho L, Zhao W, Dams-O'Connor K, Tang CY, Gordon W, Peskind ER, et al. Elevated plasma MCP-1 concentration following traumatic brain injury as a potential "predisposition" factor associated with an increased risk for subsequent development of alzheimer's disease. J Alzheimers Dis. 2012; 31:301–313. [PubMed: 22543850]
- Holsinger T, Steffens DC, Phillips C, Helms MJ, Havlik RJ, Breitner JC, et al. Head injury in early adulthood and the lifetime risk of depression. Arch Gen Psychiatry. 2002; 59:17–22. [PubMed: 11779276]
- Jordan BD, Relkin NR, Ravdin LD, Jacobs AR, Bennett A, Gandy S. Apolipoprotein E epsilon4 associated with chronic traumatic brain injury in boxing. JAMA. 1997; 278:136–140. [PubMed: 9214529]
- Jordan SE, Green GA, Galanty HL, Mandelbaum BR, Jabour BA. Acute and chronic brain injury in united states national team soccer players. Am J Sports Med. 1996; 24:205–210. [PubMed: 8775122]

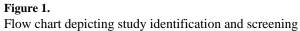
- 40. Jorge RE, Robinson RG, Starkstein SE, Arndt SV. Depression and anxiety following traumatic brain injury. J Neuropsychiatry Clin Neurosci. 1993; 5:369–374. [PubMed: 8286933]
- 41. Jun G, Naj AC, Beecham GW, Wang LS, Buros J, Gallins PJ, et al. Meta-analysis confirms CR1, CLU, and PICALM as alzheimer disease risk loci and reveals interactions with APOE genotypes. Arch Neurol. 2010; 67:1473–1484. [PubMed: 20697030]
- 42. Kay T, Harrington DE, Adams R, Anderson T, Berrol S, Cicerone K, et al. Definition of mild traumatic brain injury. J Head Trauma Rehabil. 1993; 8:86–87.
- 43. Kuopio AM, Marttila RJ, Helenius H, Rinne UK. Environmental risk factors in parkinson's disease. Mov Disord. 1999; 14:928–939. [PubMed: 10584666]
- 44. Lehman EJ, Hein MJ, Baron SL, Gersic CM. Neurodegenerative causes of death among retired national football league players. Neurology. 2012; 79:1970–1974. [PubMed: 22955124]
- 45. Lindsay J, Laurin D, Verreault R, Hébert R, Helliwell B, Hill GB, et al. Risk factors for Alzheimer's disease: A prospective analysis from the canadian study of health and aging. American Journal of Epidemiology. 2002; 156:445–453. [PubMed: 12196314]
- Lolekha P, Phanthumchinda K, Bhidayasiri R. Prevalence and risk factors of parkinson's disease in retired thai traditional boxers. Mov Disord. 2010; 25:1895–1901. [PubMed: 20669292]
- Malaspina D, Goetz RR, Friedman JH, Kaufmann CA, Faraone SV, Tsuang M, et al. Traumatic brain injury and schizophrenia in members of schizophrenia and bipolar disorder pedigrees. Am J Psychiatry. 2001; 158:440–446. [PubMed: 11229986]
- 48. Martland HS. Punch drunk. Journal of the American Medical Association. 1928; 91:1103–1107.
- Martyn CN, Osmond C. Parkinson's disease and the environment in early life. J Neurol Sci. 1995; 132:201–206. [PubMed: 8543949]
- Mayeux R, Ottman R, Maestre G, Ngai C, Tang MX, Ginsberg H, et al. Synergistic effects of traumatic head injury and apolipoprotein-epsilon 4 in patients with alzheimer's disease. Neurology. 1995; 45:555–557. [PubMed: 7898715]
- McCann SJ, LeCouteur DG, Green AC, Brayne C, Johnson AG, Chan D, et al. The epidemiology of parkinson's disease in an australian population. Neuroepidemiology. 1998; 17:310–317. [PubMed: 9778597]
- 52. McKee AC, Cantu RC, Nowinski CJ, Hedley-Whyte ET, Gavett BE, Budson AE, et al. Chronic traumatic encephalopathy in athletes: Progressive tauopathy after repetitive head injury. J Neuropathol Exp Neurol. 2009; 68:709–735. [PubMed: 19535999]
- McKee AC, Gavett BE, Stern RA, Nowinski CJ, Cantu RC, Kowall NW, et al. TDP-43 proteinopathy and motor neuron disease in chronic traumatic encephalopathy. J Neuropathol Exp Neurol. 2010; 69:918–929. [PubMed: 20720505]
- McKee AC, Stern RA, Nowinski CJ, Stein TD, Alvarez VE, Daneshvar DH, et al. The spectrum of disease in chronic traumatic encephalopathy. Brain. 2013; 136:43–64. [PubMed: 23208308]
- 55. Mehta KM, Ott A, Kalmijn S, Slooter AJ, van Duijn CM, Hofman A, et al. Head trauma and risk of dementia and alzheimer's disease: The rotterdam study. Neurology. 1999; 53:1959–1962. [PubMed: 10599765]
- 56. Moher D, Liberati A, Tetzlaff J, Altman DG. PRISMA Group. Preferred reporting items for systematic reviews and meta-analyses: The PRISMA statement. PLoS Med. 2009; 6:e1000097. [PubMed: 19621072]
- 57. Mollica RF, Lyoo IK, Chernoff MC, Bui HX, Lavelle J, Yoon SJ, et al. Brain structural abnormalities and mental health sequelae in south vietnamese ex-political detainees who survived traumatic head injury and torture. Arch Gen Psychiatry. 2009; 66:1221–1232. [PubMed: 19884610]
- 58. Mortimer JA, van Duijn CM, Chandra V, Fratiglioni L, Graves AB, Heyman A, et al. Head trauma as a risk factor for alzheimer's disease: A collaborative re-analysis of case-control studies EURODEM risk factors research group. Int J Epidemiol. 1991; 20(Suppl 2):S28–S35. [PubMed: 1833351]
- 59. National Center for Injury Prevention and Control (US). Report to Congress on Mild Traumatic Brain Injury in the United States: Steps to Prevent a Serious Public Health Problem. Centers for Disease Control and Prevention; 2003.

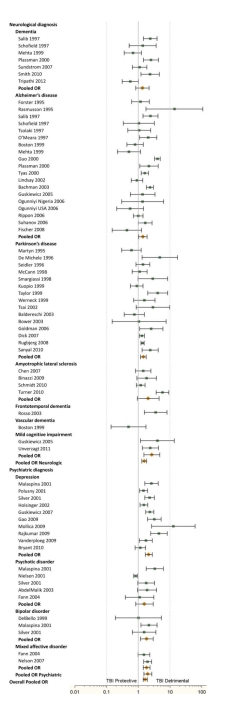
- Nelson LA, Rhoades DA, Noonan C, Manson SM. AI-SUPERPFP Team. Traumatic brain injury and mental health among two american indian populations. J Head Trauma Rehabil. 2007; 22:105–112. [PubMed: 17414312]
- Nielsen AS, Mortensen PB, O'Callaghan E, Mors O, Ewald H. Is head injury a risk factor for schizophrenia? Schizophr Res. 2002; 55:93–98. [PubMed: 11955968]
- 62. Ogunniyi A, Hall KS, Gureje O, Baiyewu O, Gao S, Unverzagt FW, et al. Risk factors for incident alzheimer's disease in african americans and yoruba. Metab Brain Dis. 2006; 21:235–240. [PubMed: 16850256]
- 63. O'Meara ES, Kukull WA, Sheppard L, Bowen JD, McCormick WC, Teri L, et al. Head injury and risk of alzheimer's disease by apolipoprotein E genotype. Am J Epidemiol. 1997; 146:373–384. [PubMed: 9290497]
- 64. Plassman BL, Havlik RJ, Steffens DC, Helms MJ, Newman TN, Drosdick D, et al. Documented head injury in early adulthood and risk of alzheimer's disease and other dementias. Neurology. 2000; 55:1158–1166. [PubMed: 11071494]
- 65. Polusny MA, Kehle SM, Nelson NW, Erbes CR, Arbisi PA, Thuras P. Longitudinal effects of mild traumatic brain injury and posttraumatic stress disorder comorbidity on postdeployment outcomes in national guard soldiers deployed to iraq. Arch Gen Psychiatry. 2011; 68:79–89. [PubMed: 21199967]
- 66. Priyadarshi A, Khuder SA, Schaub EA, Shrivastava S. A meta-analysis of parkinson's disease and exposure to pesticides. Neurotoxicology. 2000; 21:435–440. [PubMed: 11022853]
- Rajkumar AP, Thangadurai P, Senthilkumar P, Gayathri K, Prince M, Jacob KS. Nature, prevalence and factors associated with depression among the elderly in a rural south indian community. Int Psychogeriatr. 2009; 21:372–378. [PubMed: 19243657]
- Rasmusson DX, Brandt J, Martin DB, Folstein MF. Head injury as a risk factor in alzheimer's disease. Brain Inj. 1995; 9:213–219. [PubMed: 7606235]
- Rippon GA, Tang MX, Lee JH, Lantigua R, Medrano M, Mayeux R. Familial alzheimer disease in latinos: Interaction between APOE, stroke, and estrogen replacement. Neurology. 2006; 66:35–40. [PubMed: 16401842]
- Rosso SM, Landweer EJ, Houterman M, Donker Kaat L, van Duijn CM, van Swieten JC. Medical and environmental risk factors for sporadic frontotemporal dementia: A retrospective case-control study. J Neurol Neurosurg Psychiatry. 2003; 74:1574–1576. [PubMed: 14617722]
- Rugbjerg K, Ritz B, Korbo L, Martinussen N, Olsen JH. Risk of parkinson's disease after hospital contact for head injury: Population based case-control study. BMJ. 2008; 337:a2494. [PubMed: 19074944]
- 72. Salib E, Hillier V. Head injury and the risk of alzheimer's disease: A case control study. Int J Geriatr Psychiatry. 1997; 12:363–368. [PubMed: 9152722]
- Sanyal J, Chakraborty DP, Sarkar B, Banerjee TK, Mukherjee SC, Ray BC, et al. Environmental and familial risk factors of parkinsons disease: Case-control study. Can J Neurol Sci. 2010; 37:637–642. [PubMed: 21059511]
- 74. Schmidt S, Kwee LC, Allen KD, Oddone EZ. Association of ALS with head injury, cigarette smoking and APOE genotypes. J Neurol Sci. 2010; 291:22–29. [PubMed: 20129626]
- Schofield PW, Tang M, Marder K, Bell K, Dooneief G, Chun M, et al. Alzheimer's disease after remote head injury: An incidence study. J Neurol Neurosurg Psychiatry. 1997; 62:119–124. [PubMed: 9048710]
- 76. Seidler A, Hellenbrand W, Robra BP, Vieregge P, Nischan P, Joerg J, et al. Possible environmental, occupational, and other etiologic factors for parkinson's disease: A case-control study in germany. Neurology. 1996; 46:1275–1284. [PubMed: 8628466]
- 77. Shitaka Y, Tran HT, Bennett RE, Sanchez L, Levy MA, Dikranian K, et al. Repetitive closed-skull traumatic brain injury in mice causes persistent multifocal axonal injury and microglial reactivity. J Neuropathol Exp Neurol. 2011; 70:551–567. [PubMed: 21666502]
- 78. Shively S, Scher AI, Perl DP, Diaz-Arrastia R. Dementia resulting from traumatic brain injury: What is the pathology? Arch Neurol. 2012; 69:1245–1251. [PubMed: 22776913]

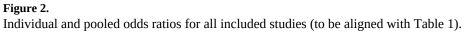
- 79. Silver JM, Kramer R, Greenwald S, Weissman M. The association between head injuries and psychiatric disorders: Findings from the new haven NIMH epidemiologic catchment area study. Brain Inj. 2001; 15:935–945. [PubMed: 11689092]
- Smargiassi A, Mutti A, De Rosa A, De Palma G, Negrotti A, Calzetti S. A case-control study of occupational and environmental risk factors for parkinson's disease in the emilia-romagna region of italy. Neurotoxicology. 1998; 19:709–712. [PubMed: 9745932]
- Smith K, Flicker L, Dwyer A, Atkinson D, Almeida OP, Lautenschlager NT, et al. Factors associated with dementia in aboriginal australians. Aust N Z J Psychiatry. 2010; 44:888–893. [PubMed: 20932202]
- Stern RA, Daneshvar DH, Baugh CM, Seichepine DR, Montenigro PH, Riley DO, et al. Clinical presentation of chronic traumatic encephalopathy. Neurology. 2013; 81:1122–1129. [PubMed: 23966253]
- Sterne JA, Egger M. Funnel plots for detecting bias in meta-analysis: Guidelines on choice of axis. J Clin Epidemiol. 2001; 54:1046–1055. [PubMed: 11576817]
- Stroup DF, Berlin JA, Morton SC, Olkin I, Williamson GD, Rennie D, et al. Meta-analysis of observational studies in epidemiology: A proposal for reporting meta-analysis of observational studies in epidemiology (MOOSE) group. JAMA. 2000; 283:2008–2012. [PubMed: 10789670]
- Suhanov AV, Pilipenko PI, Korczyn AD, Hofman A, Voevoda MI, Shishkin SV, et al. Risk factors for alzheimer's disease in russia: A case-control study. Eur J Neurol. 2006; 13:990–995. [PubMed: 16930366]
- Sundstrom A, Nilsson LG, Cruts M, Adolfsson R, Van Broeckhoven C, Nyberg L. Increased risk of dementia following mild head injury for carriers but not for non-carriers of the APOE epsilon4 allele. Int Psychogeriatr. 2007; 19:159–165. [PubMed: 16684399]
- Taylor CA, Saint-Hilaire MH, Cupples LA, Thomas CA, Burchard AE, Feldman RG, et al. Environmental, medical, and family history risk factors for parkinson's disease: A new englandbased case control study. Am J Med Genet. 1999; 88:742–749. [PubMed: 10581500]
- Terrell TR, Bostick RM, Abramson R, Xie D, Barfield W, Cantu R, et al. APOE, APOE promoter, and tau genotypes and risk for concussion in college athletes. Clin J Sport Med. 2008; 18:10–17. [PubMed: 18185033]
- Tran HT, LaFerla FM, Holtzman DM, Brody DL. Controlled cortical impact traumatic brain injury in 3xTg-AD mice causes acute intra-axonal amyloid-beta accumulation and independently accelerates the development of tau abnormalities. J Neurosci. 2011; 31:9513–9525. [PubMed: 21715616]
- 90. Tripathi M, Vibha D, Gupta P, Bhatia R, Srivastava MV, Vivekanandhan S, et al. Risk factors of dementia in north india: A case-control study. Aging Ment Health. 2012; 16:228–235. [PubMed: 21714688]
- Tsai CH, Lo SK, See LC, Chen HZ, Chen RS, Weng YH, et al. Environmental risk factors of young onset parkinson's disease: A case-control study. Clin Neurol Neurosurg. 2002; 104:328– 333. [PubMed: 12140099]
- Tsolaki M, Fountoulakis K, Chantzi E, Kazis A. Risk factors for clinically diagnosed alzheimer's disease: A case-control study of a greek population. Int Psychogeriatr. 1997; 9:327–341. [PubMed: 9513031]
- 93. Turner MR, Abisgold J, Yeates DG, Talbot K, Goldacre MJ. Head and other physical trauma requiring hospitalisation is not a significant risk factor in the development of ALS. J Neurol Sci. 2010; 288:45–48. [PubMed: 19878957]
- 94. Tyas SL, Pederson LL, Koval JJ. Is smoking associated with the risk of developing alzheimer's disease? results from three canadian data sets. Ann Epidemiol. 2000; 10:409–416. [PubMed: 11018343]
- 95. Unverzagt FW, Ogunniyi A, Taler V, Gao S, Lane KA, Baiyewu O, et al. Incidence and risk factors for cognitive impairment no dementia and mild cognitive impairment in african americans. Alzheimer Dis Assoc Disord. 2011; 25:4–10. [PubMed: 20921881]
- 96. van Reekum R, Cohen T, Wong J. Can traumatic brain injury cause psychiatric disorders? J Neuropsychiatry Clin Neurosci. 2000; 12:316–327. [PubMed: 10956565]

- Vanderploeg RD, Belanger HG, Curtiss G. Mild traumatic brain injury and posttraumatic stress disorder and their associations with health symptoms. Arch Phys Med Rehabil. 2009; 90:1084– 1093. [PubMed: 19577020]
- Werneck AL, Alvarenga H. Genetics, drugs and environmental factors in parkinson's diseaseA case-control study. Arq Neuropsiquiatr. 1999; 57:347–355. [PubMed: 10450337]
- Zhang QG, Laird MD, Han D, Nguyen K, Scott E, Dong Y, et al. Critical role of NADPH oxidase in neuronal oxidative damage and microglia activation following traumatic brain injury. PLoS One. 2012; 7:e34504. [PubMed: 22485176]









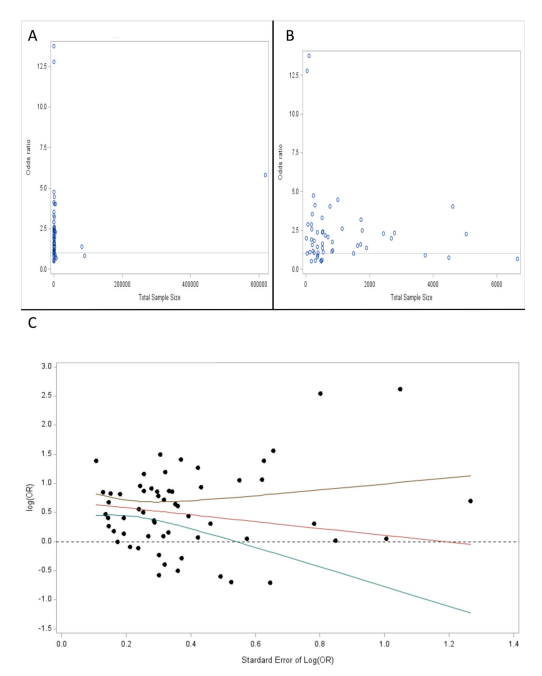


Figure 3.

Publication bias analysis. (A) Funnel plot of OR versus total sample size. (B) Funnel plot of OR versus total sample size after excluding the three studies with largest sample sizes (Rugbjerg 2008, Nielson 2001, and Turner 2010). (C) Plot of the logarithm of the OR after excluding the three largest sample size studies compared to the standard error of the logarithm of the OR showing a regression line and 95% confidence interval with slope that is not statistically significantly different from 0.

Table 1

Individual and pooled odds ratios for all included studies

Study	Cases (# with TBI/ # without TBI)	Controls (# with TBI/ # without TBI)	OR	95% CI
Neurological diagnosis		;		
Dementia				
Salib 1997 ^{<i>b</i>,<i>c</i>,<i>d</i>,<i>e</i>}	96/266	23/153	2.40	1.46-3.95
Schofield 1997 ^{b,e,g}	6/41	21/198	1.38	0.52-3.61
Mehta 1999 ^{<i>d</i>,<i>e</i>,<i>g</i>}	11/118	788/5728	0.68	0.36-1.26
Plassman 2000 ^{b,e,f,g}	28/26	520/1202	2.49	1.45-4.29
Sundstrom 2007 ^{b,e,h}	25/156	46/316	1.10	0.65-1.86
Smith 2010 ^e	31/14	154/164	2.36	1.21-4.60
Tripathi 2012 ^e	22/128	35/115	0.56	0.31-1.02
Pooled OR			1.36	0.84-2.19
Alzheimer's disease				
Forster 1995 ^b	25/84	22/87	1.18	0.62-2.25
Rasmusson 1995 ^{b,d,e}	20/48	1/33	13.75	1.76-107.53
Salib 1997 ^{<i>a</i>}	53/145	23/153	2.43	1.42-4.17
Schofield 1997 ^a	4/34	23/205	1.05	0.34-3.22
Tsolaki 1997	14/47	15/54	1.07	0.47-2.45
O'Meara 1997 ^{<i>b</i>,<i>e</i>}	32/317	16/326	2.06	1.11-3.82
Boston 1999	30/192	23/117	0.79	0.44-1.43
Mehta 1999 ^a	6/85	788/5728	0.51	0.22-1.18
Guo 2000 ^e	394/1782	127/2313	4.03	3.27-4.96
Plassman 2000 ^a	17/18	520/1202	2.18	1.12-4.27
Tyas 2000	203/821	93/605	1.61	1.23-2.10
Lindsay 2002 ^g	28/151	603/2963	0.91	0.60-1.38
Bachman 2003 ^e	397/1538	84/760	2.34	1.82-3.00
Guskiewicz 2005 ^e	15/7	1148/732	1.37	0.56-3.37
Ogunniyi Nigeria 2006 ^g	2/60	11/450	1.36	0.30-6.30
Ogunniyi USA 2006 ^g	5/84	37/344	0.55	0.21-1.45
Rippon 2006 ^e	72/78	648/700	1.00	0.71-1.40
Suhanov 2006 ^{d,e}	46/214	30/230	1.65	1.00-2.71
Fischer 2008 ^g	4/86	37/352	0.44	0.15-1.27
Pooled OR			1.40	1.02-1.90
Parkinson's disease				
Martyn 1995 ^e	11/156	35/301	0.61	0.30-1.23

De Michele 1996 ^{d,e}		TBI)		
De Michele 1996	13/103	3/113	4.75	1.32-17.16
Seidler 1996 ^e			1.40	0.85-2.30
McCann 1998 ^{d,e}			1.10	0.64–1.90
Smargiassi 1998 ^{d,e}	13/73	5/81	2.88	0.98-8.49
Kuopio 1999 ^{d,e,h}	39/84	84/162	0.90	0.56-1.42
Taylor 1999 ^b ,e	35/105	11/136	4.12	2.00-8.50
Werneck 1999	17/75	14/96	1.55	0.72-3.35
Tsai 2002 ^{<i>b</i>,<i>e</i>}	11/19	5/25	2.89	0.86–9.75
Baldereschi 2003 ^{d,e,g}	8/105	403/3980	0.75	0.36-1.56
Bower 2003 ^{b,c,e,f}	2/183	2/193	1.05	0.15-7.57
Goldman $2006^{b,c,e,h}$	20/73	9/84	2.56	1.10-5.96
Dick 2007 ^{<i>d</i>,<i>e</i>}			1.30	1.09–1.55
	409/13194	1513/66792	1.37	1.22–1.53
Rugbjerg 2008 ^{b,e,f} Sanyal 2010	27/148	25/325	2.37	1.33-4.23
Pooled OR	27/148	23/323	1.45	1.33-4.23 1.18-1.78
Amyotrophic lateral scleros	sis		1.40	1110 1110
Chen 2007 ^{<i>b</i>,<i>e</i>,<i>h</i>}	24/85	42/213	1.43	0.82-2.51
Binazzi 2009 ^b	16/61	23/162	1.85	0.91-3.73
Schmidt 2010 ^{b,e,h}	84/157	185/412	1.19	0.87-1.64
Turner $2010^{b,e,f,g}$	41/34	106552/511831	5.79	3.68-9.13
Pooled OR			2.07	0.94-4.56
Frontotemporal dementia			2.07	0194 4120
Rosso 2003 b,e	19/61	10/114	3.55	1.55-8.11
Vascular dementia				
Boston 1999	3/31	23/117	0.49	0.14–1.75
Mild cognitive impairment				
Guskiewicz 2005 ^e	19/3	450/286	4.03	1.18–13.73
Unverzagt 2011g			2.40	1.34-4.30
Pooled OR			2.69	1.51-4.77
Pooled OR Neurologic			1.55	1.31–1.83
sychiatric diagnosis				
Depression				
Malaspina 2001	107/661	22/355	2.61	1.62-4.21
Polusny 2001 ^{b,c,e,g}			1.47	1.10–1.97
Silver 2001 ^e	40/243	321/4430	2.27	1.60-3.23
Holsinger 2002 ^{b,c,e,f,g}	96/160	387/974	1.51	1.14-2.00

Study	Cases (# with TBI/ # without TBI)	Controls (# with TBI/ # without TBI)	OR	95% CI
Guskiewicz 2007 ^e	206/63	1272/893	2.30	1.71-3.08
Gao 2009	38/497	28/1174	3.21	1.95-5.28
Mollica 2009 ^{<i>d</i>,<i>e</i>}	10/3	6/23	12.78	2.65-61.56
Rajkumar 2009 ^{d,e}	19/108	33/840	4.48	2.46-8.15
Vanderploeg 2009 ^{b,e,g}	36/43	242/505	1.75	1.09-2.79
Bryant 2010 ^{b,c,e,f,g}	56/265	77/419	1.15	0.79–1.68
Pooled OR			2.14	1.65-2.77
Psychotic disorder				
Malaspina 2001	22/107	22/355	3.32	1.77-6.23
Nielsen 2001 ^{b,e,f}	278/7854	3394/78710	0.82	0.72-0.93
Silver 2001 ^a	12/89	349/4584	1.77	0.96-3.27
AbdelMalik 2003 ^b	23/44	22/80	1.90	0.95-3.79
Fann 2004 <i>a</i>			1.10	0.39–3.10
Pooled OR			1.57	0.83-2.97
Bipolar disorder				
DelBello 1999 ^{b,c,d,e,g}	4/17	3/13	1.02	0.19–5.37
Malaspina 2001	28/207	22/355	2.18	1.22-3.91
Silver 2001 ^a	6/51	355/4622	1.53	0.65-3.59
Pooled OR			1.85	1.17-2.94
Mixed affective disorder				
Fann 2004 <i>b</i> , <i>f</i> , <i>g</i>			1.50	0.98–2.30
Nelson 2007 ^b	76/248	318/2045	1.97	1.49–2.62
Pooled OR			1.84	1.44-2.36
Pooled OR Psychiatric			2.00	1.50-2.66
Overall Pooled OR			1.67	1.44-1.93

 a Studies that were not included in the overall analysis or pooled neurologic/psychiatric analyses because diagnostic groups within the study were not mutually exclusive. Studies that were included in subgroup analyses:

 $^{b}_{}$ clearest interval between TBI and symptom onset,

^c meeting mild TBI criteria for loss of consciousness,

d requiring loss of consciousness,

^e requiring at least one mild TBI feature,

 $f_{\text{TBI diagnoses not based on self-report,}}$

 g cohort studies, and

h analysis of risk of repeated TBI

Author Manuscript

Author Manuscript

Design and TBI features reported for each included study

Study Neurological diagnosis					
Neurological diagnosis	Study design	Age at head injury, mean	Interval between injury and diagnosis, mean	Required TBI characteristics	Additional TBI information
Dementia					
Salib (1997)	Case-control		7.3 years	None given	Grouped by with or without LOC
Schofield (1997)	Cohort			LOC or PTA	
Mehta (1999)	Cohort		Grouped	LOC	Grouped by LOC< or > 15 minutes
Plassman (2000)	Cohort			MC and LOC or PTA or nondisplaced skull fracture	Excluded if penetrated dura or resulted in significant sequelae 3 months after TBI, severity ranked with mild group having LOC or PTA<30 minutes and no skull fracture
Sundstrom (2007)	Case-control		5 years	MC	
Smith (2010)	Cross-sectional			None given	
Tripathi (2012)	Case-control			LOC or PTA or a symptom of PCS	
Alzheimer's disease					
Forster (1995)	Case-control	Grouped (in adulthood or childhood)		None given	
Rasmusson (1995)	Case-control	27.2 in sporadic Alzheimer's group, 45.2 in familial Alzheimer's group	>5 years (mean 33.4 years in sporadic Alzheimer's group, 18.67 in familial Alzheimer's group)	None given	Excluded if head injury resulted in "immediate, permanent cognitive or functional impairment," head injury with LOC reported separately. Distinction made between mild and severe but not defined.
Salib (1997)	Case-control		7.9 years	None given	Grouped by with or without LOC
Schofield (1997)	Cohort		14.5 years	LOC or PTA	
Tsolaki (1997)	Case-control			None given	
O'Meara (1997)	Case-control	46 (range 10–85)	34 years (range 1- 72)	MC or LOC	

J Neurosurg. Author manuscript; available in PMC 2017 February 01.

Perry et al.

		injury, mean	interval between injury and diagnosis, mean	Kequired TB1 characteristics	Additional 1 B1 miormation
Boston (1999)	Case-control			None given	
Mehta (1999)	Cohort		Grouped	LOC	Grouped by LOC< or > 15 minutes
Guo (2000)	Case-control			MC or LOC	
Plassman (2000)	Cohort			MC and LOC or PTA or nondisplaced skull fracture	Excluded if penetrated dura or resulted in significant sequelae 3 months after TBI, severity ranked with mild group having LOC or PTA<30 minutes and no skull fracture
Tyas (2000)	Cross-sectional			None given	
Lindsay (2002)	Cohort			None given	Both with and without LOC
Bachman (2003)	Case-control			MC	
Guskiewicz (2005)	Cross-sectional			AMS and one symptom of PCS	
Ogunniyi Nigeria (2006)	Cohort			None given	
Ogunniyi USA (2006)	Cohort			None given	
Rippon (2006)	Cross-sectional			LOC or PTA	
Suhanov (2006)	Case-control			LOC	
Fischer (2008)	Cohort			None given	
Parkinson's disease					
Martyn (1995)	Case-control			LOC or MC	
De Michele (1996)	Case-control			LOC	
Seidler (1996)	Case-control			PTA or PCS	
McCann (1998)	Case-control			LOC	
Smargiassi (1998)	Case-control			LOC	
Kuopio (1999)	Case-control			None given	Records number with and without LOC and duration of LOC < or > 5 minutes
Taylor (1999)	Case-control	16.3	36.5 years	LOC or AMS or ND or PCS	
Werneck (1999)	Case-control			None given	
Tsai (2002)	Case-control	18.5	17.2 years	LOC or PTA or PCS or ND	
Baldereschi (2003)	Cohort			LOC	

Bover (2003)Case-controlS3 years (range 3- ystars for TBL) ystars for TBL ystars for TBL ystars for TBL S3, median 23 ystars for TBL S3, median 23 ystars for TBL S4 years for TBL ystars for TBL S4 years for the start S4 years for the startS4 years for the start S4 years for the start S4 years for the start S4 years for the startS4 years for the start S4 years for the start S4 years for the start S4 years for the startS4 years for the start S4 years for the start S4 years for the startS4 years for the start S4 years	Study	Study design	Age at head injury, mean	Interval between injury and diagnosis, mean	Required TBI characteristics	Additional TBI information
Case-control 25.7 36.9 years (range analysis reported and reported analysis reported and reported analysis reported and reported analysis reported an	Bower (2003)	Case-control		>3 years (range 3– 55, median 29 years for TBI of all severities in study)	PTA	Excluded from this group if LOC>1 minute, PTA>30 minutes, or imaging abnormal. Mild TBI with LOC, moderate, and severe TBI analyzed separately
Case-control Case-control Grouped,>l year Case-control Case-control Grouped<>l years Case-control Grouped Grouped Cohort Case-control Grouped Control Control Control Cohort Control Case-control Cohort Control Control Case-control Case-control Case-control Case-control Case-control Case-control Case-control Case-control Case-control Case-control Case-control Case-contro Case-cont	Goldman (2006)	Case-control	25.7	36.9 years (range 2–70), separate analysis reported of only those >10 years	LOC or PTA	
Case-control Grouped, >1 year Case-control Grouped Cohort Consectional Cohort Cose-sectional Cohort Cohort Case-control Case-control	Dick (2007)	Case-control			LOC	
l Case-control Grouped Grouped Case-control Grouped Grouped Case-control Grouped (2-80+ years) Cohort Case-control	Rugbjerg (2008)	Case-control		Grouped, >1 year data used	MC	Excluded if imaging abnormal
Case-control Grouped Case-control Grouped Case-control Grouped Case-control Grouped Case-control Grouped Cohort Grouped case-control Grouped Cohort Grouped Cohort Cose-control Case-control Grouped Cohort Case-control Case-control Grouped Case-control Grouped Case-control Case-control Case-control Case-control Case-control Case-control Case-control Case-control Case-control Case-control	Sanyal (2010)	Case-control			None given	
Case-controlGroupedGroupedCase-controlGroupedGroupedCase-controlGroupedGroupedCase-controlGroupedGroupedCohortCase-controlSears)Case-controlConotCase-control <tr< td=""><td>Amyotrophic lateral sclerosis</td><td></td><td></td><td></td><td></td><td></td></tr<>	Amyotrophic lateral sclerosis					
Case-controlGroupedGroupedCase-controlGrouped (2-80+ years)cohortCohortcohortCohortcase-controlCohortCase-controlCohortCross-sectionalCohortCase-controlCohortCohortCohortCohortCohortCase-controlCohortCohortCohortCohortCohortCase-controlCohortCohortCohortCohortCohortCohortCohortCase-controlCohortCohortCohortCohortCohortCohortCohortCase-controlCohortCohor	Chen (2007)	Case-control	Grouped	Grouped	MC	
Case-control Grouped Grouped (2-80+ years) cohort Cohort Case-control Case-control Case-control Case-control Conort Control Case-control Conort Control Case-control	Binazzi (2009)	Case-control	Grouped	Grouped	None given	
Cohort Case-control Case-control Cross-sectional Cohort Cohort Case-control	Schmidt (2010)	Case-control	Grouped	Grouped (2-80+ years)	LOC or MC	
nentia Case-control Case-control Cross-sectional Cohort Case-control	Turner (2010)	Cohort			MC	
Case-control Case-control Cross-sectional Cohort Cohort Case-control	Frontotemporal dementia					
Case-control Cross-sectional Cohort Case-control	Rosso (2003)	Case-control			PCS or LOC or PTA	
Case-control Cross-sectional Cohort Case-control	Vascular dementia					
Cross-sectional Cohort Case-control	Boston (1999)	Case-control			None given	
Cross-sectional Cohort Case-control	Mild cognitive impairment					
Cohort Case-control	Guskiewicz (2005)	Cross-sectional			AMS and one symptom of PCS	
Case-control	Unverzagt (2011)	Cohort			None given	
(2001) Case-control	Psychiatric diagnosis					
Case-control	Depression					
	Malaspina (2001)	Case-control			None given	Severity grouped by LOC duration with "severe" TBI having LOC >15 minutes

Study	Study design	Age at head injury, mean	Interval between injury and diagnosis, mean	Required TBI characteristics	Additional TBI information
Polusny (2001)	Cohort		>1 year (1–2.33 years)	AMS or LOC	LOC >20 minutes excluded
Silver (2001)	Cross-sectional			LOC or AMS	
Holsinger (2002)	Cohort	20.9 (includes some not in analysis)		MC + LOC or PTA or nondisplaced skull fracture	Excluded if penetrated dura or resulted in significant sequelae 3 months after TBI
Guskiewicz (2007)	Cross-sectional			AMS and one symptom of PCS	
Gao (2009)	Cross-sectional			None given	
Mollica (2009)	Cross-sectional			LOC, PTA, and ND	
Rajkumar (2009)	Cross-sectional			LOC	
Vanderploeg (2009)	Cohort		16 years	LOC or PTA or AMS	Excluded if admitted to hospital
Bryant (2010)	Cohort	37.8	1 year	GCS 13	Excluded if focal deficit, imaging abnormal, or LOC>30 minutes
Psychotic disorder					
Malaspina (2001)	Case-control			None given	Severity grouped by duration LOC with "severe" TBI having LOC >15 minutes
Nielson (2001)	Case-control		Grouped (>1 year)	MC	ICD9 code for concussion included, excluded if skull fracture or intracranial hemorrhage
Silver (2001)	Cross-sectional			LOC or AMS	
AbdelMalik (2003)	Case-control	<17	Median 12 years		Closed head injuries without intracranial hemorrhage or other immediate sequelae
Fann (2004)	Cohort		3 years		By ICD9 codes - Excluded if imaging abnormal or LOC>1 hour
Bipolar disorder					
DelBello (1999)	Cross-sectional	10.7	6.3 years	LOC	
Malaspina (2001)	Case-control			None given	Severity grouped by duration LOC with "severe" TBI having LOC >15 minutes
Silver (2001)	Cross-sectional			LOC or AMS	i.

Author Manuscript

Study	Study design	Study design Age at head injury, mean	Interval between F injury and c diagnosis, mean	Required TBI characteristics	Additional TBI information
Mixed affective disorder					
Fann (2004)	Cohort		3 years		By ICD9 codes - Excluded if imaging abnormal or LOC>1 hour
Nelson (2007)	Cross-sectional		>1 year	None given	

"Grouped" refers to data presented in the paper by stratification or division of subjects into groups without an available mean.

AMS – alteration in mental status, GCS – Glasgow Coma Scale, LOC – loss of consciousness, MC – injury for which medical care was received, ND – Neurological deficit, PCS – post-concussion syndrome (e.g., headache, dizziness, nausea, photo- or phonophobia, fatigue, sleep difficulty, blurred vision), PTA – post-traumatic amnesia, TBI –traumatic brain injury

Table 3

Results of subgroup analyses

Analysis	Odds ratio	95 % confidence interval	Statistical significance
Risk of TBI when including only studies with clearest interval on:			
All neurologic and psychiatric outcomes	1.75	1.43–2.14	<i>p</i> <.001
All neurologic outcomes	2.05	1.55-2.71	<i>p</i> <.001
All psychiatric outcomes	1.38	0.95-2.00	<i>p</i> =.09
Risk of TBI when including only studies meeting mild TBI requirements for maximum duration of loss of consciousness	1.54	1.18-2.01	<i>p</i> =.001
Risk of TBI when including only studies requiring associated loss of consciousness on:			
All neurologic and psychiatric outcomes	1.69	1.18-2.44	<i>p</i> <.01
All neurologic outcomes	1.33	1.00-1.75	<i>p</i> <.05
All psychiatric outcomes	4.09	1.36-12.32	<i>p</i> =.01
Risk of TBI when including only studies requiring a mild TBI feature on:			
All neurololgic and psychiatric outcomes	1.70	1.42-2.05	<i>p</i> <.0001
All neurologic outcomes	1.67	1.36-2.07	<i>p</i> <.0001
All psychiatric outcomes	1.81	1.23-2.66	<i>p</i> <.01
Risk of TBI when eliminating studies with TBI diagnosis based on self-report on:			
All neurologic and psychiatric outcomes	1.62	1.14–2.31	<i>p</i> <.01
All neurologic outcomes	2.38	1.01-5.62	<i>p</i> <.05
All psychiatric outcomes	1.18	0.81-1.71	<i>p</i> =.39
Risk of TBI when including only cohort studies on:			
All neurologic and psychiatric outcomes	1.38	1.02-1.87	<i>p</i> <.05
All neurologic outcomes	1.27	0.72-2.25	<i>p</i> =.41
All psychiatric outcomes	1.45	1.23–1.71	<i>p</i> <.0001
Risk of multiple TBIs vs one TBI on any outcome diagnosis	1.10	0.72–1.70	<i>p</i> =.65