Traumatic Alterations in Consciousness: Traumatic Brain Injury

Brian J. Blyth, MD*; Jeffrey J. Bazarian, MD, MPH

The purpose of this review is to provide an overview of mild traumatic brain injury (mTBI) in a form useful for emergency physicians. mTBI is a disease with considerable public health impact and is the subject of a vast amount of current research. The authors' discussion of this disease is necessarily limited both by space constraints and the interests of their target audience. For more detailed discussions of various topics covered in this review, the reader is directed to the publications listed in Table 1.

DEFINITION OF TRAUMATIC BRAIN INJURY

Traumatic brain injury (TBI) is defined as any traumatically induced structural injury or physiologic disruption of brain function as a result of an external force. TBI is manifested by one or more clinical signs occurring immediately afterwards including a loss, decreased, or altered level of consciousness, amnesia, neurologic deficit, or intracranial lesion.1 External forces may include direct impact of the head with another object, indirect forces from acceleration/deceleration, or a blast injury. The Glasgow Coma Score (GCS) has traditionally been used to classify TBI as mild (GCS 13–15), moderate (GCS 9–12), or severe (GCS 3–8). A more recent classification scheme for TBI uses length of loss of consciousness (LOC), alteration of consciousness (AOC), and posttraumatic amnesia (PTA), as well as imaging findings to categorize TBI (Table 2).1 It is important to stress that mild TBI (mTBI) is clinically defined based solely on self-reported or observed symptoms, and often occurs with normal neuroimaging.

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Indeed, the mTBI definition included in Table 1 classifies TBI with positive neuroimaging as at least moderate in severity. Two older, but still commonly used mTBI definitions by the American Congress of Rehabilitation Medicine\(^2\) and the Centers for Disease Control and Prevention\(^3\) define patients as mild if they have positive

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**Table 1**

**Recommended further reading**

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**Table 2**

**Clinical criteria for TBI severity**

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<th>Mild</th>
<th>Moderate</th>
<th>Severe</th>
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<tr>
<td>Structural imaging</td>
<td>Definition dependent(^a)</td>
<td>Normal or abnormal</td>
<td>Normal or abnormal</td>
</tr>
<tr>
<td>Loss of consciousness (LOC)</td>
<td>0–30 min</td>
<td>&gt;30 min and &lt;24 h</td>
<td>&gt;24 h</td>
</tr>
<tr>
<td>Alteration of consciousness (AOC)(^b)</td>
<td>A moment up to 24 h</td>
<td>&gt;24 h. Severity based on other criteria</td>
<td></td>
</tr>
<tr>
<td>Posttraumatic amnesia (PTA)</td>
<td>0–1 d</td>
<td>&gt;1 and &lt;7 d</td>
<td>&gt;7 d</td>
</tr>
<tr>
<td>GCS (best score in first 24 h)</td>
<td>13–15</td>
<td>9–12</td>
<td>&lt;9</td>
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\(^a\) Patients who otherwise meet the clinical criteria for mTBI but have intracranial imaging abnormalities may be classified as complicated mTBI or moderate TBI depending on the definition used.

\(^b\) Alteration of mental status must be immediately related to the trauma to the head. Typical symptoms may include looking or feeling dazed, confusion, difficulty thinking clearly or responding appropriately to mental status questions, or inability to describe events immediately before or after the traumatic event.

neuroimaging findings but meet all other clinical criteria for mTBI. Patients who have abnormal intracranial imaging but otherwise meet a clinical definition of mTBI are referred to as complicated mTBI.4

Concussion is a common term for mTBI and will be used interchangeably within this article. Our understanding of TBI remains rudimentary relative to many other medical problems of similar magnitude. A symptom-based classification uses the description of symptoms evident on history and physical examination to classify illness. This method is imprecise, often grouping disparate pathophysiological processes together as single clinical entities. This imprecision is particularly problematic in moderate and severe TBI where multiple injury processes as evidenced by heterogeneous imaging findings are often present simultaneously. As knowledge of a disease increases and diagnostic tools improve, a more sophisticated classification emerges that may include anatomic, physiologic, metabolic, immunologic, and genetic factors. TBI has been the subject of intensive research in recent years, and recommendations for improved classification of this diverse disease are beginning to appear in the literature.5

**Epidemiology**

An estimated 1.4 million Americans presented to emergency departments for medical care after TBI each year between 1995 and 2001.6 Over 1 million of these patients had mTBI. The incidence of ED-attended concussion is 444 in 100,000 in the United States;7 however, it is thought that nearly 40% of patients suffering mTBI do not seek hospital-based care, and 25% do not report their injuries to health care providers at all.8 An estimated 5% to 25% of all patients with concussion have postconcussive symptoms or other cognitive deficits that persist beyond 1 year.9–11 This number is greater than the annual incidence of multiple sclerosis, Parkinson disease, myasthenia gravis, and Huntington disease combined.12 TBI is more common in men than women, with 60% of TBI occurring in males.6 Young children, adolescents, and the elderly suffer the highest rates of TBI.6 The most common mechanisms of TBI are falls, automobile accidents, being struck by or against an object, and assault. The rate of hospitalizations for TBI fell dramatically during the 1980s and 1990s.6 Wounded veterans from the wars in Iraq and Afghanistan may represent a nontrivial increase in patients living with mTBI. Approximately 20% of troops returning from combat deployments in Iraq have clinician-confirmed mTBI. These patients typically have TBI resulting from a blast,13 and often have comorbid posttraumatic stress disorder (PTSD) or depression.14

**Biomechanics**

Traumatic injury results from the transfer of energy from the environment to tissue that is greater than the amount that can be absorbed without dysfunction. Biomechanics is the study of the interaction of forces and physical responses in biologic systems. Traumatic insults generally occur over short periods of time and are referred to as dynamic loading. Dynamic loading includes both direct or impact loading, as well as impulsive loading whereby no physical contact occurs. The loads absorbed by the brain after trauma generally include linear and rotational components called angular loads. The rate and duration of the insult are important because loads applied at high rates tend to result in more damage.15 For example, the force involved in punching a wall can also be applied by pressing your fist against that same wall for a few minutes: the former instance results in a boxer’s fracture whereas the latter does not. Focal injury such as contusion results from direct loading and often occurs in the absence of widespread injury. In contrast, diffuse axonal injury (DAI) often occurs
as a result of the rotational acceleration accompanying indirect loading.\textsuperscript{16} Humans are particularly susceptible given their large cranium connected to the trunk by relatively weak neck musculature. Rotational acceleration produces substantial and widespread strains within the brain resulting from both acceleration and deceleration. These diffuse strains lead to differential movement of the brain relative to the skull, which can cause hemorrhage. Shear strain is most prominent after rotational injury, and brain tissue is particularly sensitive to this type of strain.\textsuperscript{17} In animal models, rotational acceleration is required to produce concussion, whereas isolated linear acceleration produced contusions and subdural hematomas but no LOC.\textsuperscript{18}

\textbf{Pathophysiology}

The initial traumatic insult results in mechanical damage including rupture of cellular and vascular membranes with release of intracellular contents, ultrastructural damage of axons, and changes in cerebral blood flow.\textsuperscript{19,20} Subsequent metabolic derangement includes widespread release of excitatory neurotransmitters such as glutamate, severe dysregulation of calcium homeostasis, energy failure due to adenosine triphosphate depletion, free radical generation, and cell death by necrotic and apoptotic pathways.\textsuperscript{20,21} More global consequences of the traumatic insult include increased intracranial pressure, decreased cerebral blood flow, tissue ischemia, cerebral edema, and functional blood-brain barrier dysfunction.\textsuperscript{22,23} Following the initial damage, repair and recovery processes begin through the removal of cellular debris, glial scar formation, and plastic changes in neural networks.\textsuperscript{24} Because of the difficulties studying human mTBI, the mechanisms described in this section were derived principally from animal studies and, to a lesser degree, from humans with severe TBI (sTBI). Similar processes are thought to occur in human mTBI.

\textbf{Putative Causes of Altered Consciousness in mTBI}

The definitive causes of altered consciousness are not known. LOC requires either loss of the function of both cerebral hemispheres or of the reticular activating system. Several plausible hypothetical mechanisms have been proposed for the AOC that occurs with mTBI. These hypotheses include the reticular, pontine-cholinergic system, centripetal, and convulsive hypotheses. The reticular activating system (RAS) resides in the brainstem reticular formation, which extends from the top of the spinal column to the rostral midbrain with extensions into the thalamus and hypothalamus. The RAS is excited by input from surrounding sensory tracts and transmits this excitation to the cortex to induce generalized cortical and behavioral arousal. In the absence of input from the RAS, consciousness is impaired.

Under the reticular hypothesis of concussion, LOC after brain trauma results from a disturbance or depression of the activity of polysynaptic pathways within the RAS.\textsuperscript{25} It is not completely understood how a traumatic dysfunction of the RAS occurs; however, it is believed to result from shearing or tensile strains on RAS pathways at the craniocervical junction. Neuropathological evidence for this is limited. The hypothesis also fails to address traumatic amnesia. A further difficulty is that EEG findings do not support depression of the RAS in concussion.

The pontine-cholinergic system hypothesis differs from the reticular activating system hypothesis in that RAS dysfunction is thought to occur as a consequence of trauma-induced activation of the inhibitory cholinergic system of the dorsal pontine tegmentum.\textsuperscript{26} In animal models, injection of cholinergic agonists into the brainstem induces unconsciousness\textsuperscript{27,28} whereas similar injections of cholinergic antagonists reduce the duration of traumatic unconsciousness.\textsuperscript{29} Furthermore, electroencephalographic (EEG) studies show widespread neuronal discharge after concussion, and
elevated acetylcholine is found in the cerebrospinal fluid of patients after TBI. However, it is not clear that activation of this system can produce LOC due to RAS suppression.

The centripetal hypothesis posits that sudden rotational forces cause shearing strains and stresses that result in functional decoupling of nerve fibers.\textsuperscript{18} The depth of this functional decoupling is directly related to the extent of rotational acceleration delivered to the brain. Also, with greater rotational acceleration the likelihood of mechanical injury to fibers increases. Lower inertial forces that result in functional decoupling between the subcortex or diencephalon and the cortex may result in amnesia or confusion without LOC. Furthermore, greater forces resulting in decoupling between more superficial structures and the mesencephalon result in LOC. This hypothesis nicely explains posttraumatic amnesia and dazed states; however, it also requires very high energy injuries to cause full LOC. Consequently, patients with LOC would often have accompanying structural brain injury that simply is not observed.

Patients with concussion have similar symptoms to those who have experienced generalized epileptic seizures or electroconvulsive therapy (ECT). This overlap of symptoms has led to speculation that similar pathophysiological events occur in all 3 conditions. Close observation of human patients and animal models shows that concussive injury generally causes an initial convulsive event followed by a longer and more prominent paralytic phase.\textsuperscript{30,31} EEG recordings from concussed animals also show initial, transient epileptiform activity. According to the convulsive hypothesis the symptoms associated with concussion are due to direct injury to neurons, resulting in hyperexcitability and widespread membrane depolarization followed by neuronal exhaustion.\textsuperscript{32} These 2 neuronal states correspond to the convulsive and paralytic phases, respectively.

The convulsive hypothesis is able to reasonably account for a broader range of postconcussive behaviors than its competitors, including LOC, amnesia, convulsive movements, autonomic disturbances, and the dazed or “dinged” state.\textsuperscript{33} While this hypothesis does a better job than the others at providing a unified explanation for the broad range of symptoms observed as an acute result of mTBI, it does not account for the structural abnormalities that occur as a result of mTBI. In summary, none of the individual hypotheses currently available explain all the findings seen with mTBI. Given the often complementary strengths and weaknesses of the 4 hypotheses discussed above, it seems likely that the mechanisms of altered consciousness after TBI may be due to a combination of processes. For a detailed explanation of these hypotheses, the reader is directed to the excellent review by Shaw.\textsuperscript{33}

**DIAGNOSIS OF MTBI**

**Clinical Presentation**

A 28-year-old man presents to the Emergency Department (ED) after a motor vehicle accident. He was the restrained driver in a car that skidded of the road in icy conditions and collided head-on with a tree at 50 mph. There was airbag deployment. The paramedic reports that he was unconscious initially but that he was alert and oriented during transport.

**Differential Diagnosis**

The diagnosis of mTBI is made clinically and relies heavily on the history obtained from the patient and any witnesses. Obtaining a reliable history is often difficult because of posttraumatic amnesia, persistent altered mental status, or intoxication, a frequent
comorbid factor in mTBI patients. Diagnoses with similar presentations include seizure, syncope, intoxication, malingering, anxiety, and other psychiatric conditions.

**Clinical Criteria**

Several clinical criteria for the diagnosis of mTBI exist.\(^1\)–\(^3\) Concussion or mTBI is defined as a LOC of less than 30 minutes or amnesia lasting less than 24 hours, or any period of altered mental status at the time of injury. In conjunction, patients must also have a GCS of 13 to 15 and normal structural imaging to meet the criteria for mTBI. Lower GCS scores classify the patient as having moderate (GCS 9–12) or severe (GCS 3–8) TBI.

**Imaging**

The imaging study of choice in mTBI is noncontrast head computed tomography (CT). This study is preferred over others because it is sensitive for traumatic injuries that require neurosurgical intervention including acute bleeding, increased intracranial pressure, and skull fracture. Although as many as 15% of mTBI patients will have an acute injury detected by noncontrast head CT, only 1% of those abnormalities require neurosurgical intervention.\(^34\)–\(^40\) Other imaging modalities are of limited use for the clinical evaluation of mTBI patients and are not recommended. Although magnetic resonance imaging (MRI) is 30% more sensitive than CT for the detection of traumatic abnormalities after mTBI,\(^41\) there is no evidence that it identifies more patients requiring neurosurgical intervention.\(^42\) More exotic imaging modalities including functional MRI, diffusion tensor imaging MRI, magnetic resonance spectroscopy, single-photon emission computed tomography, and positron emission tomography are valuable research tools but do not have proven clinical utility.

**Decision Rules**

The low rate of clinically important brain injury seen on head CT obtained acutely after mTBI has resulted in efforts to minimize unnecessary studies through the application of rigorously validated clinical decision rules. Two major decision rules applying to adult mTBI patients include the Canadian CT Head Rule\(^43\) and the New Orleans Criteria.\(^44\) For patients with a GCS of 15, both of these rules have equivalent sensitivities for detecting injuries requiring neurosurgical intervention; however, the Canadian CT Head Rule has a higher specificity for some clinical outcomes and its use may reduce imaging rates.\(^45\) In pediatric populations, increased concern for radiation exposure and the potential requirement for sedation make minimizing unnecessary CT after mTBI even more compelling than in adult populations. Kuppermann and colleagues\(^46\) recently reported a very sensitive decision rule that identifies low-risk children who do not require a head CT after mTBI. Indications for obtaining head CT after mTBI based on these decision rules are summarized in Fig. 1.

**Biomarkers**

There is substantial interest in developing protein biomarkers obtained from serum to aid the diagnosis and guide the treatment of TBI of all severities. Although several potential biomarkers have been studied,\(^47\)–\(^53\) to date only serum S100B has accepted clinical utility for mTBI. Specifically, elevated S100B has a high negative predictive value for clinically important injury on head CT after mTBI. In a large cohort, elevated S100B was 99% sensitive for the detection of injury on CT scan,\(^54\) prompting the use of this test as the clinical standard of care in several European countries. The test has the added advantage of not being affected by concomitant alcohol intoxication.\(^55\) Although not yet approved by the Food and Drug Administration (FDA) in the United
Fig. 1. Indications for obtaining noncontrast head CT after TBI. Noncontrast head CT is the study of choice to evaluate TBI patients for clinically important neurotrauma. Clinically important neurotrauma is defined as any traumatically induced intracranial injury that requires neurosurgical intervention or requires hospital admission and neurosurgical follow-up. Clinically important TBI occurs rarely after mTBI, therefore minimizing unnecessary head CT scans is desirable. This figure integrates validated decision rules for both adult and pediatric patients designed to minimize unnecessary CT scans after mTBI.

1. hematympanum, raccoon eyes, CSF otorrhea or rhinorrhea, Battle's Sign
2. agitation, somnolence, repetitive questioning, slow response to verbal communication
3. severe mechanisms include:
   a. motor vehicle accident with ejection, rollover, or death of another occupant
   b. falls 6 > 3 feet (ages 2 and over), or > 1.5 feet (under age 2)
   c. pedestrian or bicyclist struck by motor vehicle
   d. head struck by high-impact object.
States, the American College of Emergency Physicians recently issued a guideline stating that for mTBI patients with serum S100B concentrations of less than 0.1 μg/mL measured within 4 hours of injury, it is reasonable to consider not obtaining a head CT. Multicenter studies to evaluate the accuracy of this test in United States patient populations are currently underway and may provide the data necessary for FDA approval.

**TREATMENT OF MTBI**

*Clinical Presentation*

After receiving the prehospital provider’s report, you note that the patient is complaining of headache, back pain, and abdominal pain. He has no significant medical history and takes no medications. Pertinent findings on examination include a slight tachycardia, an abrasion on his forehead, mild tenderness in his upper abdomen, and diffuse tenderness over his lumbar spine. He is alert and oriented but cannot remember any events since the accident. Otherwise, he has a normal neurologic examination.

*ED Priorities*

Initial assessment of the mTBI patient in the ED is focused on identifying patients who may require medical or neurosurgical intervention for the treatment of increased intracranial pressure or an expanding mass lesion. Patients with “red flag” conditions such as altered mental status, papillary asymmetry, seizures, repeated vomiting, double vision, worsening headache, motor or sensory deficits, or ataxia should have an emergent noncontrast head CT scan performed. See Fig. 1 for further imaging recommendations.

Patients with intracranial imaging abnormalities or declining mental status require immediate neurosurgical consultation. Worsening mental status is typically caused by increasing intracranial pressure (ICP) leading to compromised cerebral blood flow and oxygen delivery. For these patients, airway management with endotracheal intubation to protect against aspiration as well as to control ventilation should be considered. Nonsurgical management also includes mitigating ICP increases by raising the head of the bed to 30° and treatment with hyperosmolar agents such as intravenous mannitol. Finally, brief periods of hyperventilation can also reduce dangerous ICP increases. Mechanistically, hyperventilation causes vasoconstriction and reduces ICP by decreasing cerebral blood flow. Overaggressive hyperventilation can result in ICP decreases at the expense of adequate tissue perfusion. Therefore, prolonged hyperventilation should be used only when other therapies have failed.

*Clinical Presentation*

You obtain imaging studies, including CT of head, neck, abdomen, and pelvis, which reveal no traumatic injuries. After returning from radiology, the patient does not recall meeting you. Although his head CT did not reveal a traumatic injury, he is admitted for observation overnight due to his persistent anterograde amnesia. He has an uneventful night and his amnesia resolves. He receives detailed discharge instructions that include a description of “red flag conditions,” common postconcussive symptoms, and reassurance that the vast majority of patients recover completely from concussion. He is advised to avoid activities that exacerbate his symptoms, to take acetaminophen as needed for headache, and to follow up with a visit to a local concussion clinic in 1 week if he is having any persistent discomfort from his concussion. He is accompanied home by his fiancée who will stay with him over the next 24 hours.
**Acute Phase: Within 1 Week of Injury**

After evaluating for “red flag” signs and symptoms (see ED Priorities), a thorough history of symptoms including loss or alteration of consciousness, headache, irritability, unsteadiness, vertigo, photophobia, or phonophobia should be obtained. The physical examination includes a focused neurologic examination including assessment of cranial nerves, postural instability, visual function, and mental status. Noncontrast head CT should be obtained when indicated (see Fig. 1). Neurosurgical consultation is necessary for patients with imaging abnormalities. These patients are often admitted for 24 hours for ongoing mental status monitoring and repeat head CT prior to discharge. Patients in whom imaging is not indicated or who have a normal head CT may be safely discharged.\(^{56–58}\)

Discharge instructions for mTBI patients include 2 principal elements: symptoms requiring immediate reevaluation (see ED Priorities) and postconcussive symptom education. Postconcussive symptoms include headache, sleep disturbances, vertigo, nausea, fatigue, sensitivity to light or noise, attention and concentration problems, depression, and emotional lability. The vast majority of adults with postconcussive symptoms recover within 3 to 12 months.\(^{59}\) Early patient education that includes likely postconcussive symptoms and reassurance about an expected positive recovery has been shown to speed recovery and decrease postconcussive symptoms.\(^{60–62}\) Headache should be managed with acetaminophen. Nonsteroidal anti-inflammatory drugs (NSAIDs) may be used in patients with negative neuroimaging but should be deferred until 48 hours after injury if imaging was not obtained. In addition, narcotics should be avoided in the treatment of posttraumatic headache.

Pharmacologic treatment of other postconcussive symptoms is not recommended in the acute phase. Rather, patients with symptoms other than headache should be advised to rest and encouraged to return to normal activity as soon as possible. However, individuals whose normal activity includes a high risk for re-injury should have careful evaluation of their symptoms and examination findings with consideration of their specific activities that result in a high injury risk. Specific limitations on activity may be recommended for these patients to mitigate their individual risk. Patients reporting fatigue may be given a graded return to work or activity. For patients with normal activities involving significant physical activity, exertional testing may be performed. If this results in a return of symptoms, a monitored progressive return to these activities as tolerated should be recommended.\(^{1}\)

**Clinical Presentation**

Two weeks after the initial injury, the patient continues to suffer from frequent headaches that are only slightly relieved by acetaminophen. He also complains of increased irritability, sleepiness, and difficulty concentrating. During an initial follow-up visit, a detailed history and physical examination fails to reveal comorbid psychiatric or physical problems including PTSD, depression, substance abuse, hypertension, cervical spine abnormalities, sinus infections, or visual acuity deficits. However, the patient indicates that he has not been sleeping well due to persistent headache. He is started on an NSAID for his headaches, and provided with education regarding good sleep hygiene and relaxation techniques. He is also advised to begin a regular exercise program. A follow-up appointment is scheduled in 4 weeks.

**Initial Management of Postconcussive Symptoms**

This section provides an overview of therapy for the initial treatment of patients with mTBI and symptoms lasting more than 1 week after injury. Patients with a delayed
initial presentation should also be treated according to these guidelines. Detailed recommendations for the evaluation and treatment of specific symptoms can be found in the VA/DoD Clinical Practice Guideline for Management of Concussion/mTBI.\(^1\)

**Symptom classification and goals of therapy**

Postconcussive symptoms generally fall into 3 categories: physical, cognitive, and behavioral or emotional. Typical physical symptoms include headache, nausea, vomiting, dizziness, fatigue, blurred vision, sleep disturbances, light or noise sensitivity, balance problems, and transient neurologic abnormalities. Cognitive symptoms may occur with attention, concentration, memory, processing speed, judgment, and executive functioning. Behavioral/emotional symptoms include depression, anxiety, agitation, irritability, and aggression.

Because there is an incomplete understanding of the etiology of symptoms after mTBI, the goal of intervention for postconcussive symptoms is to improve identified problems rather than affect a cure. It is believed that symptoms resulting from mTBI are interrelated, and alleviation of one symptom often leads to improvement in others. Postconcussive symptoms are also common to many other psychiatric ailments including depression, anxiety disorders, PTSD, and substance abuse disorders. Indeed, there is substantial evidence that affective disorders, PTSD, and substance abuse disorders are often associated with mTBI.\(^{59,63,64}\) These disorders are also associated with higher rates of persistent postconcussive symptoms.\(^{65,66}\) Consequently, aggressive treatment of any comorbid psychiatric illness may help to improve postconcussive symptoms.

**Patient evaluation**

Patient evaluation should include a thorough history, physical examination, and review of the medical record. A review of sleep habits is particularly important as poor sleep may contribute to symptoms including headache, fatigue, anxiety, irritability, depressive thoughts, poor concentration, memory difficulties, and poor decision making. TBI patients should also be screened for psychiatric conditions including PTSD, depression, and substance abuse disorders. Low-yield diagnostic testing should be minimized. There is limited evidence to support the utility of comprehensive neuropsychological/cognitive testing within the first 30 days of mTBI, and a focused clinical interview is sufficient to assess for cognitive difficulties.\(^{67}\) Laboratory studies including electrolytes, a complete blood count, and thyroid function testing may be useful, particularly when evaluating behavioral and cognitive symptoms. Imaging studies are of limited use.

Physical symptoms should prompt a search for treatable causes. Screening patients with headaches for preexisting headache conditions, hypertension, cervical spine abnormalities, sinus infections, and visual acuity deficits may provide useful avenues of treatment. Symptoms related to dizziness including poor coordination, unsteadiness, vertigo, or loss of balance may be due to medication effects, orthostatic hypotension, or peripheral vertigo. Nausea may be caused by medications or gastroesophageal reflux disease. Nasal polyps, sinus infection, and traumatic injury to the lingual or olfactory nerves may cause appetite changes. Physical injuries to the eye including corneal abrasions, lens dislocation, retinal detachment, and optic nerve injury should be considered in the evaluation of postconcussive visual complaints. Ear abnormalities including infection, tympanic membrane rupture, and auditory nerve injury may lead to phonophobia.
GENERAL TREATMENT GUIDELINES

Treatment of physical symptoms includes treating the underlying causative or contributory conditions. Interventions targeting specific patient complaints such as sleep hygiene education, physical therapy, relaxation, and modification of the environment should be used. Moreover, medications may be used to relieve pain, enable sleep, and reduce stress. Cognitive deficits are often measurable within 30 days of mTBI but generally return to normal within the same period.1 Unfortunately, many patients continue to have subjective cognitive complaints.9,71–75 Educational and cognitive-behavioral interventions consistently improve subjective cognitive complaints.61,76–79 Behavioral symptoms may improve with psychotherapeutic and pharmacologic interventions. Treatment should be based on severity and nature of the symptom presentation. Patients with atypical symptoms or with significant suspected or confirmed comorbid illnesses may benefit from specialty referral or consultation. Finally, “red flag” conditions indicating an acute neurologic condition requiring urgent neurologic or neurosurgical intervention should prompt emergent transfer to a medical facility with an appropriate level of care.

The primary goal of pharmacologic therapies for mTBI is symptomatic improvement. At present, disease-altering therapies are not available. Drug therapy for mTBI symptoms should follow several general principles. Medications that lower the seizure threshold such as bupropion and some antipsychotic medicines should be avoided. Similarly, medications such as lithium, anticholinergic agents, benzodiazepines, and others can cause altered mental status and should also be avoided. Starting doses should be as low as possible and titrated to effect under close monitoring. By contrast, maximal tolerated dosing should be trialed before switching to a new agent to avoid undertreatment. Patients should be advised to avoid alcohol, caffeine, and herbal supplements. Limited doses of medications with significant toxicity in intentional overdoses such as tricyclic antidepressants should also be considered because suicide risk is high in brain-injured patients. Finally, patients should be monitored closely for medication interactions and toxicity.

Medication therapy for patients in the first week after injury should be reserved for the treatment of headache only. Acetaminophen is the agent of choice. NSAIDs should not be used until 48 hours after injury unless there is normal neuroimaging data for the patient. Other immediate postconcussive symptoms should not be treated as they typically resolve spontaneously within the first week of injury.

Headache

Headache is the most common symptom after mTBI, affecting more than 90% of patients. Posttraumatic headaches commonly fall into 1 of 3 categories: tension, migraine, or a combination of the 2. The evaluation of posttraumatic headache should include assessment for neurologic findings suggestive of serious intracranial abnormalities. Focal neurologic deficits should prompt additional urgent investigation with appropriate neuroimaging. A medication review for patients with symptoms lasting more than 2 weeks is also important, as rebound headaches are common in with daily acetaminophen or NSAID use. Similarly, withdrawal from caffeine or nicotine may also result in headache. Patients who state that their headache improves only with opiates should be referred to a pain or headache specialist. Headache symptoms often improve after treatment of comorbid conditions such as sleep disturbances, anxiety, and depression.

Pharmacologic treatment should be selected based on the type of headache suspected. Similar to opiate-dependent patients, those with symptoms that do not
improve within 3 months of initiating therapy should be referred to a headache or pain specialist. Episodic tension type headaches may be treated with aspirin, acetaminophen, or NSAIDs. These medications typically work best when combined with other treatment modalities such as a regular exercise program, relaxation techniques, or biofeedback. Combination medications that include caffeine or a sedative may be more effective but also have a greater likelihood of rebound headache.

Migraine treatment is divided into the prevention and management of acute episodes. Awareness and avoidance of precipitating events should be encouraged. Abortive medications for acute episodes include sumatriptan and zolmatriptan. These medications are most effective when used early in the course of the episode. Some patients with established migraines may require rescue medications to break their headache. Examples of effective rescue medications include ketorolac, butorphanol, opioids, prochlorperazine, and promethazine. Patients with migraine headaches that occur more than once a month should be placed on prophylaxis. First-line agents for prophylaxis include metoprolol and topirimate. Prophylactic agents may take as long as 3 months to become maximally effective. Finally, mixed headache types may require separate agents for treatment of the tension and migraine components.

**Disequilibrium and Vertigo**

Up to 30% of patients with mTBI complain of disturbed equilibrium or vertigo. Despite this, symptoms do not correlate with objective evidence after the first week after injury. A thorough medication review should be performed for all mTBI patients complaining of dizziness. Medications such as stimulants, benzodiazepines, tricyclics, monoamine oxidase inhibitors, tetracyclics, neuroleptics, selective serotonin reuptake inhibitors, β-blockers, and cholinesterase inhibitors may cause or exacerbate dizziness. Vestibular suppression may be useful in the acute phase but has not proven effective for persistent symptoms. Vestibular suppressants should only be used if the symptoms significantly limit the patient’s functional activities as they may result in delayed improvement. Meclizine is the first agent recommended. Scopolamine and dimenhydrinate may be used if meclizine fails. Benzodiazepines should only be used after careful consideration of their sedating and habit-forming properties. Trials should be limited to 2 weeks’ duration.

**Fatigue and Sleep Disturbances**

Another common symptom after mTBI is fatigue, which may be due to central nervous system dysfunction, sleep disturbances, or depression. Proper assessment of this symptom requires a thorough history of pre- and postinjury levels of activity. There are also several validated instruments to objectively measure fatigue. Physical causes of fatigue may also be assessed with laboratory testing including metabolic panel, a complete blood count, and thyroid function testing. Review of the patient’s medication history, alcohol, caffeine, and illicit drug use should also be performed, as all of these may result in fatigue. Before initiating medications for fatigue, conservative measures such as education of the patient, initiating an exercise program, and referring the patient for physical or cognitive-behavioral therapy should be trialed. There is limited evidence for the efficacy of stimulant treatment for fatigue after mTBI. Commonly used agents include modafanil, methylphenidate, and amantadine. These medications should only be used if symptoms have lasted more than 4 weeks, the patient does not have substance abuse issues, and addressing other factors mentioned in this section have failed to improve symptoms. Trials of these medications should last at least 3 months.
Sleep disturbances are common after mTBI. The goal of therapy is to restore a regular, unbroken night-time sleep pattern and improve the perception of sleep quality. Any drug therapy for sleep disturbances should be accompanied by education regarding good sleep hygiene. Furthermore, concomitant primary sleep disorders such as obstructive sleep apnea, restless legs syndrome, and narcolepsy should be appropriately treated. In the acute phase, short-term treatment with nonbenzodiazepine sleep medications such as zolpidem may be helpful. Prazosin may be used in patients with nightmares or agitation during sleep.

**Clinical Presentation**

The patient returns to the concussion clinic 4 weeks after his initial visit and 6 weeks after his accident. He reports that he instituted the sleep hygiene recommendations given to him on the prior visit and that his headaches, sleepiness, irritability, and concentration difficulties subsequently resolved. He has required NSAIDs with decreasing frequency and has not had a headache in the last 2 weeks. The patient is sent home with instructions to contact the clinic for a future appointment if symptoms return.

**FOLLOW-UP**

All patients require a follow-up assessment within 4 to 6 weeks of initiation of therapy. Patients can be grouped into 3 categories at this second assessment: those with complete symptom resolution, those with partial resolution, and those with no improvement or worsened symptoms. Patients whose symptoms completely resolve should be given contact information to make a future appointment if symptoms return. Patients with a partial response may benefit from augmentation or adjustment of their current therapy. Those patients whose symptoms are refractory to initial treatment should be considered to have persistent postconcussive symptoms and treated according to the following guidelines.

**Management of Persistent Postconcussive Symptoms**

This section is relevant for patients who have had an initial evaluation and failed a trial of treatment for mTBI-related symptoms. Patients with delayed presentation for mTBI symptoms should first be treated according to the section Initial Management of Postconcussive Symptoms, regardless of the interval since injury. The definitive reference is the VA/DoD Clinical Practice Guideline for Management of Concussion/mTBI.¹

Patients with persistent postconcussive symptoms often have concomitant behavioral health, psychosocial support, or compensation and litigation issues. Attention should be given to addressing these issues, as this may help mitigate symptoms refractory to initial treatment. The evaluation of the patient with persistent postconcussive symptoms should include an assessment of available support systems, a mental health history including premorbid conditions, co-occurring symptoms such as chronic pain or personality disorders, substance abuse disorders, secondary gain issues, job status, and other financial or legal difficulties.¹ Finally, all patients presenting with persistent postconcussive symptoms should be assessed for any potential danger to themselves or others.

Less than 5% of patients have persistent symptoms 1 year or more after injury.¹¹ Patients typically have more physical complaints within 4 weeks of injury after which emotional complaints predominate.⁸⁷ Once a thorough assessment has been obtained, the principal goal is to identify appropriate referrals for management of the persistent symptoms. Patients with behavioral symptoms and possible comorbid
psychiatric conditions may benefit from referral to mental health professionals. Persistent physical symptoms should be evaluated by appropriate specialists. Persistent cognitive symptoms are rare and are frequently accompanied by comorbid conditions such as mood disorders, poor physical health, poor psychosocial support, or chronic pain. In addition to addressing these comorbid conditions, these patients should be referred for neuropsychiatric evaluation to determine appropriate treatment options. Cognitive rehabilitation may be helpful for patients with persistent difficulties in memory, executive function, or attention.\textsuperscript{88–90} A social work referral is appropriate for patients with poor psychosocial support, legal difficulties, or financial problems. Although there is consistent evidence of an association between mTBI-related compensation or litigation and increased symptom reporting and poor outcome,\textsuperscript{59,91–93} there is no evidence to support a therapeutic benefit of attributing persistent symptoms to these secondary gain issues.\textsuperscript{1} Consequently, clinicians should not allow symptom exaggeration by patients seeking compensation to alter their care plans.

Given the diverse group of health professionals involved in the treatment of persistent postconcussive symptoms, a multidisciplinary team approach with the referring provider as the coordinator of care is required. A designated case manager can be very helpful for coordinating care. Typical tasks benefiting from case management includes coordination of referrals, ensuring appropriate patient and family education, participation in short- and long-term goal setting, ensuring that appropriate social service and mental health screening is performed, and coordination with the multidisciplinary team. Ongoing follow-up visits should occur regularly with goals of monitoring symptom severity, reviewing symptom impact on activities, and the effectiveness of treatments.

**Return to Play after Sports Injury**

Guidelines for returning to play in an athlete differ from general instructions for return to normal activities after mTBI in that they are designed to prevent a repeat mTBI while the patient is recovering from the initial injury. In general, the risks of suffering a second and third TBI are threefold and 8- to 9-fold greater than the risk of a first TBI, respectively.\textsuperscript{94} Furthermore, case reports suggest that athletes are at increased risk for concussion in the period immediately after their initial injury.\textsuperscript{95} Therefore, consideration of the concussion risk in a sports-specific manner is important.\textsuperscript{96} Return to play guidelines are consensus rather than evidence based. The most commonly used guidelines include those by the American Academy of Neurology (AAN) and by Cantu.\textsuperscript{97,98} Both sets of guidelines use severity of the concussion and presence of postconcussive symptoms as the criteria for return to play decision making. The Cantu guidelines allow a player to return once he or she is asymptomatic for 1 week if post-traumatic amnesia lasted less than 24 hours and the initial loss of unconsciousness was less than 5 minutes. Players with more severe symptoms at the time of their concussion should not play for 1 month, and then can return after an additional 1 week without symptoms. In players with a history of multiple concussions, consideration should be given to sitting out for the remainder of the season.

Although the consensus-based back to play guidelines referenced in this section are clinically accepted as the standard of care, they are not infallible. A recent example of a second injury despite scrupulous application of these guidelines is seen with Brian Westbrook, a professional football player in the National Football League (NFL). Mr Westbrook suffered the first concussion of his 8-year NFL career on October 26, 2009, suffering a brief LOC with associated retrograde amnesia after being tackled in a game. He was held from play for 3 weeks due to lingering headache and then
suffered a second concussion in his first game back from injury. While there was widespread speculation that he might not return that season or even that his football career was over, he did play the final 2 games of the season without further injury after being out for a total of 5 weeks. Although no guideline can prevent all adverse events, it is possible that improved guidelines could result in fewer repeat injuries. Prospectively validated, evidence-based return to play guidelines are needed.

SEQUELAE OF MTBI
Postconcussive Symptoms and Cognitive Deficits

Common postconcussive symptoms include headache, dizziness, fatigue, sleep disturbances, memory problems, balance problems, sensitivity to sound or tinnitus, concentration difficulties, and irritability. These symptoms are notably nonspecific and are associated with many other diseases. Nonetheless, several studies have reported higher rates of these symptoms in patients after mTBI than in patients with no injury or extracranial trauma without TBI. The percentage of patients who suffer from persistent postconcussive symptoms diminishes with time after injury. Less than 25% of patients are likely to have problems lasting more than 12 months after injury. Although cognitive complaints are fairly common after mTBI, measurable cognitive deficits are generally only present after severe or moderate TBI. There is little evidence of objective cognitive deficits after mTBI.

Motor, Balance, and Cranial Nerve Abnormalities

In general, objective findings after mTBI are absent. Balance problems are emerging as a promising exception to this rule. In one study of 37 mTBI patients, testing of saccades, oculomotor smooth pursuit, upper limb visuomotor function, and neuropsychologic domains was performed, and the results compared with uninjured control patients. At 1 year after injury, eye and upper limb movement, but not cognitive function, remained impaired in the mTBI patients. In a more recent study, the same group found that eye movement impairment was significantly worse in mTBI patients suffering from postconcussive syndrome relative to mTBI patients with good recovery.

Psychiatric Diagnoses

Many studies have found an association between TBI of all severities and major depressive disorder. This observed association was not likely to be explained by depression prior to injury; however, prior mood disorder may be an increased risk for TBI. While there are few studies of a relationship between mania or bipolar disorder and TBI, the existing evidence suggests that there is not a strong relationship between them. There is limited evidence supporting an association between mTBI and PTSD in military populations. In a study of 2525 soldiers returning after a 1-year deployment to Iraq, researchers identified a clear association between PTSD and mild TBI with LOC (odds ratio, 2.98; 95% confidence interval [CI], 1.70–5.24). A second cross-sectional study of 2235 Afghanistan and Iraq war veterans also found an association between PTSD and mTBI. However, 2 studies of civilian populations found no relationship between mTBI and PTSD.

Second Impact Syndrome

Second impact syndrome (SIS) is a dreaded, rare complication of mTBI that occurs after a patient suffers a second mTBI while remaining symptomatic from the first. A patient typically will suffer a head injury during play resulting in postconcussive symptoms. After returning to play while still suffering symptoms the patient sustains
a second, apparently minor head trauma, and rapidly suffers depressed mental status resulting in death or a persistent vegetative state. It is postulated that this disorder is caused by disordered cerebral autoregulation resulting from the initial TBI. The condition has mainly been reported in young men who play contact sports. The term SIS was first coined by Saunders and Harbaugh; however, a similar syndrome was previously described by Schneider.

While SIS has become firmly fixed in the minds of clinicians as an important complication of mTBI, there is some question regarding whether it is a true clinical entity. A critical review of reported cases of SIS found that most did not meet a reasonable clinical definition of SIS. Cases often lacked neuropathologic evidence of unexplained cerebral swelling. Even more problematic, most of the reported cases of precipitous neurologic collapse after a seemingly minor trauma occurred in the absence of any documented “first impact.” Of the 17 cases reviewed, only 5 were classified as “probable SIS.” Given this analysis it is reasonable to conclude that the term SIS is inaccurate. Diffuse cerebral swelling can very rarely occur after mTBI, principally in children and adolescents; however, a second mTBI is not required.

Seizures

Although there is sufficient evidence to support a causal relationship between moderate or severe TBI and the development of unprovoked seizures, the evidence is limited for an association between seizures and mTBI. In nonmilitary TBI populations, there is a 3.6-fold increase in the incidence of seizures relative to noninjured patients after TBI of all severities. After severe TBI there was a 17-fold increase in seizure incidence, which declined to 2.9-fold in moderate TBI patients. For mTBI patients with LOC or posttraumatic amnesia, the incidence of seizures was 1.5 times that of controls (95% CI 1.0–2.2). These studies were limited in that pediatric patients, who have a higher baseline incidence of seizures than adults, were not analyzed separately from adults. Posttraumatic seizure risk is greatest in the first year after injury. After 4 years, TBI patients are no longer at increased risk relative to uninjured subjects.

Dementia and Neurodegeneration

Alzheimer disease is the most common neurodegenerative disease, resulting in progressive dementia and eventual death. Familial or early-onset Alzheimer disease is caused by specific mutations and comprises approximately 10% of cases. The remaining 90% of cases are referred to as sporadic. Although the mechanisms of disease progression in sporadic Alzheimer disease are not known, it likely results from a combination of genetic and environmental factors. TBI is the strongest known environmental exposure associated with subsequent development of sporadic Alzheimer disease. A retrospective cohort study of World War II veterans with documented closed head injury demonstrated an increase risk of Alzheimer type dementia relative to nonhead-injured controls (hazard ratio 2.00, 95% CI 1.03–3.90). A meta-analysis of 7 case-control studies revealed similar results.

Dementia pugilistica, also known as chronic traumatic encephalopathy, is a neurodegenerative condition that affects athletes in sports that involve repeated head trauma such as boxing and mixed martial arts. Characteristic neuropathologic changes include cerebellar damage, cortical damage, and other scarring of the brain; substantia nigral degeneration; neurofibrillary tangles in the cerebral cortex and temporal horn areas; and abnormalities of the septum pellucidum. Autopsy of professional football players who died in their forties after developing dementia also showed neurodegenerative changes consistent with chronic traumatic encephalopathy.
Neuropsychologic deficits associated with dementia pugilistica have been found in some studies\textsuperscript{128,129} but not others.\textsuperscript{130,131}

Parkinsonism is a constellation of symptoms including tremor, rigidity, bradykinesia, and postural instability, and is caused by loss of central dopamine. Very little has been reported regarding association between TBI and parkinsonism; however, several case-control studies have shown an increased risk after mTBI with LOC or posttraumatic amnesia.\textsuperscript{132,133} The risk for the development of parkinsonism seems to increase with severity of TBI.\textsuperscript{132,134}

**SUMMARY**

mTBI is a widespread problem. Because of our limited understanding of the injury pathophysiology, the diagnosis of mTBI is based entirely on clinical symptoms, and often occurs in the absence of objective findings. The central feature of mTBI is a transiently altered state of consciousness after a traumatic injury to the head. The priority of emergency care is to identify potentially life-threatening intracranial injuries through the judicious application of appropriate imaging studies and neurosurgical consultation. Although post-mTBI symptoms quickly and completely resolve in the vast majority of cases, a significant number of patients will complain of lasting problems. Postconcussive complaints tend to be interrelated, and relief of one may have beneficial effects on others. Although the evidence is not definitive, longer-term sequelae of mTBI may include seizure disorders and neurodegeneration. Recognizing the potentially life-changing aspects of mTBI should be an important priority for the emergency physician because simple, early interventions, such as education regarding the expected positive outcome from the injury and prompt treatment, can prevent chronic symptoms from occurring.

**REFERENCES**


