An integrated perspective linking physiological and psychological consequences of mild traumatic brain injury

Harm Jan van der Horn1 · Manon L. Out1 · Myrthe E. de Koning2 · Andrew R. Mayer3 · Jacoba M. Spikman4 · Iris E. Sommer5 · Joukje van der Naalt1

Received: 18 March 2019 / Revised: 22 April 2019 / Accepted: 23 April 2019 / Published online: 27 April 2019
© The Author(s) 2019

Abstract
Despite the often seemingly innocuous nature of a mild traumatic brain injury (mTBI), its consequences can be devastating, comprising debilitating symptoms that interfere with daily functioning. Currently, it is still difficult to pinpoint the exact cause of adverse outcome after mTBI. In fact, extensive research suggests that the underlying etiology is multifactorial. In the acute and early sub-acute stages, the pathophysiology of mTBI is likely to be dominated by complex physiological alterations including cellular injury, inflammation, and the acute stress response, which could lead to neural network dysfunction. In this stage, patients often report symptoms such as fatigue, headache, unstable mood and poor concentration. When time passes, psychological processes, such as coping styles, personality and emotion regulation, become increasingly influential. Disadvantageous, maladaptive, psychological mechanisms likely result in chronic stress which facilitates the development of long-lasting symptoms, possibly via persistent neural network dysfunction. So far, a systemic understanding of the coupling between these physiological and psychological factors that in concert define outcome after mTBI is lacking. The purpose of this narrative review article is to address how psychophysiological interactions may lead to poor outcome after mTBI. In addition, a framework is presented that may serve as a template for future studies on this subject.

Keywords Mild traumatic brain injury · Biomarkers · Cytokines · Cortisol · Psychology

Introduction
The vast majority (85–90%) of patients with traumatic brain injury (TBI) sustain a mild traumatic brain injury (mTBI). With an estimated annual worldwide incidence of approximately 600/100,000 persons (42,000,000 persons worldwide), mTBI is the most common neurologic disorder [1, 2]. For a considerable percentage of patients (≈ 25%) it takes months or even years to recover due to persistent cognitive and emotional symptoms that interfere with resumption of work, social activities and studies [3–5]. Hence, this patient group imposes a tremendous burden on health services and economy. Although acute symptoms are thought to result from the brain injury itself, a comprehensive physiological substrate for the persistence of symptoms has not been empirically established. In fact, for the majority of patients conventional computed tomography (CT) or magnetic resonance imaging (MRI) do not show traumatic abnormalities, and if abnormalities are identified, they tend to correlate poorly with persisting symptom severity [6–8]. The application of advanced imaging modalities, such as functional MRI (fMRI) and diffusion-weighted imaging (DWI), has provided evidence of altered brain network connectivity after mTBI, although a clear-cut mechanism underlying continued symptoms has not been found [9, 10]. There is also increasing evidence that certain protein biomarkers of
cellular injury in the acute phase post-injury are informative of injury severity and clinical outcome after mTBI [11, 12]. However, it is unclear if these markers have a role in the pathophysiology of persistent symptoms, and how this process may be influenced by other acute physiological sequelae such as inflammation or the acute stress response. Furthermore, there still exists a lively debate as to whether the influence of psychological factors may outweigh that of the physiological consequences of the injury itself in the causation of persistent symptoms and poor outcome after mTBI. Psychological factors, such as the presence of anxiety and depression, and the employment of certain coping styles, are known to exert a strong influence on recovery after mTBI [5, 13–15]. Not surprisingly, stressful events such as sustaining a traumatic brain injury, can result in an increased demand on coping skills, especially in the current busy and often stressful modern environment. One of the most important elements of coping is the ability to regulate negative emotions and stress, which is closely related to our personality [16, 17]. Inadequate emotion regulation could lead to emotional distress, such as anxiety and depression, which may enhance the persistence of post-traumatic symptoms.

The pathophysiology of mTBI encompasses various (mostly acute) physiological and (mostly sub-acute/chronic) psychological processes, which are intricately linked. Disentangling these relationships forms one of the major challenges in mTBI research. For example, it is still poorly understood whether or not physiological disturbances in the acute phase post-injury, such as cellular injury, inflammation and the acute stress response, are related to psychological problems at a later stage after mTBI, and whether there is an association with perturbations in neural networks that are necessary for emotion regulation and adequate coping skills. In the current narrative review, we aim to provide an overview of these physiological and psychological factors, and theoretically explore the interaction between these factors in the etiology of persistent symptoms and poor outcome after mTBI.

### Cellular injury

Traumatic injury to neuronal, axonal, glial, and vascular structures causes the release of brain-specific proteins [11]. Promising results have been published suggesting that serum levels of brain-specific proteins, such as S100 calcium-binding protein B (S100B; primarily found in astroglial cells, but also in melanocytes), glial fibrillary acidic protein (GFAP; present in the cytoskeleton of several cells in the central nervous system, for example astrocytes), tau (protein that stabilizes microtubuli that make up the cytoskeleton of axons), neurofilament light (NF-L; protein that is also part of the cytoskeleton of neurons), and ubiquitin C-terminal hydrolase-L1 (UCH-L1; an enzyme which plays a role in the repair of neurons and axons via regulation of protein degradation), are predictive of the presence of lesions after mTBI, and poor outcome after mTBI [11, 18–23]. In Scandinavia, S100B measurement in the acute phase post-injury has already been added to the Neurotrauma guidelines as a biomarker to reduce unnecessary CT-scans and associated costs [24]. In the United States, UCH-L1 and GFAP have been approved for the same purpose [23]. In sports-related mTBI, elevated tau-protein is related to recovery time and return to play [18]. It is thought that tau measurements might be used as a tool to detect residual neural damage, even in the absence of acute symptoms, which may protect a player from premature return to play, thereby reducing the risk of sustaining additional injury. Furthermore, hyperphosphorylated tau protein is a hallmark of the pathophysiology of chronic traumatic encephalopathy (CTE), which is a neurodegenerative disorder that can occur even in patients with repetitive mild TBI [25].

Despite these findings, we still know little about the exact role of these proteins in the development of persistent symptoms. The kinetics of these biomarkers vary, with S100B and GFAP reaching a peak serum concentration in the first 24 h after injury (‘acute biomarkers’), tau staying elevated for days to weeks (‘subacute biomarkers’), and NF-L continue rising for weeks to months after injury (‘chronic biomarkers’) [26]. Therefore, it would be interesting to investigate whether ‘acute’ biomarkers predict, and if ‘chronic’ biomarkers accompany long-lasting symptoms.

### Inflammation

Under normal circumstances, inflammation is closely regulated, and helps to repair damaged tissue and to fight infections [27, 28]. However, excessive and prolonged inflammation, for example after TBI, has the opposite effect [28]. It has been shown that neuroinflammation occurs within seconds to minutes after a TBI, and involves a complex cascade of microglia activation, pro- and anti-inflammatory cytokine release, and up- and downregulation of neurotrophic factors [28, 29]. Neuropathological research has demonstrated signs of neuroinflammation up to 18 years post-injury in patients with moderate/severe TBI, and these findings were related to white-matter degeneration [30]. Furthermore, a positron emission tomography (PET) study on American football players has shown microglia activation many years after the last mTBI [31], and chronic microglia activation is associated with the development of CTE [32]. Studies using rodent models have yielded data on the relationship between inflammation and behavioural deficits after mild TBI [29, 33–37]. However, relatively little is known about the role of inflammation in the pathophysiology of sequelae.
following mild TBI in humans, as compared to moderate-severe TBI [29]. A retrospective study on a human mTBI sample has shown an association between persistent post-traumatic symptoms and elevated C-reactive protein within the first 72 h after injury [38]. Another recent retrospective study demonstrated a relation between interleukin (IL)-1β and IL-6 gene polymorphisms and the number and duration of post-traumatic symptoms after sports-related concussion [39]. In addition, reviews have been published that posit the possible influence of earlier (pre-injury) pro-inflammatory states, extracranial injury, infections and immunosenescence (i.e., deterioration of the immune system due to aging) on the occurrence of (neuro)inflammation in the pathophysiology of mTBI [40, 41]. For example, similarities exist between post-traumatic symptomatology and cognitive or behavioral changes that arise from other inflammatory conditions, such as sepsis [40]. Interestingly, a recent study has shown that using an univariate multivariate approach applied to a multi-analyte proteomic panel, including several inflammatory markers, it was possible to fairly accurately predict positive CT-findings and outcome in a study sample that consisted predominantly of mild TBI patients, although 42% of these patients had lesions on CT [42].

It is important for future studies on the role of acute and chronic inflammation in the development of persistent symptoms and poor outcome, to compare patients with mild TBI to patients with peripheral inflammatory conditions, such as orthopedic injury.

**Stress**

During an acute stress response, for example elicited by a traumatic brain injury, brain structures such as the hypothalamus, amygdala, and insula become activated, which engage the sympathetic autonomic nervous system and the hypothalamic–pituitary–adrenal (HPA) axis, leading to increased heart action and the release of catecholamines and cortisol. The release of cortisol is needed to initiate adaptive metabolic and mental processes to deal with the acute stressor [43]. However, an exaggerated and protracted stress response, which may be due to poor emotion regulation skills in case of stressful situations, is harmful to an individual. Chronically changed cortisol levels are associated with various physical and mental diseases [44]. Furthermore, excessive or prolonged stress can induce inflammation, via microglia activation, and the release of pro-inflammatory cytokines such as interleukin (IL)-1β and 6, and tumor necrosis factor (TNF)-α [27, 45]. These cytokines have also been found to be related to emotion regulation deficits in healthy young adults [46]. An important causative factor in the pro-inflammatory effect of (especially chronic) stress is glucocorticoid resistance, which is the decrease in sensitivity of immune cells to the effects of glucocorticoids [47].

There have been some data published on cortisol in patients with mTBI, and patients with mild to severe TBI [48–52]. These studies, although limited by sample size and lack of control groups, demonstrated that associations are present between cortisol levels, injury severity, and cognitive deficits at various stages post-injury. On the one hand, alterations in cortisol levels after TBI can be caused by HPA-axis dysfunction due to the injury itself, which is common in patients with TBI, even following mild TBI [53, 54]. On the other hand, alterations in cortisol levels (i.e., increased levels) after mTBI can be viewed as a physiological response to the psychological effects of sustaining a TBI, which is a stressful event. It is still unknown if acute changes in cortisol levels are involved in the causation and/or persistence of symptoms, and if the effects of stress are mediated via inflammation. In contrast to acute stress, it is thought that psychological mechanisms are responsible for chronic stress and persistent symptoms after mTBI, although to our knowledge this has not yet been corroborated by results from longitudinal cortisol measurements. Similar to cytokine analyses, acute changes in cortisol can be reliably measured in both serum and saliva. Interestingly, chronic stress can (also) be measured in hair samples. In fact, differences in long-term hair cortisol levels have been found between various mental illness such as major depressive disorder, anxiety disorders and post-traumatic stress disorder, illnesses which are also quite common after mild TBI [44, 55]. Since hair grows approximately 1 cm per month, hair analysis is also informative of the cortisol levels before and after a stressful event, which makes it possible to investigate stress before and after mTBI, and more specifically, to investigate whether pre-injury stress levels are associated with chronic stress after injury.

**Neural networks and emotion regulation**

Adequate emotion regulation skills are a prerequisite to deal with stressful conditions, such as a mTBI. Although many fMRI-studies have been conducted in non-TBI study samples to investigate neural networks during tasks that challenge certain emotion regulation skills, or during tasks that contain stressful conditions, not much is known about functioning of neural networks during emotion regulation after mTBI [10, 17, 56]. There are two major types of emotion regulation: top–down (explicit) and bottom–up (implicit) emotion regulation [57]. One of the most important top–down emotion regulation strategies is cognitive reappraisal, which refers to the process of rethinking negative thoughts and changing them into more neutral or perhaps even positive thoughts [58]. Top–down emotion regulation is mainly conducted...
using the executive network, a network that is also important for cognitive performance [17, 57, 59]. Neural networks that play a pivotal role in emotion regulation are depicted in Fig. 1; notably, the prefrontal cortex occupies a central position in these networks. The default mode network is generally considered a task-negative network, as areas of this network (e.g., PCC, precuneus and medial prefrontal cortex) are consistently deactivated during externally directed mental tasks [60]. Accumulating evidence suggests that these areas are also important for top–down emotion regulation, especially when emotion regulation is stimulated via social support from family-members, friends, or psychotherapists [17, 61]. However, over-activity in this network is often associated with psychopathology such as major depressive disorder [62]. Recently, a study on healthy adults suggested that there is an association between pro-inflammatory cytokines and functional connectivity of the default mode network, which highlights the interaction between stress, inflammation and network dysfunction in psychopathology [63].

In contrast to top–down emotion regulation, bottom–up regulation refers to the implicit processing of salient information that generates emotions, for example a stressful event or stimulus, without engagement of higher order cognitive mechanisms [57, 64]. An example of a bottom–up mental process is when attention is focused to the present internal sensations (e.g., pain, emotions) without cognitively analyzing these sensations. An important network for bottom–up emotion regulation is the salience network, which consists of the insulae and anterior cingulate cortex. The mechanism by which the salience network contributes to effective emotion regulation during stressful conditions, is through generation of emotional awareness and subsequent modulation of executive and default mode network activity [57, 65–67]. The salience network forms an intricate and dynamic link between autonomic bodily arousal responses that accompany emotional and stressful states, and cognitive emotion regulation [68]. Furthermore, HPA-axis reactivity is positively linked to connectivity of the salience network, which highlights the role of this network in modulation of stress responses [67].

Mindfulness, which has also shown to be beneficial in mTBI, engages bottom–up as well as top–down emotion regulation mechanisms, depending on the duration of practice [64, 69]. Mindfulness is a form of meditation, which is derived from Buddhism and involves focusing one’s attention to the present moment, maintaining awareness and control of present thoughts, feelings and sensations, and observing them without judgment [70]. Interestingly, fMRI studies have shown that mindfulness increases activity of areas within the salience network [71], reduces default mode network activity [72], and alters the functional collaboration between the salience, executive, and default mode networks [65].

Resting state functional MRI (fMRI) research from several research groups, including our own, has shown the involvement of the aforementioned neural networks in the presence and persistence of post-traumatic symptoms, anxiety and depression after mTBI [73–77]. Global increases in resting-state functional connectivity in symptomatic patients with sports-related mild TBI seem to have a delayed

Fig. 1 Spatial maps representing neural networks that are important for emotion regulation. The executive network is depicted in yellow (key areas: ventro- and dorsolateral prefrontal cortex, supplementary motor area, and posterior parietal cortex), the salience network in red (key areas: insula and anterior cingulate cortex) and the default mode network in blue (key areas: medial prefrontal cortex, posterior cingulate cortex, and precuneus). Maps are derived from fMRI-data of our own department.
onset, peaking at approximately 1 week post-injury [78]. Our research group has found that higher resting-state connectivity within the default mode network in the sub-acute phase post-injury, is a possible biomarker for the development of persistent symptoms [74]. However, little is known about neural networks during the actual process of emotion regulation, which may be difficult to perform in daily life for a subgroup of patients with mTBI. This can be tested using emotion regulation fMRI tasks, which are comprised of conditions that evoke either positive or negative emotions. Participants are instructed to apply certain emotion-regulation techniques such as enhancing or suppressing these emotions during these conditions. To our best knowledge, only two studies have been published that used emotion regulation fMRI paradigms in mTBI. One study demonstrated lower activity in the orbitofrontal cortex and superior parietal lobe during bottom-up emotion induction (i.e., condition in which subjects had to judge whether an angry, fearful or neutral face belonged to a male or female), possibly indicating impaired emotion regulation [59]. These findings were related to higher post-traumatic stress scores. The other study showed higher activity of the insula, peri-central gyri, parietal and temporal cortex in emotion enhancement (i.e., condition in which subjects had to try to amplify their negative feelings towards a negatively valenced picture), which may indicate increased vigilance or bottom-up emotional processing, or poor recruitment of areas for emotion regulation [79]. It is still unclear how the executive, default mode, and salience networks function during emotion regulation after mTBI.

### The heart as a biomarker of emotion regulation and stress

So far, it can be concluded that the complex interplay between neural networks that are important for emotion regulation (i.e., executive, default mode, and salience network), the autonomic nervous system, and the HPA-axis, determines an individual’s capacity to deal with stressful situations [56, 80, 81]. The neurovisceral integration model (NIM) states that resting-state heart rate variability (HRV), defined as the variability of time between heartbeats, is an indirect marker of the function of these neural networks [80, 82, 83]. The heart is under constant inhibitory control of the parasympathetic nervous system (vagal nerve tone), which dominates the sympathetic nervous system, keeping the heart rhythm in balance [81]. Simplistically stated, high resting-state HRV primarily reflects higher parasympathetic activity, whereas low HRV reflects activity of the sympathetic nervous system. With respect to the NIM, high resting state HRV is considered a marker of high vagal tone and thus of adequate function of the neural networks. Emotion regulation difficulties and psychopathological states, such as anxiety, depression and post-traumatic stress disorder, are related to lower HRV [80, 81, 83–86]. In addition, higher resting state HRV is associated with better working memory performance in healthy people [87].

The last few years, several studies have been published that report changes in HRV after traumatic brain injury, both at the severe as well as at the mild end of the TBI spectrum [51, 88–93]. Among other things, it has been shown that lower HRV is related to anxiety and depression after TBI [92]. However, most of these studies were sub-optimally designed, and to our knowledge, no study so far focused on HRV in the acute phase after injury, or on the relationship between HRV and function of neural networks (as measured with fMRI).

### The psychophysiological linkage in mTBI

It is still unknown to what extent the various acute biochemical effects are responsible for persistent symptoms and poor outcome after mTBI, and if there is a correlation with psychological factors. More research is required to understand this complex pathophysiological process.

Previous research has shown that traumatic injury to neural, glial or vascular tissue causes the release of brain-specific proteins, and that measurement of levels of these proteins in the acute phase after injury offers some diagnostic value [18–20, 24]. However, not much is known about the role of these proteins in the development of persistent symptoms after mTBI. Another consequence of cellular injury is neuroinflammation, which has barely been investigated in clinical mTBI samples. Neuroinflammation may also develop secondary to high cortisol levels associated with stress. Inter-individual differences in cortisol release acutely after injury can be either linked to the degree of injury to neural structures that regulate the HPA-axis, or to pre-existent hyper- or hypo-reactivity of the HPA-axis [53, 54]. The latter could be associated with pre-existent psychological factors such as the ability to cope with stress or the experience of stressful situations or psychotrauma in childhood. Thus, the extent of the acute stress response probably depends on both physiological as well as psychological factors, although this has never been thoroughly investigated in patients with mTBI.

The interaction between cellular injury, inflammation and stress, mediated by pre-existent coping style or personality, might well be a key mechanism in the persistence of post-traumatic symptoms. Although psychological mechanisms most likely play a dominant role in the development of persistent symptoms, it could be hypothesized that acute physiological effects related to the injury lead to dysfunction of neural networks that are important for emotion...
Acute symptoms are probably the result of trauma-induced physiological changes, such as cell injury, inflammation and acute stress. It is thought that maladaptive pre-existent psychological factors (e.g., neuroticism, passive coping, and pre-injury mental distress) impede the ability to cope with acute symptoms leading to the persistence of symptoms. In this infographic we propose a scientific framework that can be used to further study this subject. Infographic was made by Rikkert Veltman Media Producties.

Fig. 2

Regulation. Patients with network dysfunction after mTBI may be prone to developing chronic stress, which facilitates the persistence of symptoms. It is also possible that patients with specific psychological characteristics (for example high neuroticism) have a stronger stress and pro-inflammatory response in the acute phase post-injury. These complex topics can be approached using a combination of personality and coping questionnaires, serum and salivary protein, cortisol and cytokine analyses, and emotion regulation fMRI-paradigms (e.g., cognitive reappraisal and mindfulness tasks). In addition to administering questionnaires, pre-injury stress levels can be assessed using hair cortisol analyses. A prolonged pro-inflammatory state, perhaps due to chronic elevations in cortisol, could contribute to persistent network dysfunction. This can be measured using longitudinal serum or salivary cytokine samples. It has to be realized that causality is a difficult issue (which is inherent in mTBI research), as neural network dysfunction itself may also lead to inflammation. Nonetheless, an intriguing question is to what extent the variance associated with network...
dysfunction and prolonged inflammation is explained by the injury itself, or by (pre-existent) psychological factors. This directly links to another fascinating question: can mTBI and subsequent stress and inflammatory responses alter coping skills, vulnerability to stress and personality?

Heart rate variability is reflective of autonomic nervous system reactivity related to acute stress, and is also a derivative of the function of neural networks that are responsible for emotion regulation and the ability to deal with stress [83]. Therefore, HRV can be used alongside fMRI-experiments to measure emotion and stress regulation in the acute or later stages after mTBI. For example, a key question is whether acute changes in HRV, measured during presentation at the emergency department, are related to dysfunction of neural networks at a later stage after mTBI.

In Fig. 2 we present a possible framework that can be used to investigate the interaction between aforementioned physiological and psychological factors. For the sake of clarity, a subdivision was made between the acute ‘physiological phase’ and the more sub-acute/chronic ‘psychological phase’, although we realize that there is much overlap between factors that fall under these phases. The key feature of this infographic is (subjective) post-traumatic symptoms, which develop in the acute phase, and the ultimate goal is to unravel the mechanisms that lead to persistent symptoms and poor outcome.

Conclusion

In summary, studies so far have suggested that sequelae after mTBI have a complex multifactorial etiology. A better understanding of the interaction between factors is warranted, especially regarding the interaction between acute (neuro)physiological and psychological factors. This understanding is needed to develop tailored psychological or pharmacological interventions in patients with mTBI, which are important future research goals. In our opinion, it is worthwhile to study the influence of acute physiological effects (i.e., cellular damage, inflammation, acute stress) after mTBI on neural networks that are important for emotion regulation, and to examine the possible interactions between these physiological effects and psychological factors such as personality characteristics, coping style, childhood psychotrauma or pre-injury mental distress.

Compliance with ethical standards

Conflicts of interest The authors declare that they have no conflict of interest.

Ethical standard An approval by an ethics committee was not applicable.

References

1. Bazarian JJ, McClung J, Shah MN et al (2005) Mild traumatic brain injury in the United States, 1998–2000. Brain Inj 19:85–91
2. Gardner RC, Yaffe K (2015) Epidemiology of mild traumatic brain injury and neurodegenerative disease. Mol Cell Neurosci 66:75–80. https://doi.org/10.1016/j.mcn.2015.03.001.Epidemiology
3. de Koning ME, Gareb B, el Moumni M et al (2016) Subacute post-traumatic complaints and psychological distress in trauma patients with or without mild traumatic brain injury. Injury 47:2041–2047. https://doi.org/10.1016/j.injury.2016.04.036
4. Dischinger PC, Ryb GE, Kufera JA, Auman KM (2009) Early predictors of postconcussive syndrome in a population of trauma patients with mild traumatic brain injury. J Trauma 66:287–289. https://doi.org/10.1097/TA.0b013e3181961da2
5. van der Naalt J, Timmerman ME, de Koning ME et al (2017) Early predictors of outcome after mild traumatic brain injury (UPFRONT): an observational cohort study. Lancet Neurol 16:532–540. https://doi.org/10.1016/S1474-4422(17)30117-5
6. Huang Y-L, Kuo Y-S, Tseng Y-C et al (2015) Susceptibility-weighted MRI in mild traumatic brain injury. Neurology 84:580–585. https://doi.org/10.1212/WNL.0000000000001237
7. Yuh EL, Mukherjee P, Lingsma HF et al (2013) Magnetic resonance imaging improves 3-month outcome prediction in mild traumatic brain injury. Ann Neurol 73:224–235. https://doi.org/10.1002/ana.23783
8. van der Horn HJ, de Haan S, Spikman JM et al (2018) Clinical relevance of microhemorrhagic lesions in subacute mild traumatic brain injury. Brain Imaging Behav 12:912–916. https://doi.org/10.1007/s11682-017-9743-6
9. Mayer AR, Bellgowan PS, Hanlon FM (2015) Functional magnetic resonance imaging of mild traumatic brain injury. Neurosci Biobehav Rev 49:8–18
10. van der Horn HJ, Liemburg EJ, Aleman A et al (2016) Brain networks subserving emotion regulation and adaptation after mild traumatic brain injury. J Neurotrauma 33:1–9. https://doi.org/10.1089/neu.2015.3905
11. Mercier E, Tardif P-A, Emond M et al (2017) Characteristics of patients included and enrolled in studies on the prognostic value of serum biomarkers for prediction of postconcussion symptoms following a mild traumatic brain injury: a systematic review. BMJ Open 7:e017848. https://doi.org/10.1136/bmjopen-2017-017848
12. Kim HJ, Tsao JW, Stanfill AG (2018) The current state of biomarkers of mild traumatic brain injury. JCI Insight 3:1–10. https://doi.org/10.1172/jci.insight.97105
13. Lingsma HF, Yue JK, Maas AI et al (2015) Outcome prediction after mild and complicated mild traumatic brain injury: external validation of existing models and identification of new predictors using the TRACK-TBI pilot study. J Neurotrauma 32:83–94. https://doi.org/10.1089/neu.2014.3384
14. Wáljas M, Iverson G, Lange R et al (2014) A prospective biopsychosocial study of the persistent post-concussion symptoms following mild traumatic brain injury. J Neurotrauma 54:1–54. https://doi.org/10.1089/neu.2014.3339
15. Zahniser E, Nelson LD, Dikmen SS et al (2019) The temporal relationship of mental health problems and functional limitations following mTBI: a TRACK-TBI and TED Study. J Neurotrauma. https://doi.org/10.1089/neu.2018.6172

16. Alarcon G, Eschleman KJ, Bowling NA (2009) Relationships between personality variables and burnout: a meta-analysis. Work Stress 23:244–263. https://doi.org/10.1080/02678379030282600

17. Charvat C, Bode S, Dernl B, Heekeren HR (2017) The effect of strategies, goals and stimulus material on the neural mechanisms of emotion regulation: a meta-analysis of fMRI studies. Neurosci Biobehav Rev 72:111–128. https://doi.org/10.1016/j.neubiorev.2016.11.014

18. Gill J, Merchant-Borna K, Jeromin A et al (2017) Acute plasma tau relates to prolonged return to play after concussion. Neurology 88:595–602. https://doi.org/10.1212/WNL.000000000003587

19. Shahim P, Tegner Y, Marklund N et al (2018) Neurofilament light and tau as guidelines for management of mild head injury is potentially cost saving. BMC Neurol 16:1–7. https://doi.org/10.1186/s12883-016-0723-z

20. Metting Z, Wilczak N, Rodiger LA et al (2012) GFAP and S100B in the acute phase of traumatic brain injury. Neurology 78:1428–1433. https://doi.org/10.1212/WNL.0b013e318253d5c7

21. Gardner RC, Rubenstein R, Wang KKW et al (2018) Age-related differences in diagnostic accuracy of plasma glial fibrillary acidic protein and tau for identifying acute intracranial trauma on computed tomography: a TRACK-TBI study. J Neurotrauma. https://doi.org/10.1089/neu.2017.54694

22. Cordiglia O, Anell A, Undén J (2016) The addition of S100B and immunocytochemical localization of S-100 antigen in normal human skin. Nature 294:85–87. https://doi.org/10.1038/294085a0

23. Bazarjan JJ, Biberthaler P, Welch RD et al (2018) Serum GFAP and UCH-L1 for prediction of absence of intracranial injuries on head CT (ALERT-TBI): a multicentre observational study. Lancet Neurol 17:782–789. https://doi.org/10.1016/S1474-4223(18)30231-X

24. Calcagnile O, Anell A, Undén J (2016) The addition of S100B and tau as blood biomarkers for sports-related concussion. Neurology 90:e1780–e1788. https://doi.org/10.1212/NEURO.0000000000005518

25. Paterson ZR, Holahan MR (2012) Understanding the neuroinflammatory response following concussion to develop treatment strategies. Front Cell Neurosci 6:1–10. https://doi.org/10.3389/fncel.2012.00058

26. Baratz R, Tweedie D, Wang J-Y et al (2015) Transiently lowering tumor necrosis factor-α synthesis ameliorates neuronal cell loss and cognitive impairments induced by minimal traumatic brain injury in mice. J Neuroinflammation 12:45. https://doi.org/10.1186/s12974-015-0237-4

27. Su S-H, Xu W, Li M et al (2014) Elevated C-reactive protein levels may be a predictor of persistent unfavourable symptoms in patients with mild traumatic brain injury: a preliminary study. Brain Behav Immun 38:111–117. https://doi.org/10.1016/j.bbi.2014.01.009

28. Mc Fie S, Abrahams S, Patricios J et al (2018) Inflammatory and apoptotic signalling pathways and concussion severity: a genetic association study. J Sports Sci 36:2226–2234. https://doi.org/10.1080/02640414.2018.1448570

29. Rathbone ATL, Tharmaradinam S, Jiang S et al (2015) A review of the neuro- and systemic inflammatory responses in post concussion symptoms: introduction of the “post-inflammation brain syndrome” PIBS. Brain Behav Immun 46:1–16. https://doi.org/10.1016/j.bbi.2015.02.009

30. Sun M, McDonald SJ, Brady RD et al (2018) The influence of immunological stressors on traumatic brain injury. Brain Behav Immun 69:618–628. https://doi.org/10.1016/j.bbi.2018.01.007

31. Huie JR, Diaz-Arrastia R, Yue JK et al (2019) Testing a multivariate proteomic panel for traumatic brain injury biomarker discovery: a TRACK-TBI pilot study. J Neurotrauma 36:100–110. https://doi.org/10.1089/neu.2017.5449

32. de Kloet ER, Joëls M, Holsboer F et al (2005) Stress and the brain: mechanisms of emotion regulation: a meta-analysis of fMRI studies. Neuropsychoendocrinology 30:294–303. https://doi.org/10.1038/sj.nen.1740151

33. de Kloet ER, Joëls M, Holsboer F et al (2005) Stress and the brain: mechanisms of emotion regulation: a meta-analysis of fMRI studies. Neuropsychoendocrinology 30:294–303. https://doi.org/10.1038/sj.nen.1740151

34. Broussard J, Acion L, De Jesús-Cortés H et al (2018) Repeated cell activation and white matter integrity in brains of active and recently retired national football league players. JAMA Neurol. https://doi.org/10.1001/jamaneurol.2018.003587

35. Broussard J, Acion L, De Jesús-Cortés H et al (2018) Repeated cell activation and white matter integrity in brains of active and recently retired national football league players. JAMA Neurol. https://doi.org/10.1001/jamaneurol.2018.003587

36. Patterson ZR, Holahan MR (2012) Understanding the neuroinflammatory response following concussion to develop treatment strategies. Front Cell Neurosci 6:1–10. https://doi.org/10.3389/fncel.2012.00058

37. Broussard J, Acion L, De Jesús-Cortés H et al (2018) Repeated cell activation and white matter integrity in brains of active and recently retired national football league players. JAMA Neurol. https://doi.org/10.1001/jamaneurol.2018.003587

38. Su S-H, Xu W, Li M et al (2014) Elevated C-reactive protein levels may be a predictor of persistent unfavourable symptoms in patients with mild traumatic brain injury: a preliminary study. Brain Behav Immun 38:111–117. https://doi.org/10.1016/j.bbi.2014.01.009

39. Mc Fie S, Abrahams S, Patricios J et al (2018) Inflammatory and apoptotic signalling pathways and concussion severity: a genetic association study. J Sports Sci 36:2226–2234. https://doi.org/10.1080/02640414.2018.1448570

40. Rathbone ATL, Tharmaradinam S, Jiang S et al (2015) A review of the neuro- and systemic inflammatory responses in post concussion symptoms: introduction of the “post-inflammation brain syndrome” PIBS. Brain Behav Immun 46:1–16. https://doi.org/10.1016/j.bbi.2015.02.009

41. Sun M, McDonald SJ, Brady RD et al (2018) The influence of immunological stressors on traumatic brain injury. Brain Behav Immun 69:618–628. https://doi.org/10.1016/j.bbi.2018.01.007

42. Huie JR, Diaz-Arrastia R, Yue JK et al (2019) Testing a multivariate proteomic panel for traumatic brain injury biomarker discovery: a TRACK-TBI pilot study. J Neurotrauma 36:100–110. https://doi.org/10.1089/neu.2017.5449

43. de Kloet ER, Joëls M, Holsboer F et al (2005) Stress and the brain: mechanisms of emotion regulation: a meta-analysis of fMRI studies. Neuropsychoendocrinology 30:294–303. https://doi.org/10.1038/sj.nen.1740151

44. de Kloet ER, Joëls M, Holsboer F et al (2005) Stress and the brain: mechanisms of emotion regulation: a meta-analysis of fMRI studies. Neuropsychoendocrinology 30:294–303. https://doi.org/10.1038/sj.nen.1740151

45. Slavich DC, Graham-Engeland JE, Smyth JM, Engeland CG (2015) Salivary markers of inflammation in response to acute stress. Brain Behav Immun 44:253–269. https://doi.org/10.1016/J.BBI.2014.08.008

46. Newton TL, Fernandez-botran R, Lyle KB et al (2017) Salivary cytokine response in the aftermath of stress: an emotion regulation perspective. Emotion 17:1007–1020

47. Slavich GM, Irwin MR (2014) From stress to inflammation and major depressive disorder: a social signal transduction theory of depression. Psychol Bull 140:774–815. https://doi.org/10.1037/a0035302

48. Bohnen N, Jolles J, Twijnsstra A et al (1992) Coping styles, cortisol reactivity, and performance in a vigilance task of patients with persistent postconcussive symptoms after a mild head injury. Int
49. Sojka P, Stålnacke B-M, Björnstig U, Karlsson K (2006) One-year follow-up of patients with mild traumatic brain injury: occurrence of post-traumatic stress-related symptoms at follow-up and serum levels of cortisol, S-100B and neuron-specific enolase in acute phase. Brain Inj 20:613–620. https://doi.org/10.1080/0269905060676982

50. Woolf PD, Cox C, Kelly M et al (1990) The adrenocortical response to brain injury: correlation with the severity of neurologic dysfunction, effects of intoxication, and patient outcome. Alcohol Clin Exp Res 14:917–921

51. Sung C-W, Chen K-Y, Chiang Y-H et al (2016) Heart rate variability and serum level of insulin-like growth factor-1 are correlated with symptoms of emotional disorders in patients suffering a mild traumatic brain injury. Clin Neurophysiol 127:1629–1638. https://doi.org/10.1016/j.clinph.2015.08.006

52. Bay E, Xie Y (2009) Psychological and biological correlates of fatigue after mild-to-moderate traumatic brain injury. West J Nurs Res 31:731–747. https://doi.org/10.1177/0193945909334856

53. Rothman MS, Arciniegas DB, Filley CM, Wierman ME (2007) The neuroendocrine effects of traumatic brain injury. J Neuropsychiatry Clin Neurosci 19:363–372. https://doi.org/10.1176/jnp.2007.19.4.363

54. Tanriverdi F, Unluhisarcikli K, Kelestimur F (2010) Pituitary function in subjects with mild traumatic brain injury: a review of literature and proposal of a screening strategy. Pituitary 13:146–153. https://doi.org/10.1007/s11195-010-0215-x

55. Stein MB, Jain S, Giacino JT et al (2019) Risk of posttraumatic stress disorder and major depression in civilian patients after mild traumatic brain injury. JAMA Psychiatry 76:249. https://doi.org/10.1001/jamapsychiatry.2018.4288

56. Veer IM, Oei NY, Spinholven P et al (2011) Beyond acute social stress: increased functional connectivity between amygdala and cortical midline structures. Neuroimage 57:1534–1541. https://doi.org/10.1016/j.neuroimage.2011.05.074

57. Guendelman S, Medeiros S, Ramesh H (2017) Mindfulness and emotion regulation: insights from neurobiological, psychological, and clinical studies. Front Psychol 8:220. https://doi.org/10.3389/fpsyg.2017.00220

58. Ochsner KN, Gross JJ (2008) Cognitive emotion regulation: insights from social cognitive and affective neuroscience. Curr Dir Psychol Sci 17:153–158. https://doi.org/10.1111/j.1469-7610.2008.00566.x

59. Wang X, Xie H, Cotton AS et al (2017) Early changes in cortical emotion processing circuits after mild traumatic brain injury from motor vehicle collision. J Neurotrauma 34:273–280. https://doi.org/10.1089/neu.2015.4392

60. Raichle ME (2015) The brain’s default mode network. Annu Rev Neurosci 38:413–427. https://doi.org/10.1146/annurev-neuro-070113-014030

61. Xie X, Mulej Bratec S, Schmid G et al (2016) How do you make me feel better? Social cognitive emotion regulation and the default mode network. Neuroimage 134:270–280. https://doi.org/10.1016/j.neuroimage.2016.04.015

62. Whitfield-Gabrieli S, Ford JM (2012) Default mode network activity and connectivity in psychopathology. Annu Rev Clin Psychol 8:49–76. https://doi.org/10.1146/annurev-clinpsych-032511-143049

63. Marsland AL, Kuan DC-H, Sheu LK et al (2017) Systemic inflammation and resting state connectivity of the default mode network. Brain Behav Immun 62:162–170. https://doi.org/10.1016/j.bbi.2017.01.013

64. Chiesa A, Serretti A, Jakobsen JC (2013) Mindfulness: top-down or bottom-up emotion regulation strategy? Clin Psychol Rev 33:82–96

65. Doll A, Höfel BK, Boucard CC et al (2015) Mindfulness is associated with intrinsic functional connectivity between default mode and salience networks. Front Hum Neurosci 9:1–11. https://doi.org/10.3389/fnhum.2015.00461

66. Seeley WW, Menon V, Schatzberg AF et al (2007) Dissociable intrinsic connectivity networks for salience processing and executive control. J Neurosci 27:2349–2356. https://doi.org/10.1523/JNEUROSCI.5587-06.2007

67. Thomason ME, Hamilton JP, Gotlib IH (2011) Stress-induced activation of the HPA axis predicts connectivity between subgenual cingulate and salience network during rest in adolescents. J Child Psychother Psychiatry 52:1026–1034. https://doi.org/10.1111/j.1469-7610.2011.02422.x

68. Critchley HD (2005) Neural mechanisms of autonomic, affective, and cognitive integration. J Comp Neurol 493:154–166. https://doi.org/10.1002/cne.20749

69. Azulay J, Smart C, Mott T, Cicerone K (2013) A pilot study examining the effect of mindfulness-based stress reduction on symptoms of chronic mild traumatic brain injury/postconcussive syndrome. J Head Trauma Rehabil 28:323–331. https://doi.org/10.1097/HTR.0b013e318250ebda

70. van der Horn HJ (2017) Adaptation after mild traumatic brain injury: the role of structural and functional brain networks. Dissertation

71. Young KS, van der Velden AM, Craske MG et al (2018) The impact of mindfulness-based interventions on brain activity: a systematic review of functional magnetic resonance imaging studies. Neurosci Biobehav Rev 84:424–433. https://doi.org/10.1016/j.neubiorev.2017.08.003

72. Simon R, Engström M (2015) The default mode network as a biomarker for monitoring the therapeutic effects of meditation. Front Psychol 6:776. https://doi.org/10.3389/fpsyg.2015.00776

73. van der Horn HJ, Liemburg EJ, Scheenen ME et al (2016) Brain network dysregulation, emotion, and complaints after mild traumatic brain injury. Hum Brain Mapp 37:1645–1654. https://doi.org/10.1002/hbm.23126

74. van der Horn HJ, Scheenen ME, de Koning ME et al (2017) The default mode network as a biomarker of persistent complaints after mild traumatic brain injury: a longitudinal fMRI study. J Neurotrauma. https://doi.org/10.1089/neu.2017.5185

75. McCuddy WT, España LY, Nelson LD et al (2018) Association of acute depressive symptoms and functional connectivity of emotional processing regions following sport-related concussion. NeuroImage Clin 19:434–442. https://doi.org/10.1016/j.nicli.2018.05.011

76. Mayer AR, Mannell MV, Ling J et al (2011) Functional connectivity in mild traumatic brain injury. Hum Brain Mapp 32:1825–1835. https://doi.org/10.1002/hbm.21151

77. Palacios EM, Yuh EL, Chang Y-S et al (2017) Resting-state functional connectivity alterations associated with six-month outcomes in mild traumatic brain injury. J Neurotrauma. https://doi.org/10.1089/neu.2016.4752

78. Kaushal M, España LY, Nencka AS et al (2019) Resting-state functional connectivity after concussion is associated with clinical recovery. Hum Brain Mapp 40:1211–1220. https://doi.org/10.1002/hbm.24440

79. Dretsch MN, Daniel TA, Goodman AM et al (2017) Differential neural activation when voluntarily regulating emotions in service members with chronic mild traumatic brain injury. Appl Neuropsychol. https://doi.org/10.1080/23279095.2017.1562406

80. Gillie BL, Thayer JF (2014) Individual differences in resting heart rate variability and cognitive control in posttraumatic stress disorder. Front Psychol 5:1–7. https://doi.org/10.3389/fpsyg.2014.00758

81. Williams DP, Cash C, Rankin C et al (2015) Resting heart rate variability predicts self-reported difficulties in emotion regulation.
regulation: a focus on different facets of emotion regulation. Front Psychol 6:1–8. https://doi.org/10.3389/fpsyg.2015.00261
82. Koshino H, Minamoto T, Yaoi K et al (2014) Coactivation of the default mode network regions and working memory network regions during task preparation. Sci Rep 4:5954. https://doi.org/10.1038/srep05954
83. Thayer JF, Lane RD (2000) A model of neurovisceral integration in emotion regulation and dysregulation. J Affect Disord 61:201–216
84. Friedman BH, Thayer JF (1998) Autonomic balance revisited: panic anxiety and heart rate variability. J Psychosom Res 44:133–151
85. Friedman BH (2007) An autonomic flexibility-neurovisceral integration model of anxiety and cardiac vagal tone. Biol Psychol 74:185–199. https://doi.org/10.1016/j.biopsycho.2005.08.009
86. Brosschot JF, Gerin W, Thayer JF (2006) The perseverative cognition hypothesis: a review of worry, prolonged stress-related physiological activation, and health. J Psychosom Res 60:113–124. https://doi.org/10.1016/j.jpsychosomat.res.2005.06.074
87. Hansen AL, Johnsen BH, Sollers JJ et al (2004) Heart rate variability and its relation to prefrontal cognitive function: the effects of training and detraining. Eur J Appl Physiol 93:263–272. https://doi.org/10.1007/s00421-004-1208-0
88. Abaji JP, Curnier D, Moore RD, Ellemberg D (2016) Persisting effects of concussion on heart rate variability during physical exertion. J Neurotrauma 33:811–817. https://doi.org/10.1089/neu.2015.3989
89. Bishop SA, Dech RT, Guzik P, Neary JP (2017) Heart rate variability and implication for sport concussion. Clin Physiol Funct Imaging. https://doi.org/10.1111/cpf.12487
90. Sentihathan A, Mainwaring LM, Psych C, Hutchison M (2017) Heart rate variability of athletes across concussion recovery milestones: a preliminary study setting: interuniversity sports teams at a single institution. Clin J Sport Med 27:288–295. https://doi.org/10.1097/JSM.0000000000000337
91. Francis HM, Fisher A, Rushby JA, McDonald S (2016) Reduced heart rate variability in chronic severe traumatic brain injury: association with impaired emotional and social functioning, and potential for treatment using biofeedback. Neuropsychol Rehabil 26:103–125. https://doi.org/10.1080/09602011.2014.1003246
92. Liao KH, Sung CW, Chu SF et al (2016) Reduced power spectra of heart rate variability are correlated with anxiety in patients with mild traumatic brain injury. Psychiatry Res 243:349–356. https://doi.org/10.1016/j.psychres.2016.07.001
93. Deepika A, Devi BI, Shukla DP et al (2018) Neuroimmunology of traumatic brain injury: a longitudinal study of interdependency of inflammatory markers and heart rate variability in severe traumatic brain injury. J Neurotrauma. https://doi.org/10.1089/neu.2017.5151