			<ul> <li>the acutely injured brain is vulnerable to damage from systemic hypotension, cerebral hypoperfusion, hypercarbia, hypoxemia &amp; elevated ICP</li> <li>care of the TBI victim should begin with evaluating and securing a patent airway and restoring normal breathing and circulation</li> </ul>
			- early intubation is probably of benefit; however, the benefits of prehospital
			intubation have yet to be demonstrated in a randomised controlled trial
(i) avoidance of hypoxaemia: Fi00 should be distributed maintain portrait except a			the patient should be sedated and pharmacologically paralysed before intubation     because initiation of the orophapma causes transient bunchesions (CP)
- FIO2 should be traded maintain normal oxygen saturation - High levels of PEEP may increase ICP; however, clinical studies have shown that			- supplemental oxygen should be provided before intubation
the use of PEEP of up to 15cmH20 in patients with ARDS does not increase ICP		nrehosnital	- therapeutic hyperventilation is inadvisable unless neurological deterioration is clearly evident during
(ii) maintenance of normocarbia		care	evaluation and transport; aggressive hyperventilation can cause cerebral ischaemia via vasoconstriction
<ul> <li>maintaining an arterial PCO2 of approximately 35 is advised to avoid the cerebral vasoconstriction associated with aggressive hyperventilation</li> </ul>			<ul> <li>- rapid fluid resuscitation and restoration of normal blood pressure are critical in the prenospital setting because hypotension has been associated with doubling of mortality after severe traumatic brain injury.</li> </ul>
(iii) avoidance of nypotension and nypovolaemia - hypotension should be agressively treated with normovolaemia achieved by infusing			- hypovolaemia is the likely mechanism and therefore normal saline or Hartmanns should be infused as rapidly as possible
normal saline [human albumin is associated with increased morbidity and mortality in severe TBI)			- although preclinical studies suggested hypertonic saline may be more effective for rapid volume resuscitation in head injured patients,
- hypotension which is refractory to volume replacement should be treated with vasopressors or inotropes			several clinical trais have failed to demonstrate a benefit
- anaemia should be treated; however, the precise level at which transfusion should occur is not clear		1	<ul> <li>- an patients with a ostracting injury (including need injury) should be related as in they have a cervical spine injury estimate about the isotracting in to a lowal, be 11 forume on the ostraction as in they have a cervical spine injury</li> </ul>
(iv) maintenance of CPP - some advocate use of induced hypertension to maintain a CPP above 70mmHo: however a randomised trial of patients with TRI comparing a			- patients should be transported to a rever for in tradina centre (ensuing the immediate availability of neurostroical care when the patient arrives)
group whose CPP was kept above 70mmHg to a group whose CPP was allowed to drift to 60mmHg showed no difference in outcome at six months			
between the two groups and more use of vasopressors and a higher incidence of ARDs in the group whose CPP was maintained above 70mmHg			- upon arrival at the trauma centre. the emergency medical personnel should report their prehospital assessment and management including
<ul> <li>others have round that or an ussue PO2 in patients with 1bi does not rail until the CPP dops below ourning</li> <li>based on the above findings the current recommendation is to maintain a CPP above 60mmHg</li> </ul>			mechanism of injury, stabilising maneouvres, medications given, initial vital signs and GCS and haemodynamic stability during transport
(v) avoidance of intracranial hypertension			<ul> <li>- immediate management should proceed according to the principles of the ATLS</li> </ul>
- intracranial hypertension is defined as a sustained ICP greater than 20mmHg			protocol which is designed to identify and treat immediately line threatening injuries
<ul> <li>several clinical studies have found that persistent infracranial hypertension is associated with significantly worse morbity and mortality based on the association with worse outdowns and the provise that intercential hypertension can compressive that</li> </ul>			<ul> <li>The alrway should be reassessed and the need to initioate the patient should be considered; for patients intubated in the read the proper position of the ET tube is verified both clinically and radiologically as well as with end tidal CO2</li> </ul>
<ul> <li>based on the association with worse buckets and the perfusion and induce ischaema to an experimentation of intracranial hypertension is almost universally endorsed</li> </ul>			- when the airway is secured adequate oxygenation is confirmed using percutaneous oxygen saturation and arterial blood gas analysis
- always considere physiological causes of raised ICP including seizures, fever, jugular outflow obstruction and agitation			- two large bore iv catheters are inserted to provide sufficient venous access for high volume
(vi) sedation:		emergency	fluid resuscitation and isotonic crystalloid should be continued to replace volume loss
<ul> <li>increasing sedation may lead to rapid control of intracranial hypertension particularly in a patient who is posturing or agitated the maior disadvantages of sedation are that the ability to determine an accurate GCS</li> </ul>		department	I - If e threatening injuries such as tension pneumothorax, cardiac tamponade and overt
is lost and sedative agents often induce hypotension		Care	- a brief peurological evaluation is performed including assessment of the GCS punits & extent of extremity movements
(vii) venting of CSF			<ul> <li>careful inspection of the head should reveal haemotympanum, periorbital or mastoid ecchymosis and CSF rhinorrhoea or otorrhoea</li> </ul>
- in a patient with an external ventricular drain intermittent or continuous venting of CSF is useful intermittent vertication of entry of the section of entry of the section of the sect		1	- oxygen saturation is continually monitored and blood pressure frequently or continuously measured during the primary examination
- intermittent venting has the advantage of allowing reliable measurement of ICP		1	<ul> <li>a foley catheter is placed to help monitor the fluid status and an orogastric tube is inserted to decompress the stomach blood generinsers are abbiased and anshired for always called blood for the storage arbitration and the storage and the</li></ul>
I. mannitol		1	Induce specimients are obtained and analysed for glucose, electrolytes, full blood count, coags, and cross match; serum toxicology may be appropriate and women of childbearing age should undergo a breanancy test
- intermittent boluses of mannitol (0.25-1g/kg every 3-4 hours as needed) lowers ICP & increases CBF by expanding		1	- a CT brain should be performed unless haemodynamic instability necessitates an emergent laparotomy or thoracotomy; in these circumstances.
intravascular volume and reducing blood viscosity within a few minutes of administration; its duration of action is 3-5 hours			diagnostic burrholes may be appropriate in theatre if the patient has lateralising neurological deficits particularly a unilateral fixed and dilated pupil
<ul> <li>- continuous intrasion on maintenis tess destable train docess declares de tate is tess intery to read to extravasation of drug into the brain causing a reverse osmotic gradient and increased oedema</li> </ul>			
- the serum osmolarity and sodium level should be monitored frequently during mannitol administration to minimise the risk of renal			- critical to detemining the severity of the brain injury and the appropriate treatment are CT findings
failure from ATN; the drug should be discontinued if the serum sodium exceeds 160mmol/L or the osmolarity exceeds 320mosm			combined with a reliable post-resuscitation GCS score and assessment of pupil size and reactivity
<ul> <li>a. hyperionic same</li> <li>- 3% saline can be administered as an osmotherapy and titrated to serum sodium</li> </ul>			<ul> <li>other determining factors include the size and location of the haematoma, the presence and and other disputcies of the second the results of powerland examination</li> </ul>
- principle advantages of hypertonic saline in this setting are:			- neurological deterioration success enlargement of the haematoma and a new CT scan should be performed promptly
(i) rapid effect which peaks in 10 minutes and wanes after 1 hour			- haematomas less than 10mm thick that cause midline shift of less than 5mm can usually be observed especially if they do not involve the middle
(ii) end point for merapy is serum socialm which is 145-155 and easily monitored through ABGS (iii) there is less potential for hypovolaemia than with mannitol	medical managem	ent	cranial fossa; a haematoma that compresses the termporal lobe is particularly ominous and can rapidly can uncal herniation so that such lesions
(iv) there may be a better effect on CBF for a given reduction in ICP	treatment of trauma	tic	warrant a lower threshold for evacuation - if a clot is small enough not to require evacuation it should be monitored with frequent CT scans over the first several days after injury. Enlaming
(v) HS is inexpensive	brain inju	ry	middle fossa hæmatomas large enough to cause hernlation do not always lead to a rise in ICP
<ul> <li>virial operation operation in modulating the minimization response</li> <li>principle disadvantages of hypertonic saline in this setting are:</li> </ul>	Created	by definitive	- patients with small or deep seated contusions without mass effect can be managed non-operatively initially. Contusions should be followed serially
(i) need for central access	Paul You	ng treatment	With CT scanning as there is a 20-30% chance of significant enlargement in the first 24-48 hours a temporal contribution can exist of uncel homical branching without a significant
(iii) hypokalaemia & hypercholoraemic acidosis (iii) kapt of outeness data	11/11/07	]	- a temporar contestor can emarge to the point of ancal remained without a significant rise in ICP: thus the threshold for exacution of these lesions should be lower
(iii) lack of outcome data (iv) increase in circulating volume and risk of CCF			I - unilateral frontal or temporal lobectomies are usually well tolerated, do not cause
(v) coagulopathy - HS may affect APTT & INR as well as platelet aggregation	/		measurable neurological deficit and provide space for the brain to swell
(vi) rapid changes in serum sodium concentrations may result in seizures and encephalopathy (vii) some suggest that HS affects promat brain more than injurad brain which theoretically warsans benefation			penetrating injuries: - high velocity projectiles such as bullets generally cause massive destruction of brain tissue, severe brain swelling & often death
(viii) hypervaliation			- low veolicity massiles such as knives or arrives do not cause the massive brain injury associated with bullet wounds and usually
- the use of hyperventilation to lower ICP is controversial because of its association			only the tissue in the immediate path of the missile is damaged
with cerebral vasoconstriction and potential for worsening of brain ischaemia			<ul> <li>- prophytactic antibiotics should be administered because the missile usually carries skin and hair into the brain</li> </ul>
injury by casing and increase in extracellular lactate and glutamate levels			
- its only role is probably in the patient in whom other therapies have failed in whom emergent surgery is planned to control ICP			general:
(ix) paralysis and cooling			(i) continual end tidal CO2 and frequent analyses of ABGs allow early detection of deteriorating ventilatory status
<ul> <li>paralysis may nelp control ICP where other measures have failed; however, it is associated with an increased risk of neuronia and critical care myoneuronathies</li> </ul>			(ii) oxygen saturation should be continuously monitored with pulse oximetry
- therapeutic hypothermia to 32-34 degrees has been studied in the 1st 24-48 hours after TBI. While it has not been convincingly demonstrated to			(iii) blood pressure should be investigation of a second structure of the seco
improve outcome, it does consistently reduce ICP. In patients who are cool at arrival to hospital it appears to confer benefit in subgroup analyses			(iv) urine output is continuously monitoring via an indwelling catheter
(x) barbiturate coma - barbiturates are thought to be effective through their shilling to reduce combral metabolic rate and blood flow		1	/ icp monitoring
<ul> <li>- balanciates are indigined by encoding the second many of earlies and the second many and block many of the second many and the seco</li></ul>			<ul> <li>continuous icp monitoring should be mandatory for all patient with severe 161 and abnormal</li> <li>CT findings because intracrenial hypertension develops in 55-63% of such patients. Monitoring of ICP and MAP allows</li> </ul>
(xi) avoidance of hyperthermia			calculation of CPP which may a more important value than MAP or ICP.
- there is a log increase in neuronal death in ischaemic brain regions for every degree above 39 for at least 24 hours after brain injury;			- the gold standard for icp monitoring is an ventricular catheter which has a
aggressive treatment or sources or tever should be pursued and tever should be treated. Whether aggressive cooling and paralysis to achieve normothermia is warranted is unknown		physiologic	al (i) ventricular provenual advantages over atternative systems:
(xi) seizure prophylaxis		monitoring	<ul> <li>(ii) subdural, extradural or subarachnoid catheters are more prone to occlusion</li> </ul>
- contusions and subdural haematomas are well known to cause generalised seizures and anticonvulsant			(iii) ventriculostomies can be rezeroed after insertion
prophylaxis is therefore recommended for patients with these lesions (usually phenytoin is given)			(iv) ventriculostomies allow drainage of CSF to treat intractantial hypertension - the overall complication rate of EVDs is 7 7% with infection occurring in 6.3% & baemorthage occurring in 1.4% [some
<ul> <li>a proscepticitive randomised trial mas found no benefit in continuing seizure prophytaxis beyond / days</li> <li>seizures may not be evident in patients who are paralysed therefore seizure prophytaxis should be</li> </ul>			studies indicate that infection rate increases markedly after catheters have been in situ for 5 days]
continued in these patients and continuous EEG monitoring should be considered			- alternatives to ventriculostomy include devices that contain a pressure sensing transduced within the tip of the catheter (eg Codmans).
(xii) DVT prophylaxis			Auroniteges att. (i) they provide relatively accurate measurements of global ICP,
<ul> <li>patients with 1BI, particularly those or are comatose or have associated injuries such as pelyic or long hone fractures are at high risk of thrombourbuils events</li> </ul>			(ii) they are easier to insert than EVDs
<ul> <li>- patients should be devine devine and the prophylaxis including the use of sequential calf compression devices</li> </ul>			(iii) they may cause fewer complications than EVDs
- early use of both heparin and enoxaparin (within 2 to 3 days of injury) has been demonstrated to be		1	<ul> <li>- usadvantages or mese systems are trutturery can only be canonated at insertion and measurement drift may be significant over the course of a few days</li> </ul>
sate in clinical trials and has not been demonstrated to cause or worsen intracranial haemorrhage after TBI	1	1	
- malnutrition is common after TBI with metabolic expenditure increasing significantly; early enteral nutrition should be instituted	1	1	(i) evacuation of mass lesions
	1	1	- the first response to a rise in ICP should be to repeat a CT brain to exclude a
- rehabilition of TBI patients should begin in the ICU within the first few days of iniury with passive	1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1		new or worsening mass lesion that might be amenable to surgical intervention
range of movement exercise and mobilisation to prevent deep vein thrombosis	renabilitation	surgical	(II) decompressive craniectomy - evidence surrounding decompressive craniectomy is contradictory
		treatment	- while one study of patients with severe TBI demonstrated that 6 month outcomes were similar among patient given large decompressive
		<u>۱</u>	cranicactomies than among patients that did not despite lower GCS & more severe radiologoical abnormalities in the craniectomy group another advalues from the did did patience unclude LCD. CDP are producible to a producible to the craniectomy control of the craniectomy of the c
		\	suby his round that is do not intribute for, or or initiality rates - another study suggested that for young patients decompressive temporal lobectomy, improves outcome
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