Pathophysiological to Clinical Aspects of Head Injury: Narrative Literature Review

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ABSTRACT

Head injury is a change in brain function or brain pathology, caused by external forces on the head. Changes in brain function consist of any period of loss or loss of consciousness, anterograde or retrograde amnesia, neurological deficits, or mental changes following a head injury.¹,² A study reported that the highest incidence of head injury was in the Americas, which was 1299 cases per 100,000 population, followed by Europe, which was 1012 cases per 100,000 population. The lowest incidence of head injury occurred on the African continent, which was 801 cases per 100,000 population. Based on these data, the overall incidence of head injuries in the world is 939 cases per 100,000 population with most of them being mild head injuries (55.9 million cases per year) and 5.28 million people experiencing severe head injuries.

INTRODUCTION

Head injury is a change in brain function or brain pathology, caused by external forces on the head. Changes in brain function consist of any period of loss or loss of consciousness, anterograde or retrograde amnesia, neurological deficits, or mental changes following a head injury.¹,² A study reported the highest incidence of head injuries was in the Americas, namely 1299 cases per 100,000 population, followed by Europe, which was 1012 cases per 100,000 population. The lowest incidence of head injury occurred on the African continent, which was 801 cases per 100,000 population. Based on these data, the overall incidence of head injuries in the world is 939 cases per 100,000 population with most of them being mild head injuries (55.9 million cases per year) and 5.28 million people experiencing severe head injuries. At the regional level, the highest incidence of head injury in the world is in Southeast Asia, which is 18.3 million cases.³⁻⁵ In Indonesia, there have been several reports of head injuries, during June-December 2018 there were 118 cases of head injuries recorded at H. Adam Malik Hospital, North Sumatra with patient ages ranging from 18 years to 35 years and dominated by the male gender. In this study, most cases were epidural bleeds. Based on medical record data at the Central General Hospital Dr. M. Djamil Padang there were 356 head injury cases in 2017 and 505 cases in 2018.⁵

Classification of head injury

Classification of head injuries was made based on
clinical severity and was assessed using the Glasgow Coma Scale (GCS). The Glasgow coma scale consists of 3 components that allow a rapid assessment of the severity of the head injury: eye, verbal, and motor. Scores of 14-15, 9-13, and 3-8 were classified as mild head injury, moderate head injury, and severe head injury, respectively. Table 1 displays the scores on the GCS.6,7

Table 1 Glasgow coma scale

| Response | Score |
|----------|-------|
|          | 1     | 2     | 3     | 4     | 5     | 6     |
| Eyes     |       |       |       |       |       |       |
| Does not open eyes | Opening the eyes with pain stimulation | Opens eyes with sound | Opens eyes spontaneously | - | - |
| Verbals  |       |       |       |       |       |       |
| Does not make sound | Unclear voice | Inappropriate words | Confused, disoriented | Good orientation, normal | - |
| Motor    |       |       |       |       |       |       |
| No movement | Extension due to painful stimulation | Abnormal flexion to painful stimulation | Flexion/avoids pain stimulation | Localizing pain stimulation | Following orders |

Other classifications of head injuries are classified according to morphology:

a. Fracture of the skull: cranium (depressive and non-depressive fractures) and the cranial base.
b. Intracranial lesions: focal (epidural, subdural, and intracerebral/contusion), diffuse (concussion, ischemic injury, diffuse axon injury).8

Main causes of head injuries are traffic accidents, falls, and assaults. Based on the mechanism, head trauma is classified as blunt (a most common mechanism), penetrating (most fatal) and explosive. The most serious injuries result from motor vehicle collisions and falls.9,10

Pathophysiology of head injuries

Head injuries can be classified into primary and secondary injuries. Primary injuries include injuries due to shifting of the brain due to direct impact, rapid acceleration-deceleration, or penetration. These injuries can lead to contusions, hematomas, or axonal injuries. The secondary injury occurs after primary injury in the form of the release of biochemical compounds such as glutamate which causes further damage to mitochondria and cell death and necrosis. A secondary head injury causes systemic hypotension, hypoxia, and increased intracranial pressure leading to brain herniation.11-13

The initial phase

The initial phase occurs immediately after trauma due to impaired blood flow and ischemic conditions leading to conditions of decreased adenosine triphosphate, oxygenation, glucose consumption and distribution, depolarization of calcium ion channels, increased lactate, and neuronal cell death. After neuronal cell death, glutamate exits the damaged presynaptic vesicles causing excitotoxicity. Glutamate binds to N-methyl-D-aspartate receptors, increases Ca2þ and Naþ in cells, and activates enzymes responsible for tissue damage. Excessive accumulation of calcium in neurons also stimulates nitric oxide and causes oxidative stress. This exacerbates cell death.14 Necrotic areas of nerve cells and glial cells are concentrated in areas where the blood supply is impaired, which can occur due to epidural hemorrhage, subdural hemorrhage, and intracerebral hemorrhage. Secondary contusions can
occur in opposing brain tissues because of the coup and counter-coup. Cognitive deficits, behavioral changes, and hemiparesis depend on the severity of the injury. In contrast to focal injury, the main mechanism of diffuse brain injury is rapid acceleration and deceleration forces that cause shear and strain injuries to cerebral brain tissue. This results in injury to axons, oligodendrocytes, and blood vessels, which leads to cerebral edema and ischemic brain damage. The degree of axonal injury and neuronal degeneration determines the severity of the head injury.

**Intermediate phase**

Abnormal function caused by mechanical damage and neurotrauma results in an inflammatory process. This inflammation can increase brain injury and activate microglia, which further prolongs neuroinflammation. An increase in proinflammatory cytokines and chemokines occurs approximately 1 hour after nerve injury. One of the inflammatory cytokines, IL-1b, acts on astrocytes, activates the intracellular ERK pathway, and releases matrix metalloproteinase-9. Matrix metalloproteinases will erode the extracellular matrix, impair the integrity and function of the BBB, and induce chronic neuroinflammation. The brain is normally protected from immune cells and pathogens due to the presence of the Blood-Brain Barrier (BBB). However, damage to the BBB following head injury results in leakage of prostaglandins, nitric oxide, cytokines, and inflammatory mediators into brain tissue. After the primary injury, the inflammatory response is activated by the invasion of monocytes, neutrophils, and lymphocytes across the BBB.

**Final phase**

The inflammatory process is to remove pathogens from the site of injury, regenerate damaged cells and improve nerve cell function. However, the recovery process leaves some sequelae such as seizures and epilepsy. The reduced expression of Kv.4.2 increases the excitability of the nerves thought to mediate

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Figure 1. Schematic of the pathophysiology of head injury
seizures. In addition, injury-induced epilepsy is caused by activation of the trkBERK1/2-CREB/Elk-1 pathway and GAP-43 expression.\textsuperscript{14}

**Diagnosis of head injury**

**Anamnesis**

Anamnesis is asked the mechanism of injury, symptoms and findings of physical examination associated with head injury.\textsuperscript{16} The mechanism of injury was associated with immediate intervention which included observation, admission to hospital or ICU, and/or neurosurgical intervention. The mechanisms of injury that most often result in severe head injuries include pedestrians being hit by vehicles, passengers being thrown from motorized vehicles, and people falling from a height of more than 1 meter. Motorcycle collisions, not wearing a helmet, and chronic alcoholism are associated with an increased risk of severe intracranial injury even with mild initial symptoms.\textsuperscript{16,17} The presence of vomiting, especially more than two episodes, has a higher probability of developing a severe head injury. Posttraumatic seizures are also associated with a higher degree of severity.\textsuperscript{16,17}

**Physical examination**

In patients with mild and moderate head injuries, physical examination findings can help predict serious injury. Focal neurologic deficits are associated with a severe head injury. Signs of skull fracture (open, depressed, or base) such as hemotympanum, otorrhea, peri-orbital ecchymosis, or skull base fracture identified by postauricular ecchymosis (*Battle sign*) are also associated with a severe head injury. A severe injury in an unconscious drunk patient may be associated with alcohol intoxication. In mild head injury, initial GCS 13, GCS worsening, and GCS $<$14 2 hours post-injury were all associated with more severe intracranial injury.\textsuperscript{16}

**Imaging**

Radiological examination plays an important role in identifying patients with head injuries. Common imaging techniques include a CT scan of the head and an MRI. Imaging will help differentiate patients who require immediate neurosurgical intervention or who may be discharged from patients under observation. When there is a clinical indication for imaging, a non-contrast head CT scan is the first choice. MRI is superior in identifying small, focal traumatic lesions.\textsuperscript{17} Non-contrast head CT scan is recommended as the imaging choice and can identify subdural hemorrhage, epidural hemorrhage, subarachnoid hemorrhage, intracerebral hemorrhage, cerebral contusion, skull fracture, pneumocephalus, and cerebral edema. Indications for CT scanning are divided into two criteria, namely moderate risk of intracranial injury and high risk of intracranial injury.\textsuperscript{16}

|                          | Moderate risk                                                                 | High risk                                                   |
|--------------------------|-------------------------------------------------------------------------------|-------------------------------------------------------------|
|                          | Loss of consciousness after injury                                           | Unclear decrease in consciousness                           |
|                          | Drug intoxication                                                            | Focal neurologic deficit                                    |
|                          | Post-traumatic seizures                                                      | Depressive fracture                                         |
|                          | The mechanism of trauma is unclear                                           | Translucent skull injury                                    |
|                          | Age less than 2 years                                                        |                                                             |
|                          | Vomiting                                                                     |                                                             |
|                          | Amnesia after head injury                                                    |                                                             |
|                          | Multiple trauma                                                              |                                                             |
|                          | Signs of base fracture                                                       |                                                             |
|                          | Severe facial injury                                                         |                                                             |
|                          | Suspicion of depression or fractured skull                                   |                                                             |
|                          | With GCS $\leq$14                                                            |                                                             |
Management of head injury

Prehospital management

Care for head-injured patients should begin at the site of injury to maintain the airway and maintain adequate circulation and ventilation. Patients with moderate to severe head injuries should be immediately sent to a medical center with a neurosurgical facility. The initial goal of management was the prevention of hypoxia and hypotension because both of these doubled mortality.\(^\text{18}\)

- Airway management/oxygenation
  - Prevention, identification and therapy of hypoxia (\(O_2\) saturation < 90% and/or cyanosis).
  - Airway maneuver.
  - Ventilation with NRM.
  - Endotracheal intubation is indicated.

- Ventilation Management
  - Intubated and mechanically ventilated patient: if available, target \(\text{PaCO}_2\) of 40 mmHg.
  - Prophylactic hyperventilation in the prevention of ICP is not recommended.

- Management of blood pressure
  - Hypotension: at systolic blood pressure < 90 mmHg, fluid resuscitation was performed with an initial bolus of 1 liter of normal saline or RL with a target TDS of 90 mmHg.
  - Hypertension: therapy for acute hypertension is not recommended in cases of head injury. However, IV fluids are restricted at a minimal rate if the TDS is 140 mmHg.\(^\text{18}\)

Medical Interventions

Head elevation

Elevation in head injuries generally has a rapid effect on lowering intracranial pressure. Intracranial pressure (ICP) is reduced by displacement of cerebrospinal fluid from the intracranial compartment and promoting venous outflow. Although the mean carotid pressure is reduced during head elevation, ICP is reduced and cerebral blood flow is not affected.\(^\text{14}\)

Hyperventilation

Hyperventilation lowers ICP by reducing the intraarterial partial pressure of carbon dioxide (\(\text{PaCO}_2\)) and causing vasoconstriction. However, this action ultimately causes a decrease in cerebral blood volume. Prophylactic hyperventilation is not recommended, because vasoconstriction reduces blood flow to the brain. The use of hyperventilation in the setting of severe head injury is usually only used for a short time during acute neurologic deterioration. Hyperventilate to \(\text{PaCO}_2\) 25 mm Hg to lower ICP.\(^\text{19-21}\)

Seizure prophylaxis

Current head injury guidelines state that 1 week of prophylactic antiepileptic use is acceptable to help prevent early seizures. Currently, the recommended drug is phenytoin. However, there has been no proven benefit in the long-term prevention of seizures after the head injury, so prophylaxis was discontinued after 7 days.\(^\text{22-24}\)

Hyperosmolar Therapy

Hyperosmolar therapy in head injuries can be given as a bolus or infusion. Administration of mannitol is recommended in euvolemic severe head injury patients.\(^\text{22}\) Mannitol is used to reduce the increase in ICP. The most common preparation is a 20% solution (20 g of mannitol per 100 ml of solution). Strong indications for administering mannitol in euvolemic patients are pupillary dilation, hemiparesis, and decreased consciousness.\(^\text{22}\)

Medications that cause coma status

The patient is brought into a coma by infusion of a benzodiazepine or barbiturate. The administration is carried out with a continuous electroencephalogram. This therapy serves to reduce the metabolic needs of
the brain. This administration is only recommended in cases of severe refractory increase in ICP after medical therapy and maximal reduction in ICP.22,23

**Therapeutic hypothermia**

This therapy can reduce oxidative injury due to the effect of decreasing the metabolic demands of the brain but at risk of changes in blood sugar, platelet count, and coagulation factors. This therapy is performed on severe head injuries.24,23

**Surgical Intervention**

Surgical intervention is generally required when there is a mass effect of either an epidural hemorrhage, subdural hemorrhage, intracerebral hemorrhage, or contusion with significant blood volume. The principal management of epidural hemorrhage is craniotomy of the traumatized area, with the evacuation of the hematoma and cauterization of the torn vessel, often the middle meningeal artery. Acute subdural hemorrhage is usually associated with more severe brain injury.24,25

**Conclusion**

Head injury is a change in brain function or brain pathology, caused by external forces on the head. Changes in brain function consist of any period of loss or loss of consciousness, anterograde or retrograde amnesia, neurological deficits, or mental changes following a head injury.

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