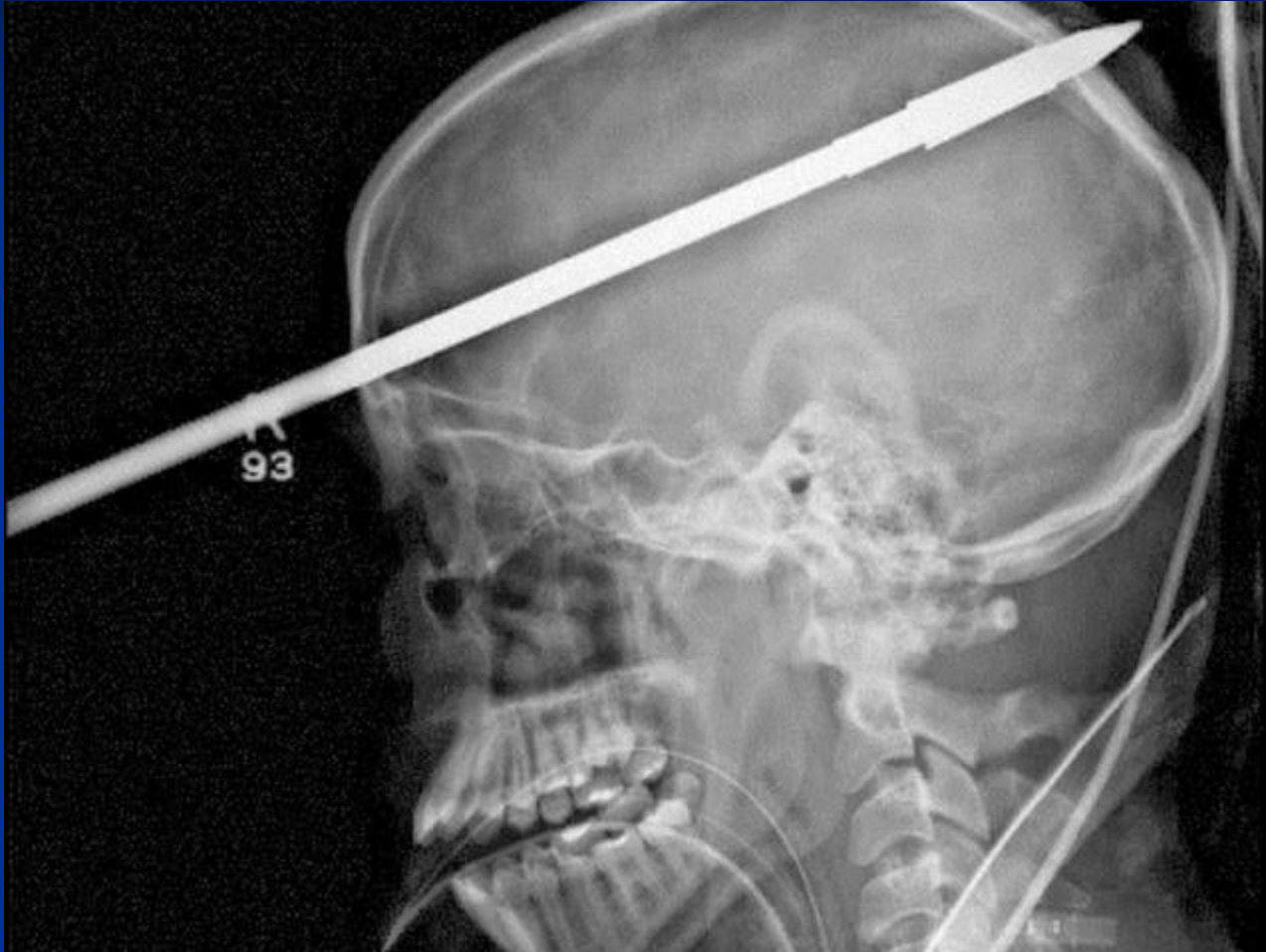


# Neurotraumatology

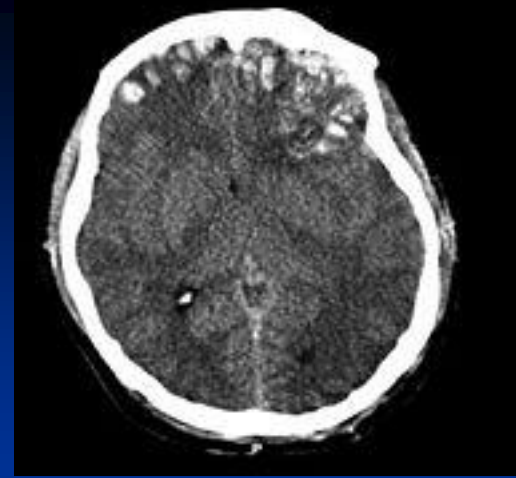


SE Dept. of Neurosurgery and Neurointervention, Budapest

# Causes of traumatic brain injury (TBI)

- Auto, motorcycle or bicycle crashes
- Falls
- Violence: gun shots, abuse
- Explosive blasts
  
- Classification: Scalp injury, skull injury, brain injury

# Traumatic brain injury



- **Primary:** direct force, skull fracture, contusion, haematomas, laceration, nerve damage
- **Secondary(consequence):** brain swelling (oedema), increased pressure inside of the skull ( high intracranial pressure), epilepsy, haematomas
- The aim: management of secondary brain injury to decrease of mortality  
(the primary has already occurred)

# Traumatic brain injury

- Healthy , young patients,  
The major cause of death and disability worldwide under 45 y
- Serious consequences (personal and social)
- Importance of primary injury supply





# Statistics on TBI

14 000/year patients hospitalized in Hungary cause of TBI

## Symptoms

-71,3% mild

-19,4% moderate

-9,4% severe



55% of patients die with severe TBI

40% of survivors have lasting damage or vegetative state.

60%-of survivors have mild symptoms or asymptomatic.

# Symptoms of TBI

- Light-headedness, dizziness
- Feeling tired, with no motivation
- Urge to vomit
- Changes in sleep patterns (more sleeping)
- Mood changes (sad or angry for no reason)
- Increased sensitivity to lights, sounds,
- Loss of sense of smell or taste, ringing in the ear

# Prehospital treatment for TBI

- Checking of vital parameters
- **Oxygenation and blood pressure**



***Correct CPP (brain perfusion).*** management of secondary brain injury to decrease of mortality

- Inspektion – cranium
- Palpation – cranium
- Neurological examination
- „Glasgow Coma Scale” determining
- Known of anamnesis
- Polytrauma (80% the probability of head injury)!
- Always need to exclusion the injury of cervical spine

# Glasgow Coma Scale (GCS)

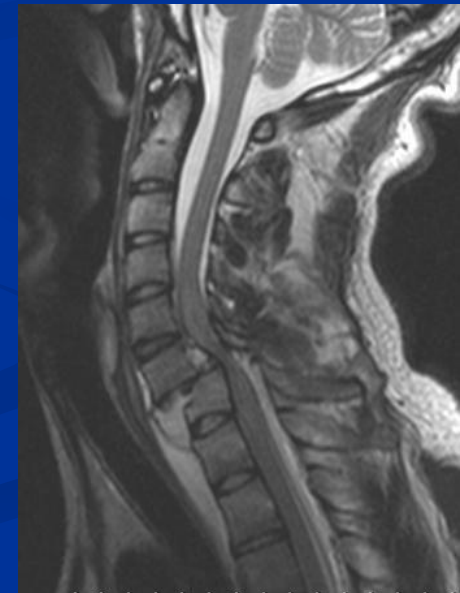
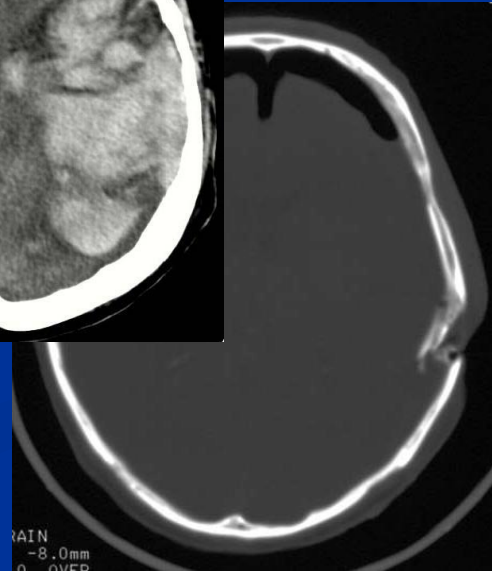
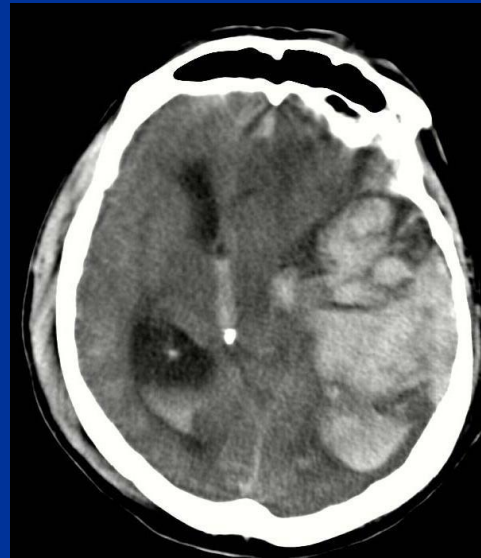
Glasgow Coma Scale		
Response	Scale	Score
Eye Opening Response	Eyes open spontaneously	4 Points
	Eyes open to verbal command, speech, or shout	3 Points
	Eyes open to pain (not applied to face)	2 Points
	No eye opening	1 Point
Verbal Response	Oriented	5 Points
	Confused conversation, but able to answer questions	4 Points
	Inappropriate responses, words discernible	3 Points
	Incomprehensible sounds or speech	2 Points
	No verbal response	1 Point
Motor Response	Obeys commands for movement	6 Points
	Purposeful movement to painful stimulus	5 Points
	Withdraws from pain	4 Points
	Abnormal (spastic) flexion, decorticate posture	3 Points
	Extensor (rigid) response, decerebrate posture	2 Points
	No motor response	1 Point
<b>Minor Brain Injury = 13-15 points; Moderate Brain Injury = 9-12 points; Severe Brain Injury = 3-8 points</b>		

# Radiological methods

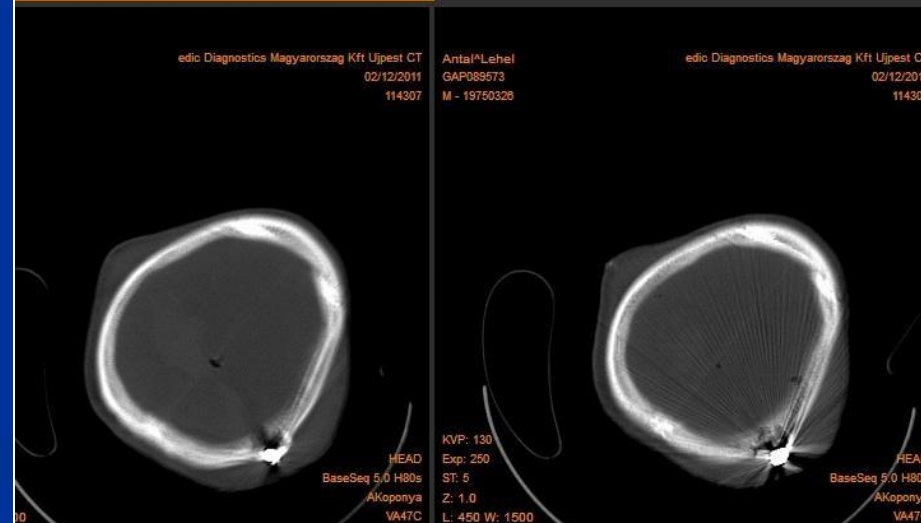
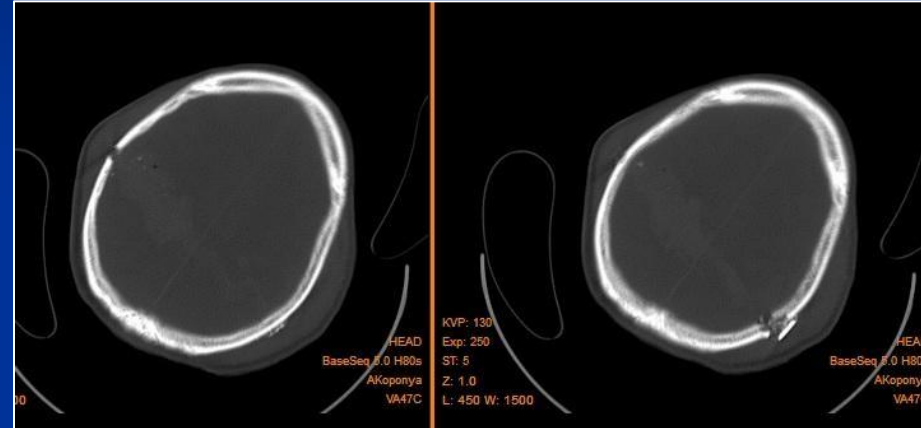
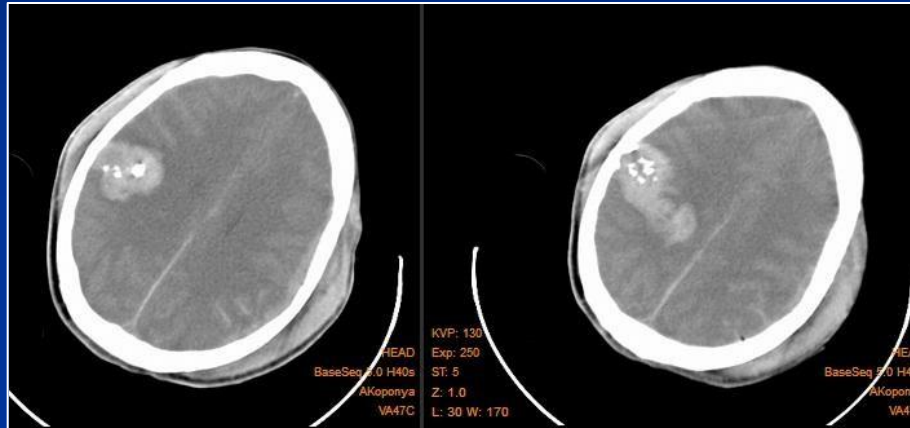
X-ray

CT

MRI (rare)

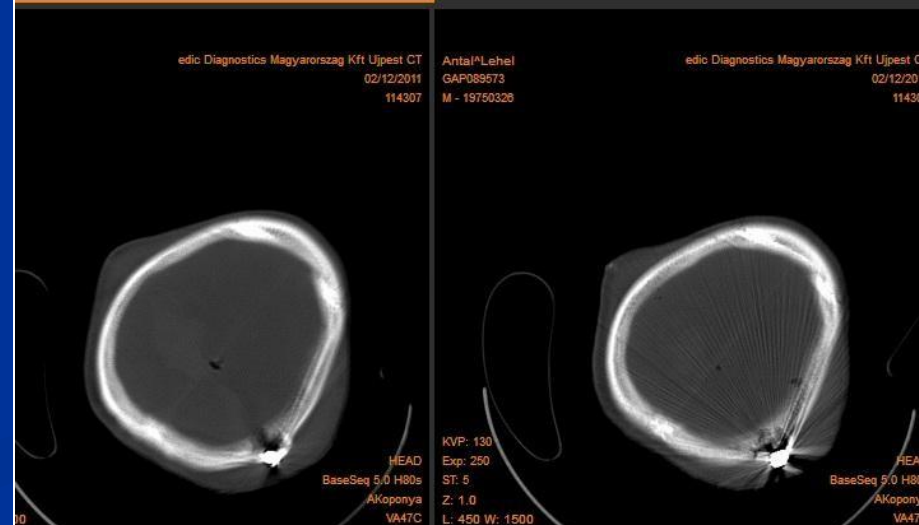
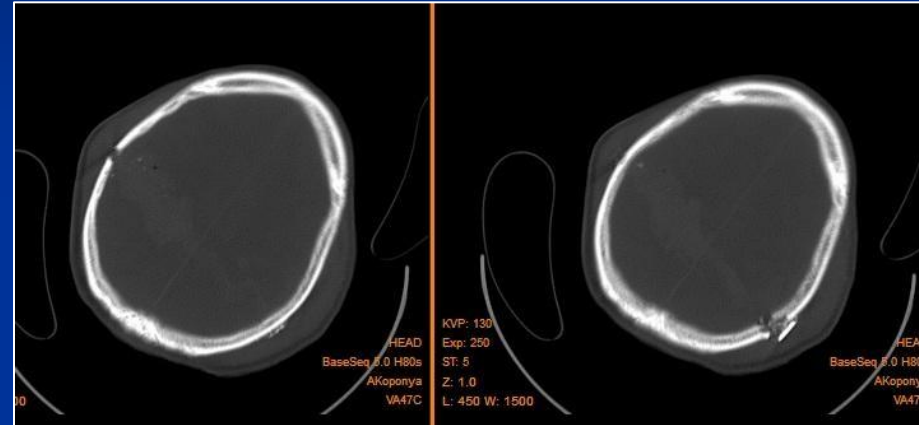
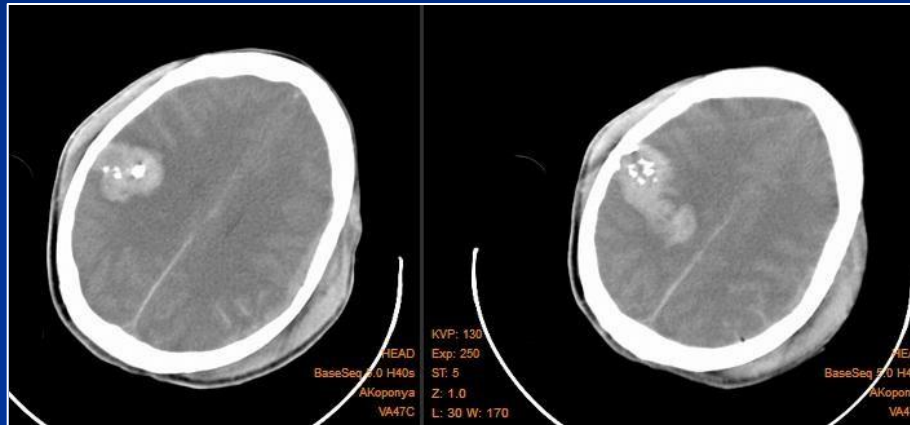


# CT examination



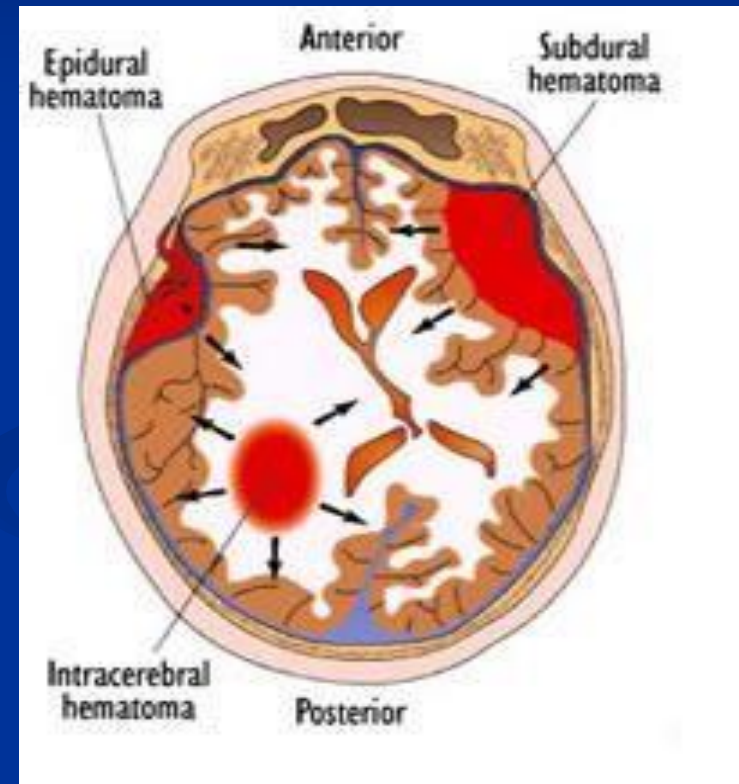


# The bullet comes from the.?..side



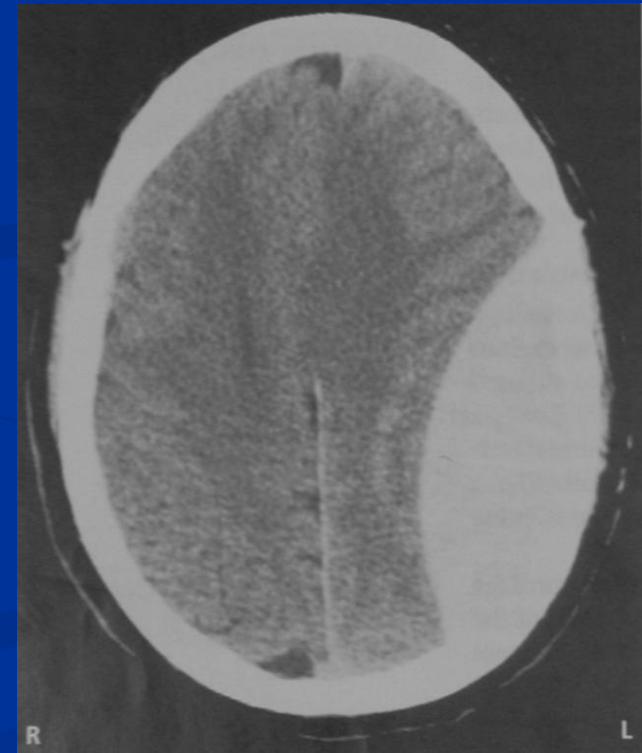
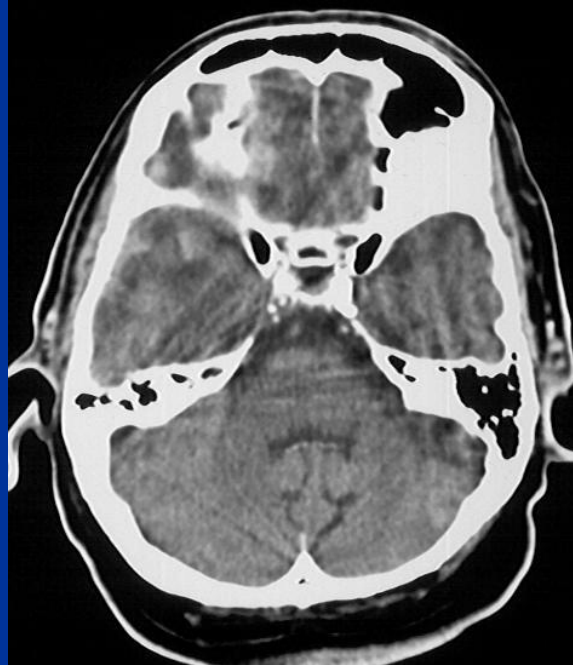
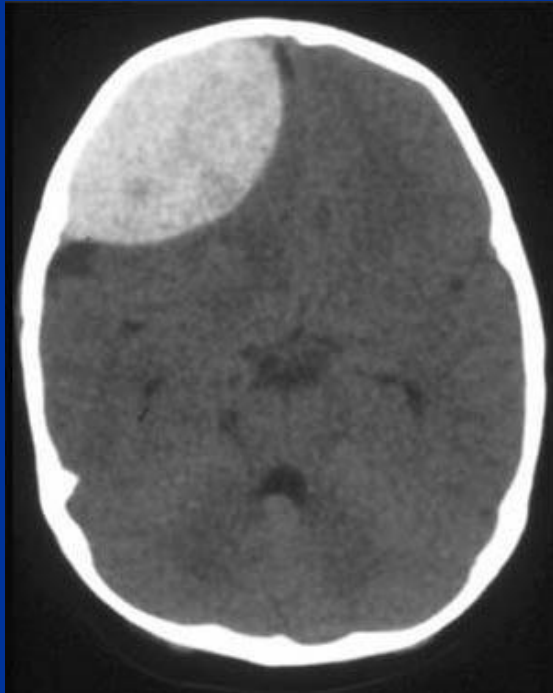
# Consequences of TBI

Brain oedema  
Fracture of skull  
Epidural haematoma  
Subdural haematoma  
Intracerebral haematoma  
Subarachnoideal bleeding



# TBI

- Focal damage (the area of the force)



# TBI

- Diffuse axonal injuries

- Damage to the pathways (axons) that connect the different areas of the brain. This occurs when there is twisting and turning of the brain tissue secondary to unrestricted head movement at the time of blast.
- Affects white matter of the cerebrum, corpus callosum, deep grey matter, internal capsule, upper brainstem and the cortico-medullary junctions of cerebral cortex
- Damage to rats axonal cytoskeleton results on loss of their elasticity and impaired transport and accumulation of axonal transport proteins within axonal swellings.
- Axonal injuries cause consciousness.

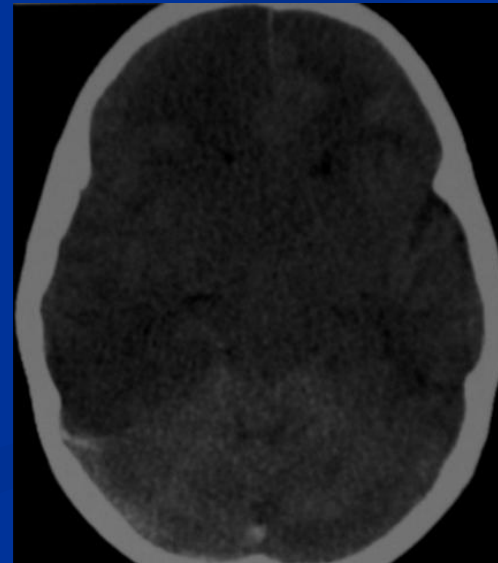
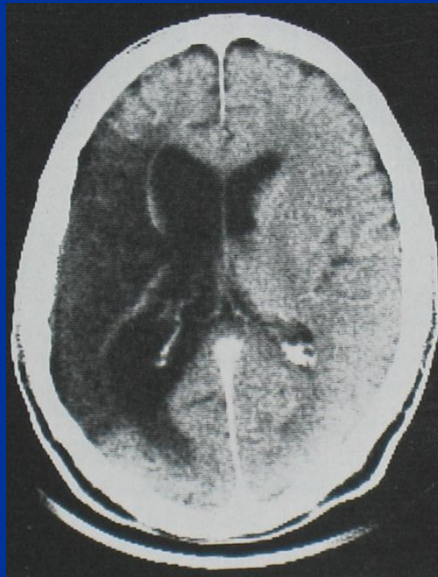


# Diffuse axonal injuries



# Closed skull-brain injury

Hypoxia, ischaemia (penumbra)  
Diffuse microvascular damage  
(damage of autoregulation , CO2  
reactivity) - oedema

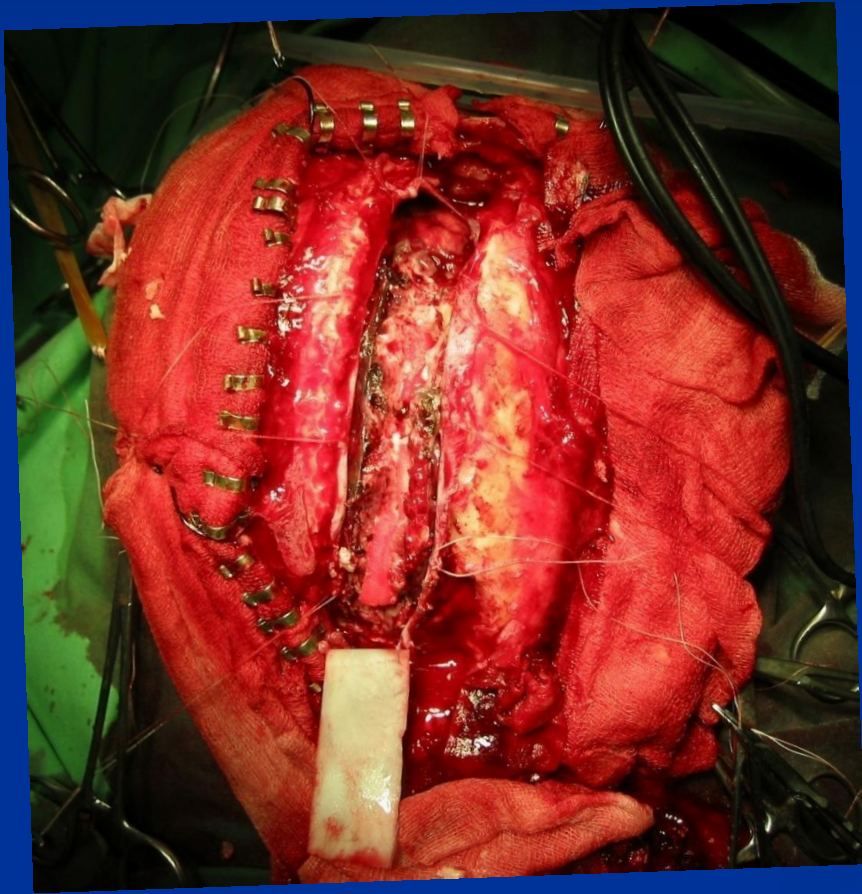




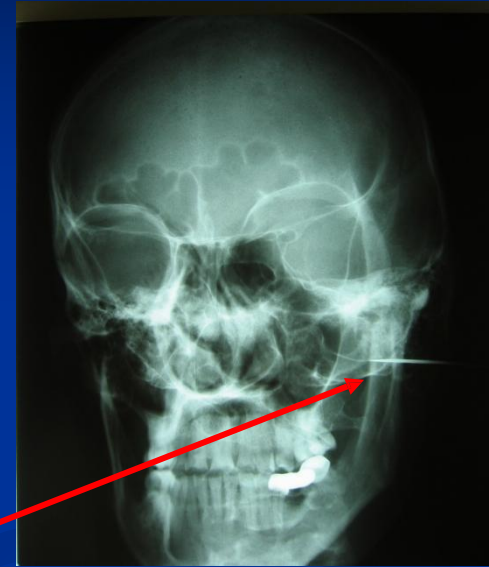
# Open skull-brain injury (foreign body)



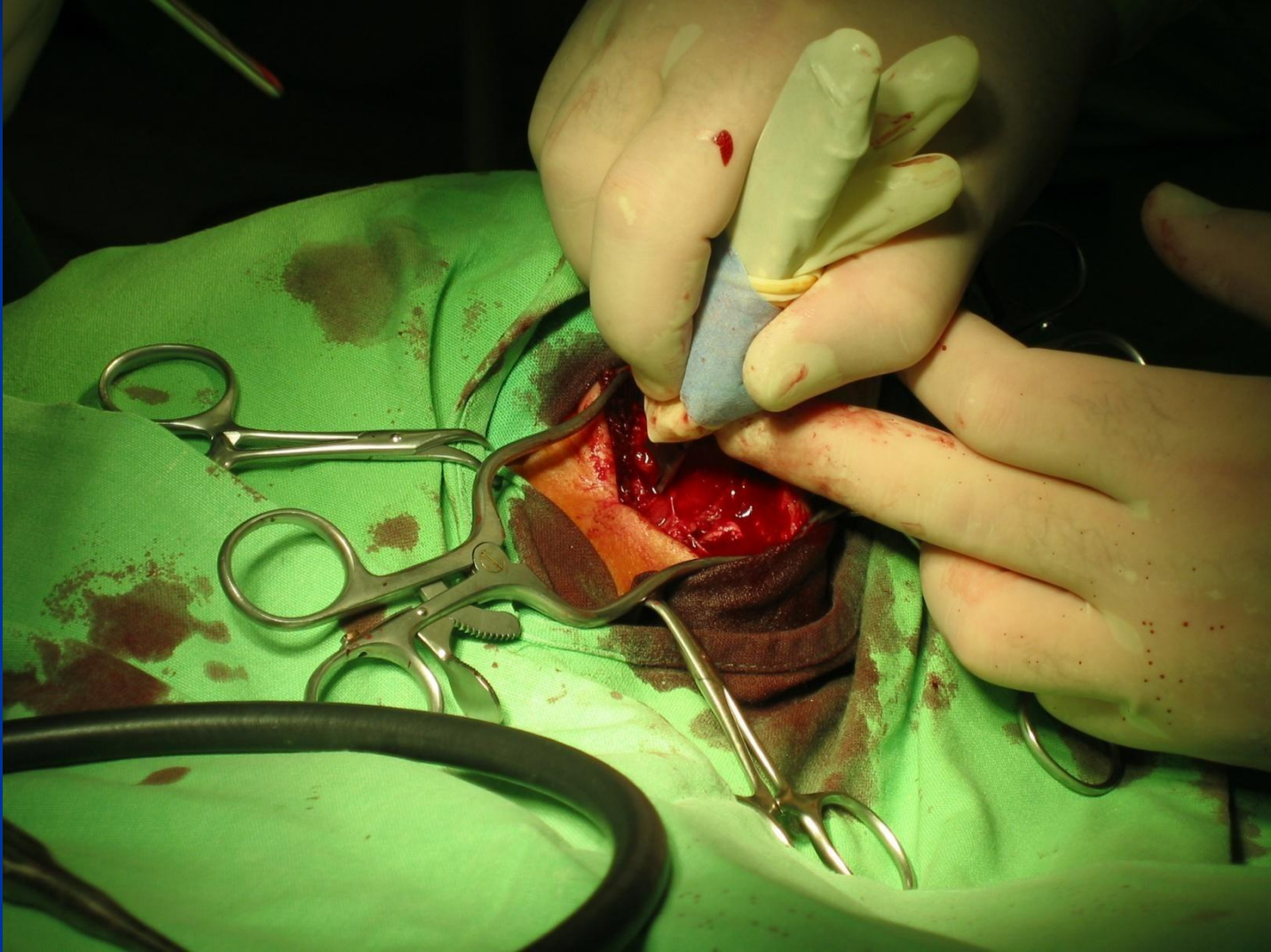




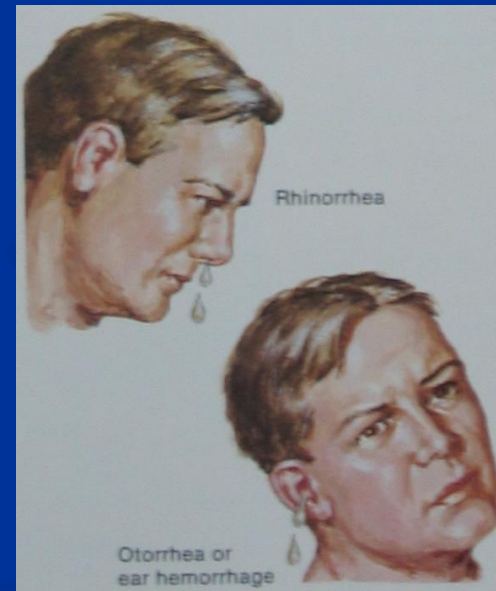
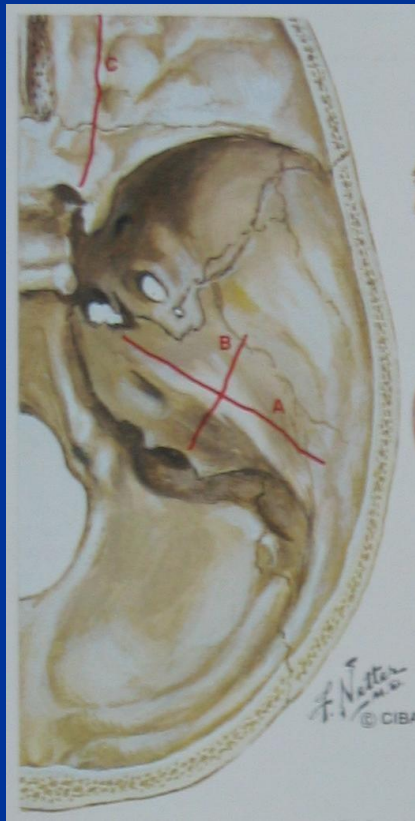
# The seriousness of injury:





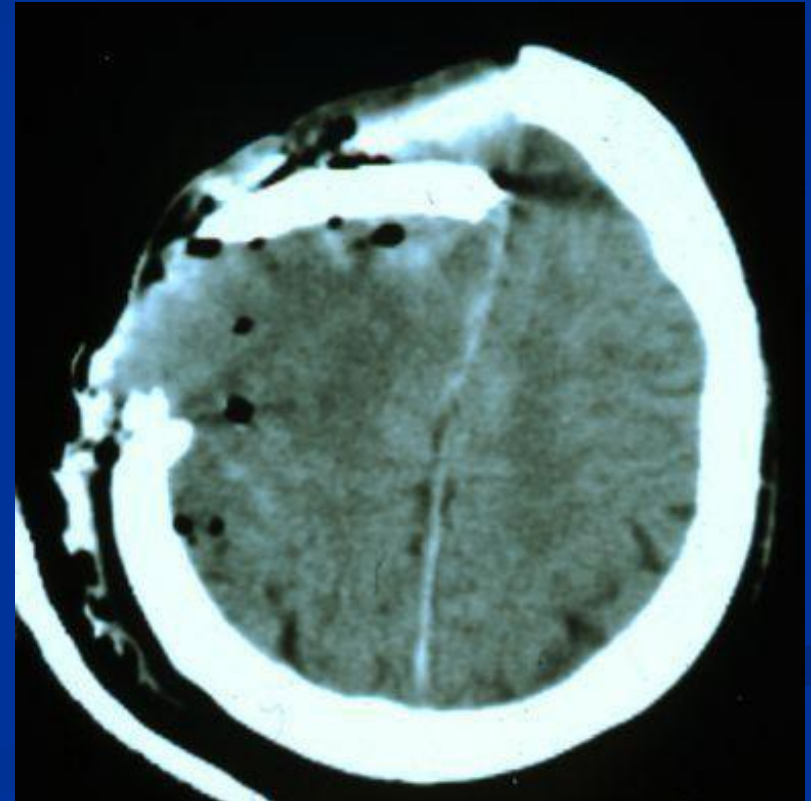
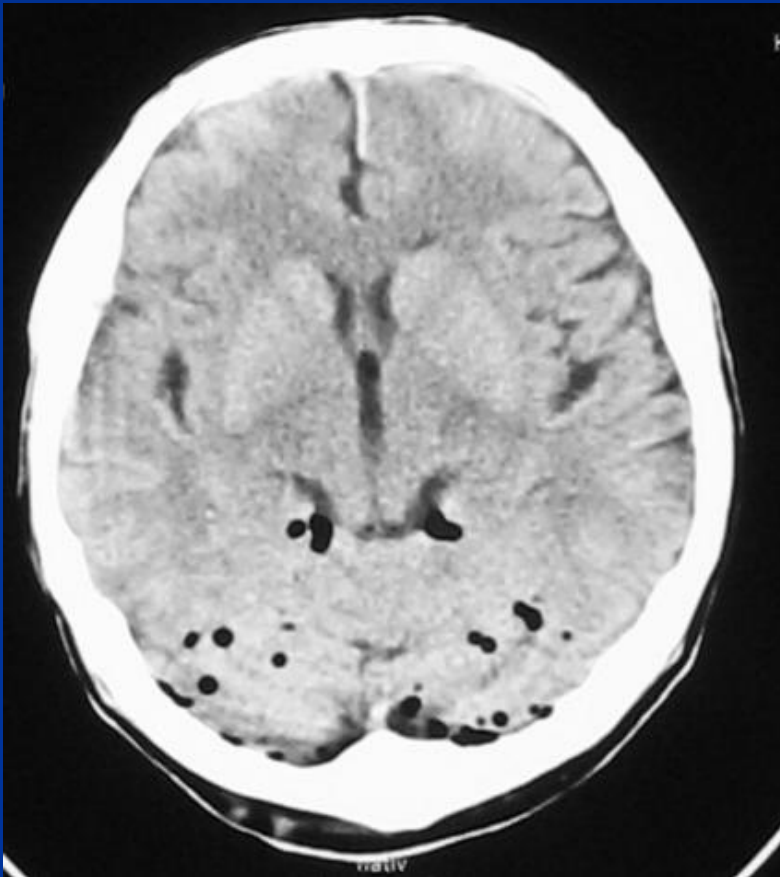


- Skull base fractures (frontale, pyramis)



# Open skull-brain injury

Pneumocephalus: intracranial air





# Head injury



# Comotio cerebri

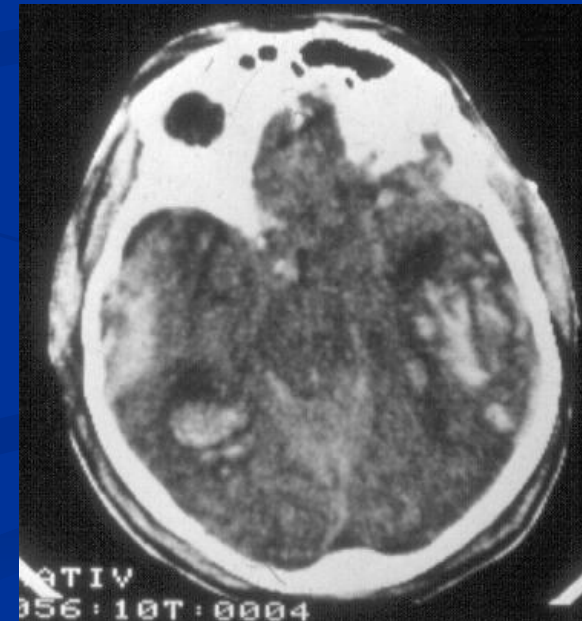


- Lost of consciousness (2-10 min)
- Apnoe (6-12 sec)
- Bradycardia (1-2 min)
- Amnesia: congrad, retrograd, anterograd
- CT negative
- Treatment: rest in bed, observatio, dehydration?



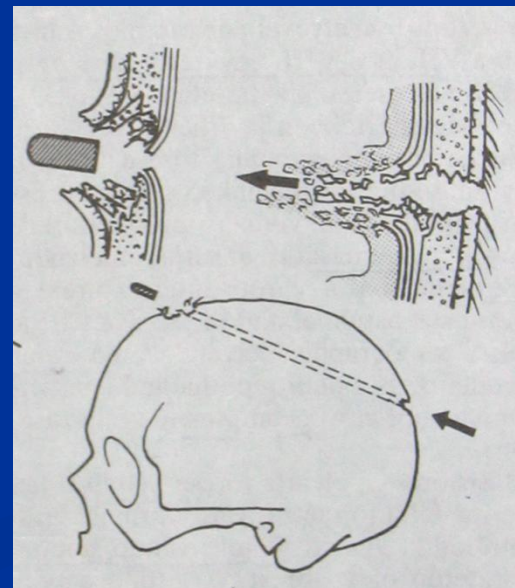
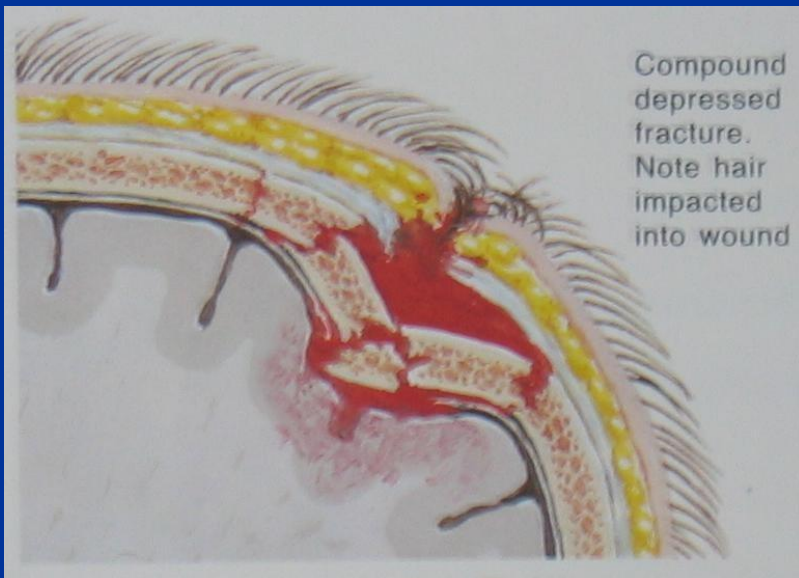
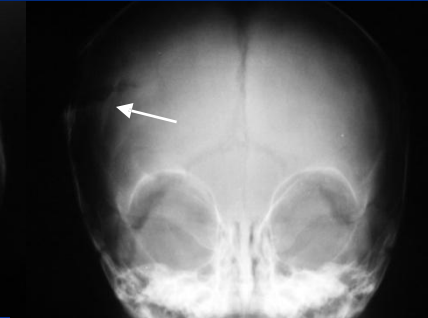
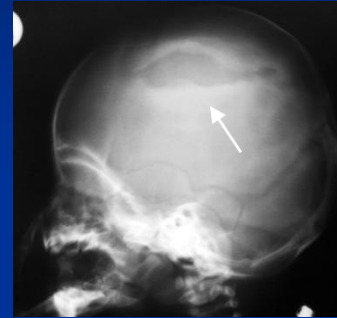
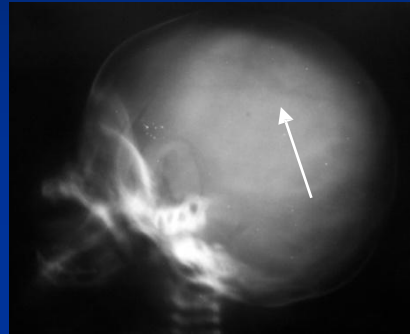
# Contusio cerebri

- Lost of consciousness
- Other neur. deficit
- Damage of tissue of the brain  
haematoma, or oedema
- Treatment: oedema ↓ if GCS <8  
measure of ICP clin. and radiol. observation  
haematoma evacuatio?

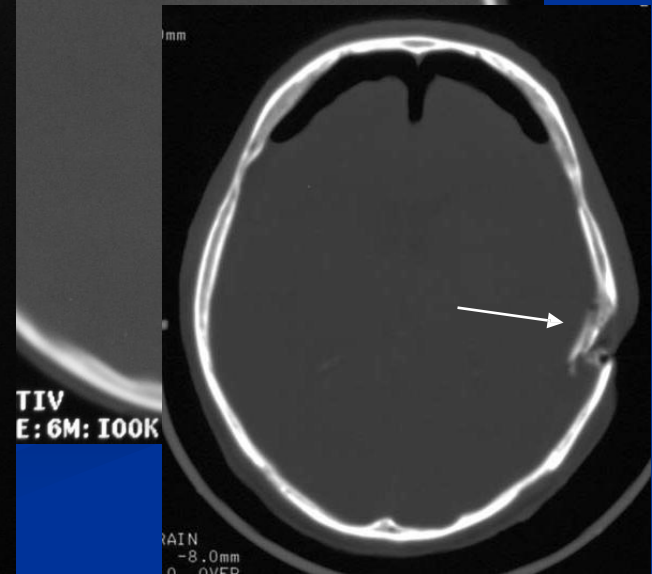
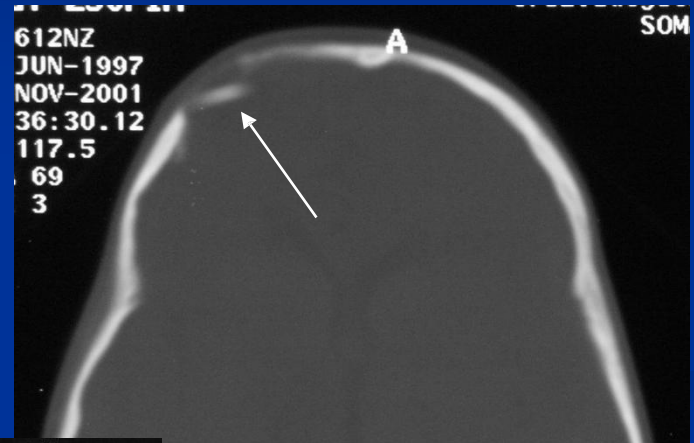
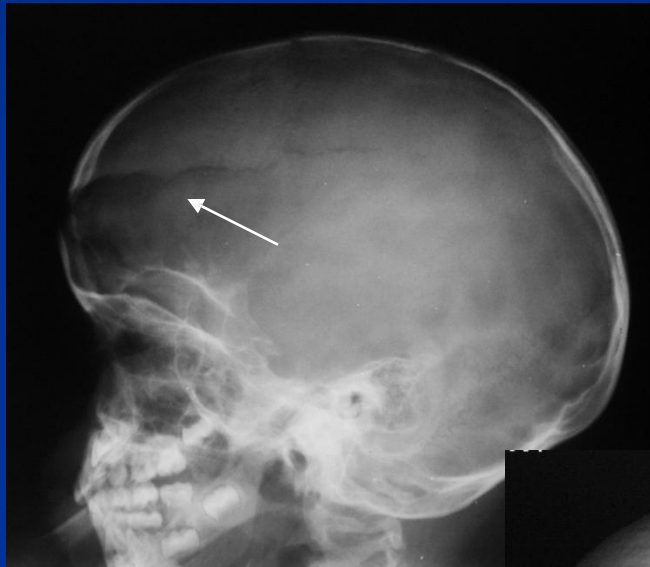


# Skull bone fracture

Linear  
Impressive  
Perforating

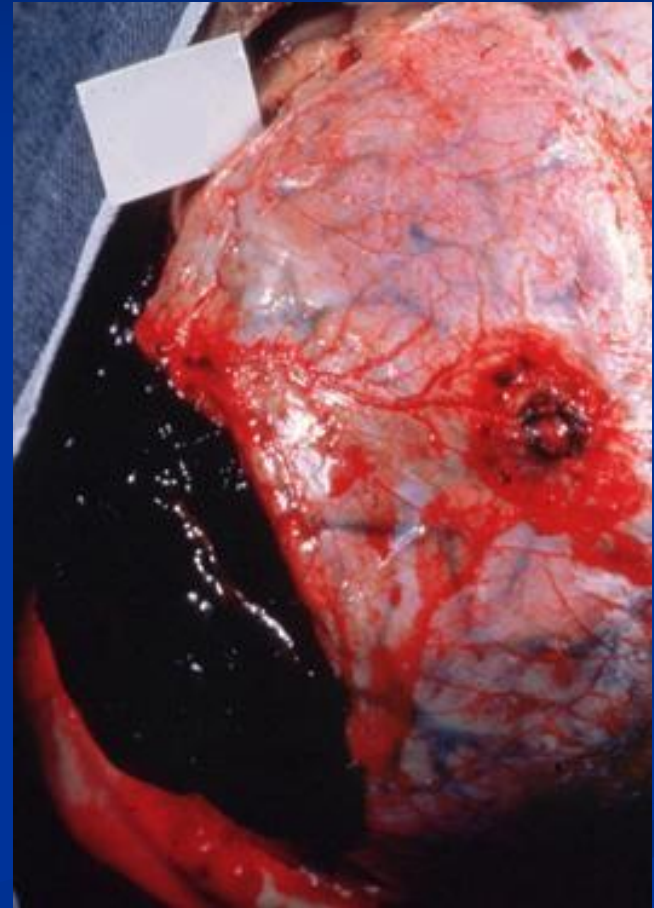
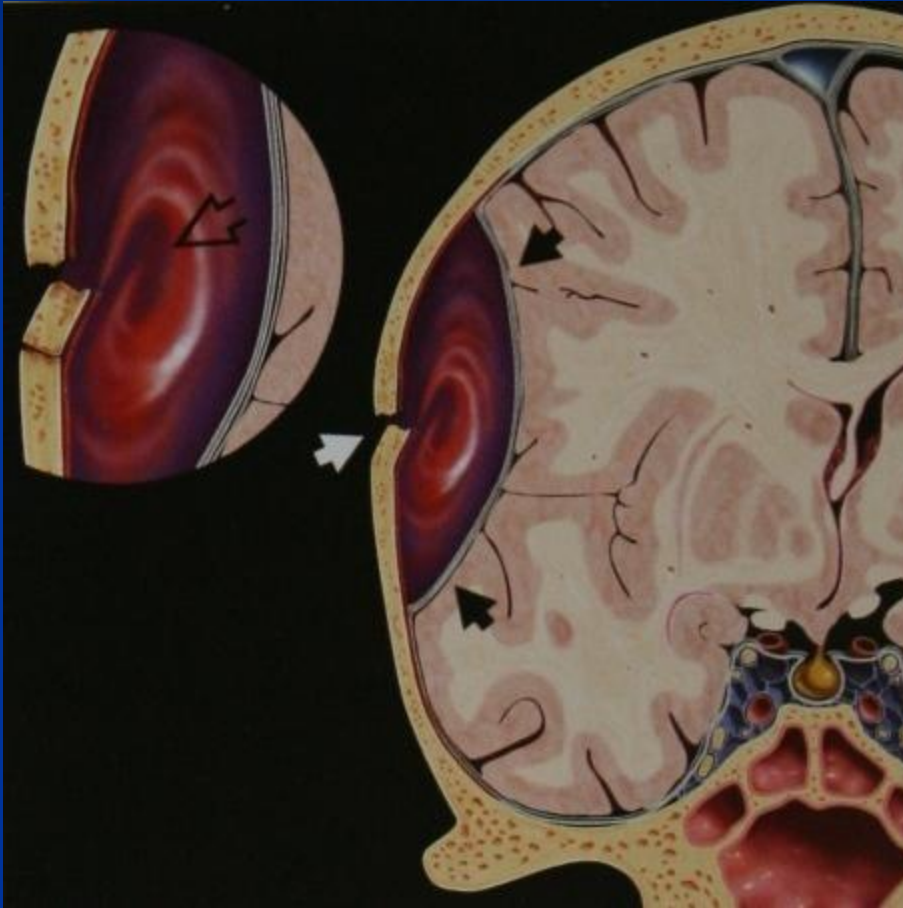


# Skull bone fracture



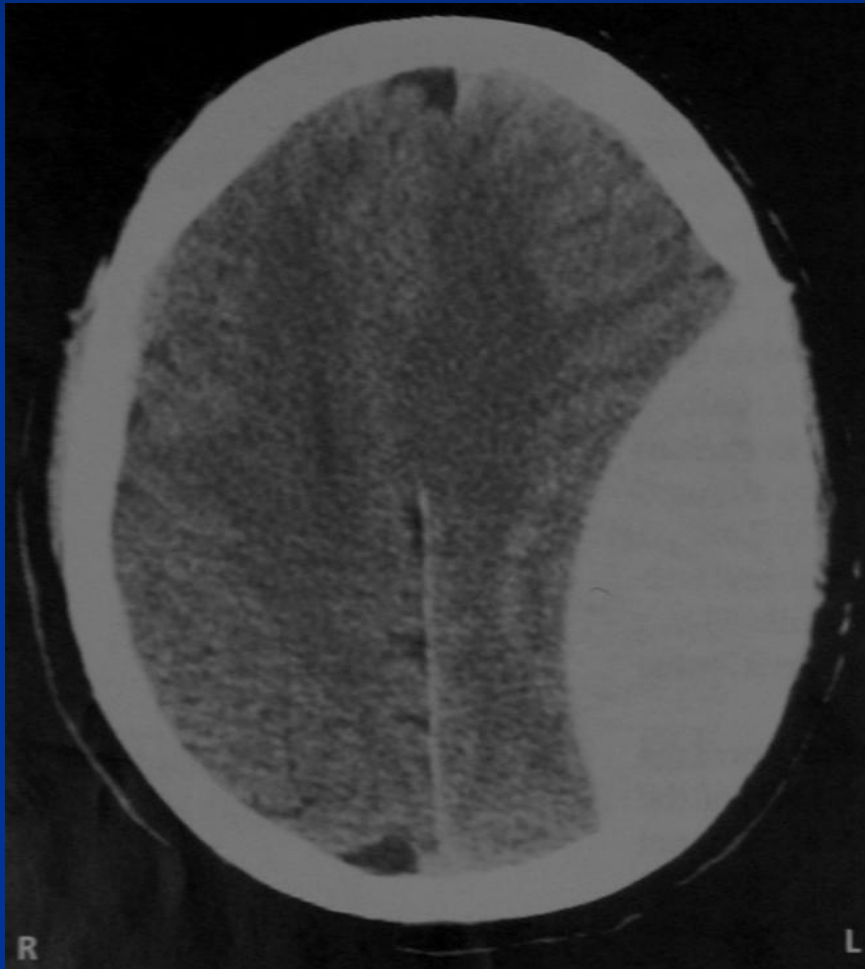


# Epidural haematoma



a. meningea media damage, CT: biconvex (lentiform sign)

# Epidural haematoma



Treatment: haematoma evacuatio, supply of bleeding

# Subdural haematoma

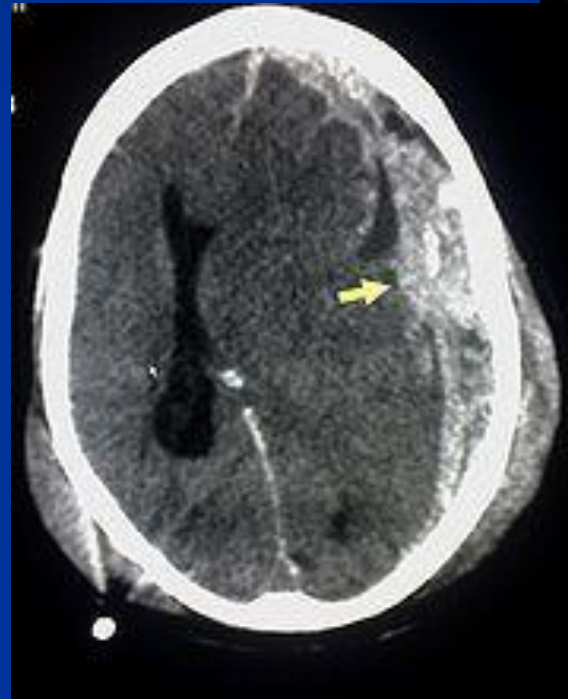
Result from the tearing of bridging veins crossing the subdural space

- acute
- subacute
- chronic  
(reduced density)





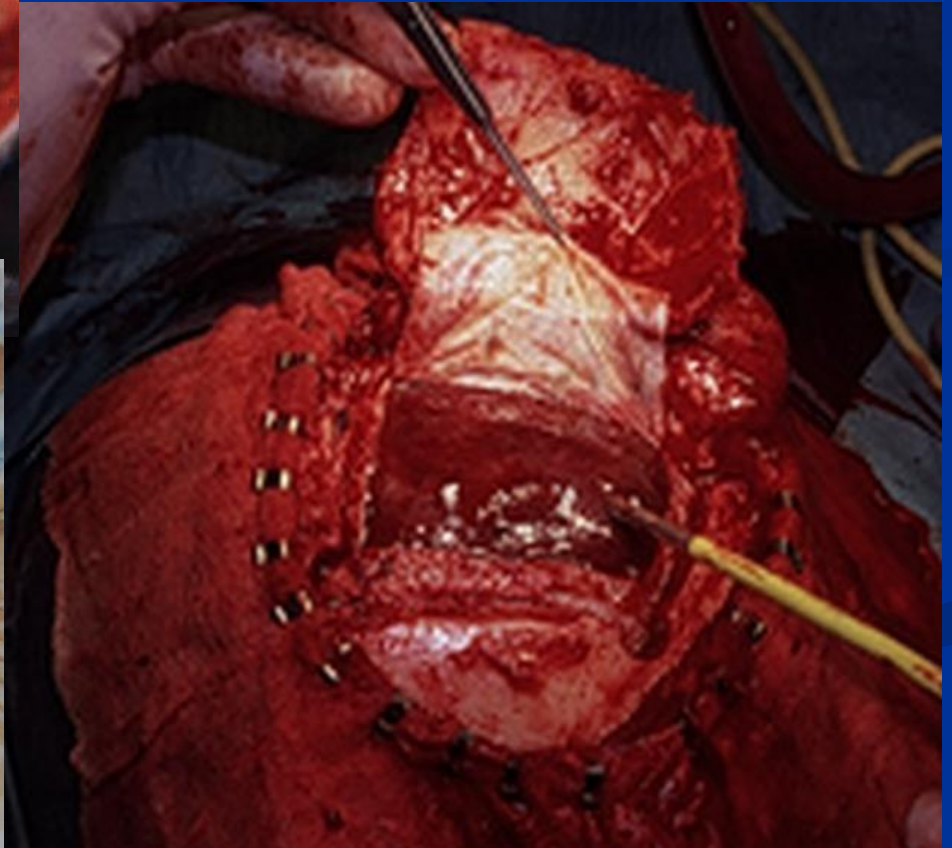
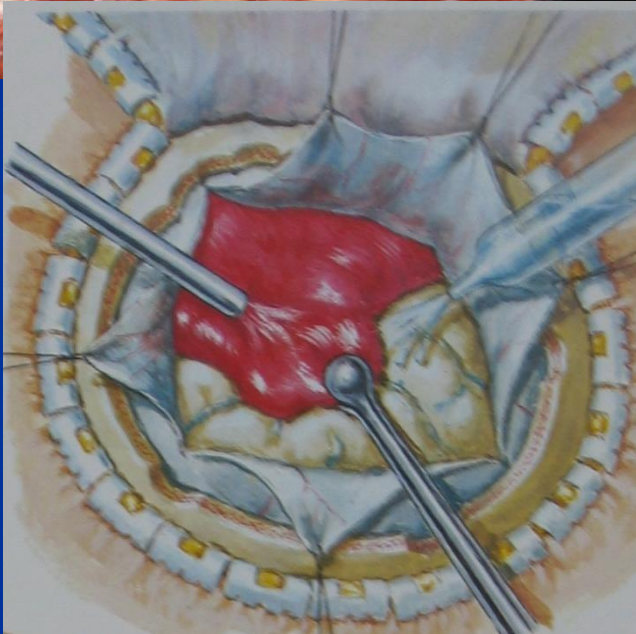
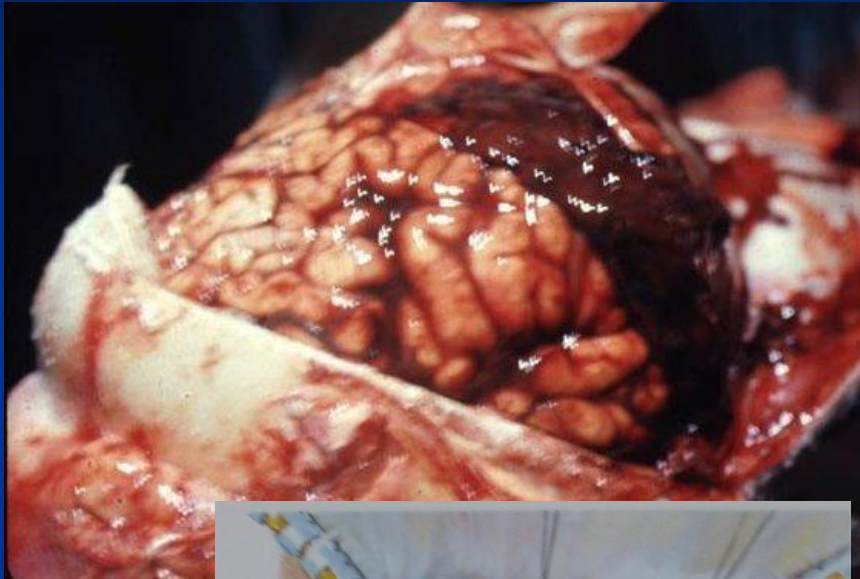
# Subdural haematoma: acut



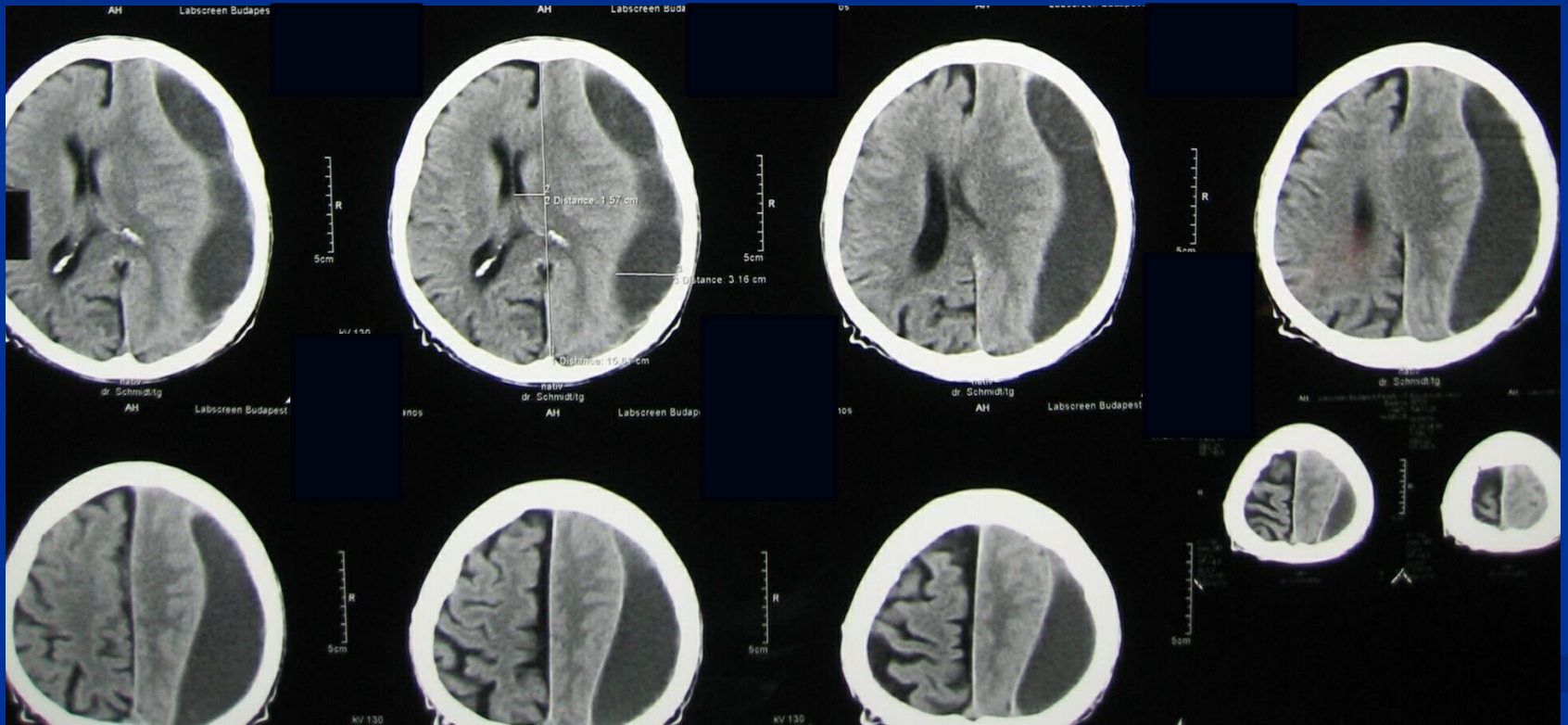
Treatment: haematoma evacuation (craniotomy, burr hole)



# Subdural haematoma



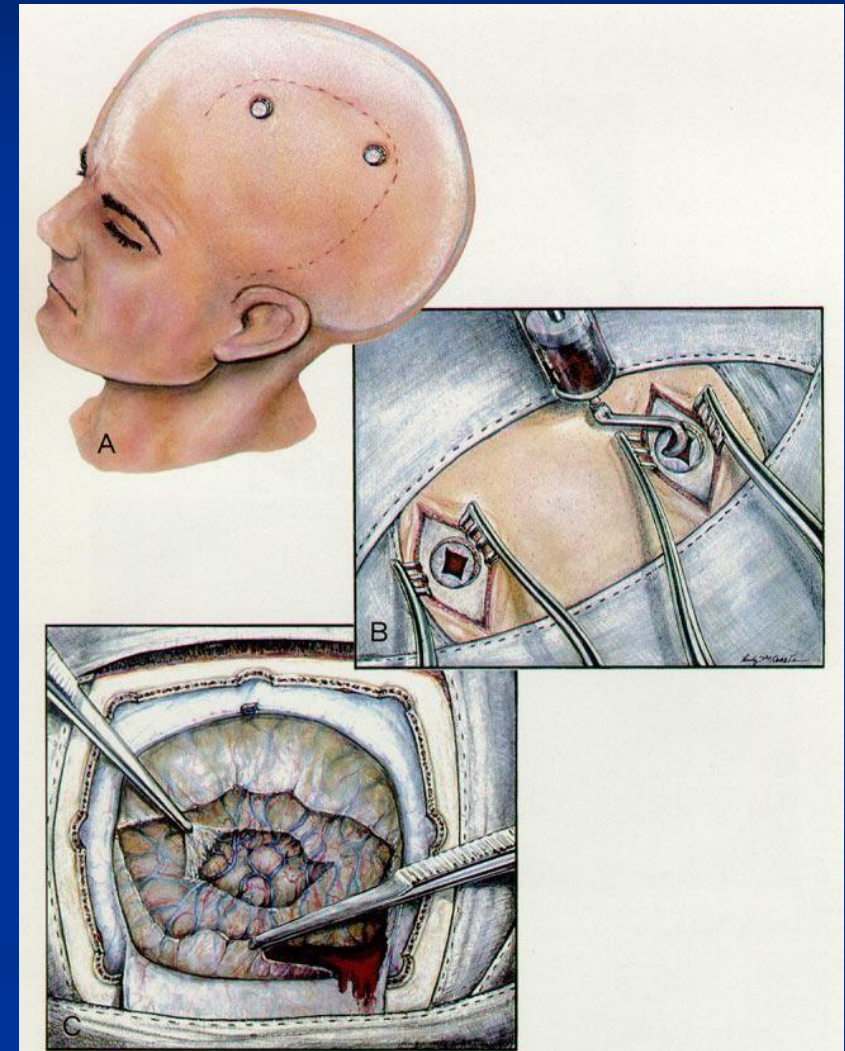
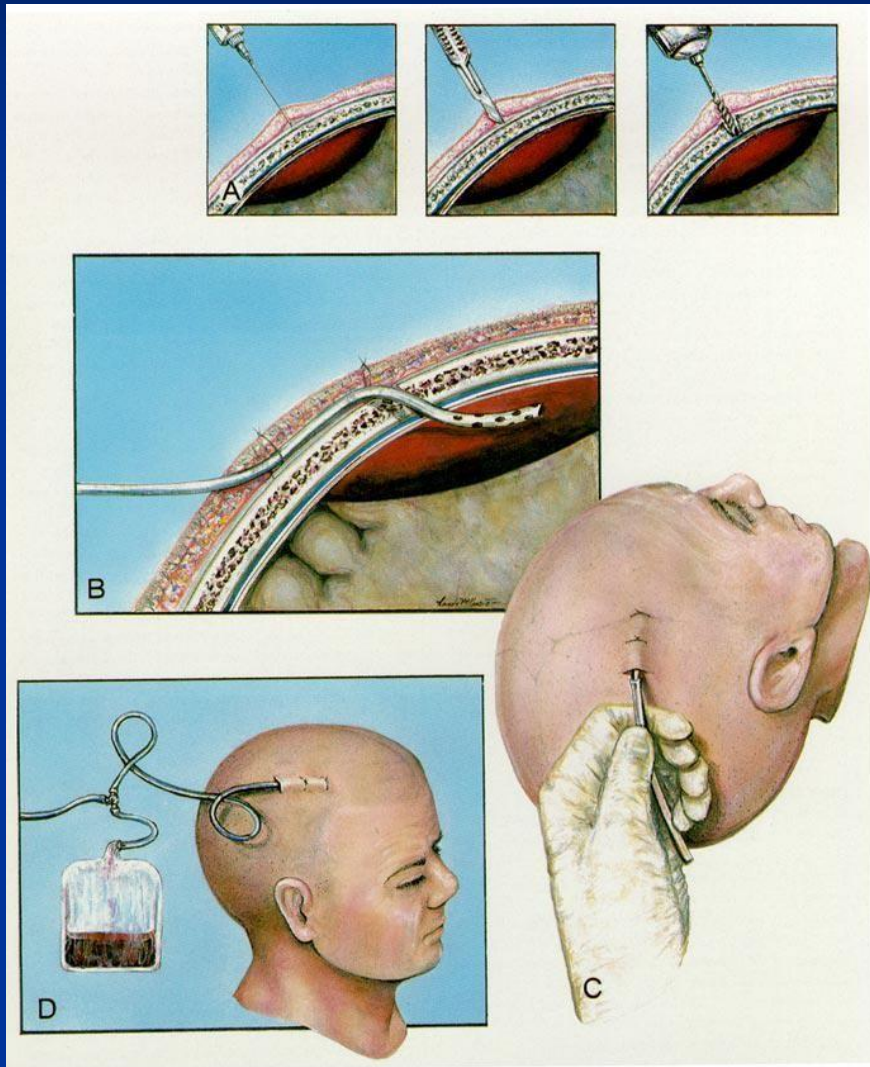
# Subdural haematoma: chr.



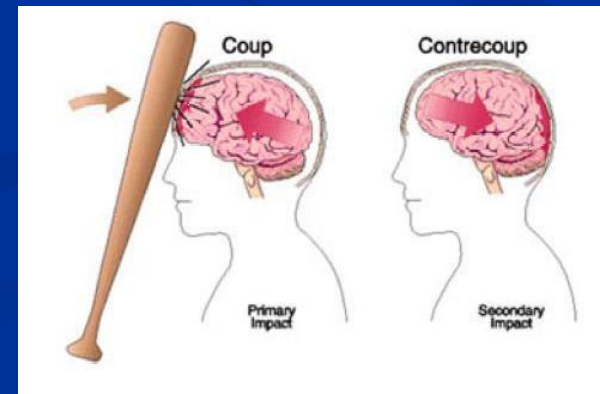
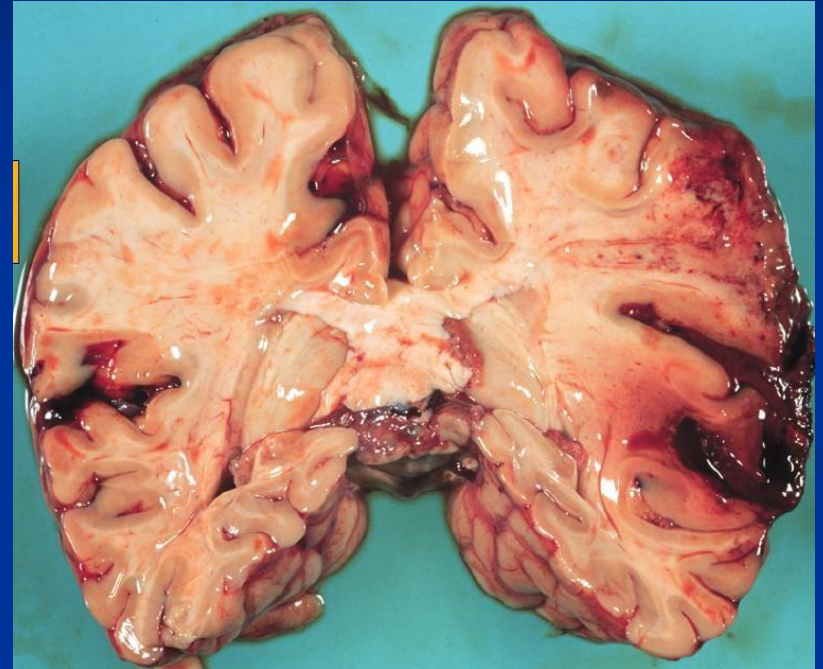
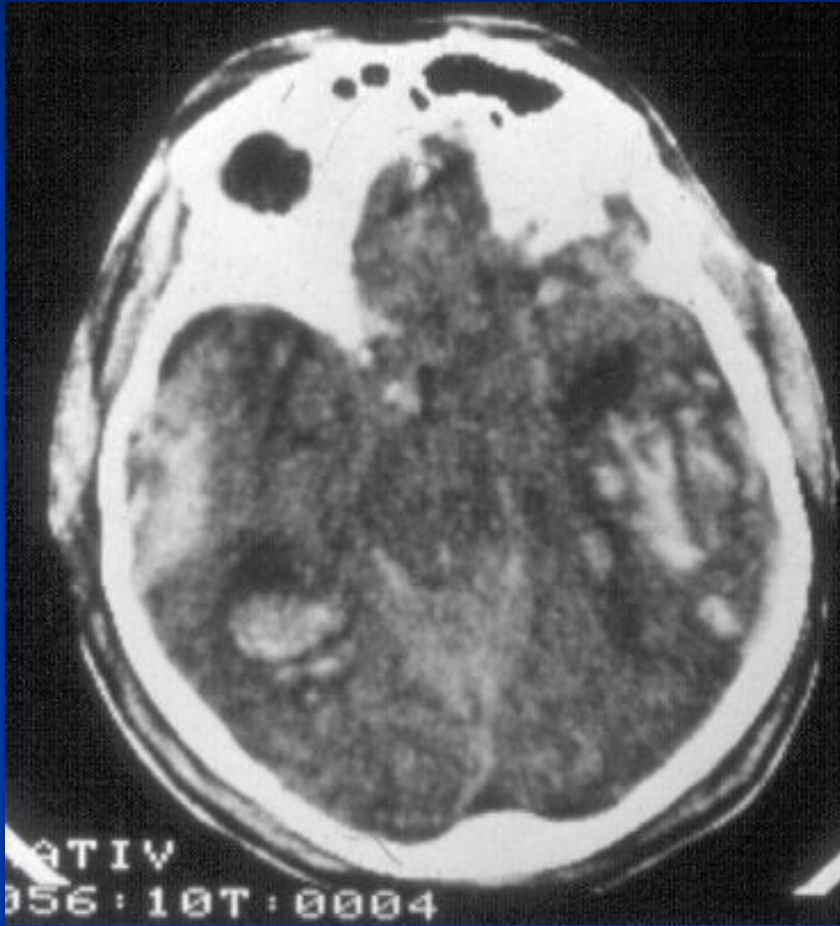
Treatment: haematoma evacuation (burr hole, craniotomy)



# Subduralis haematoma: chr.



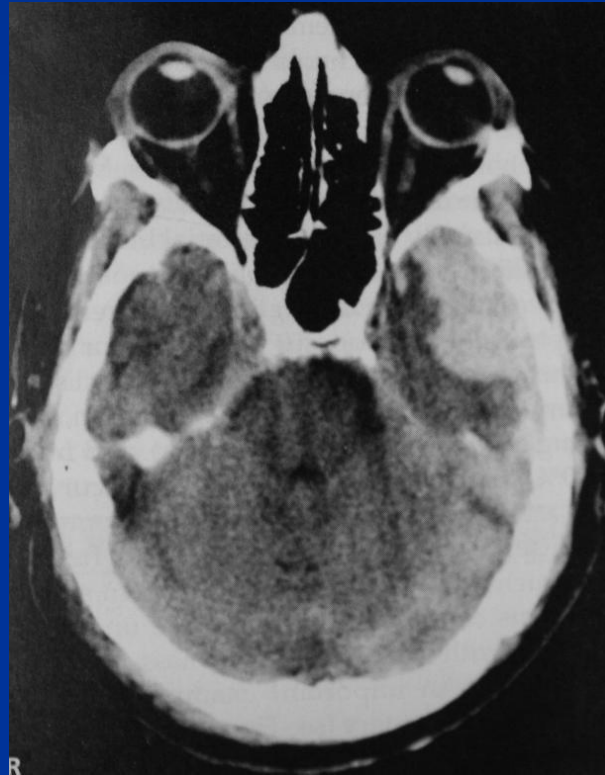
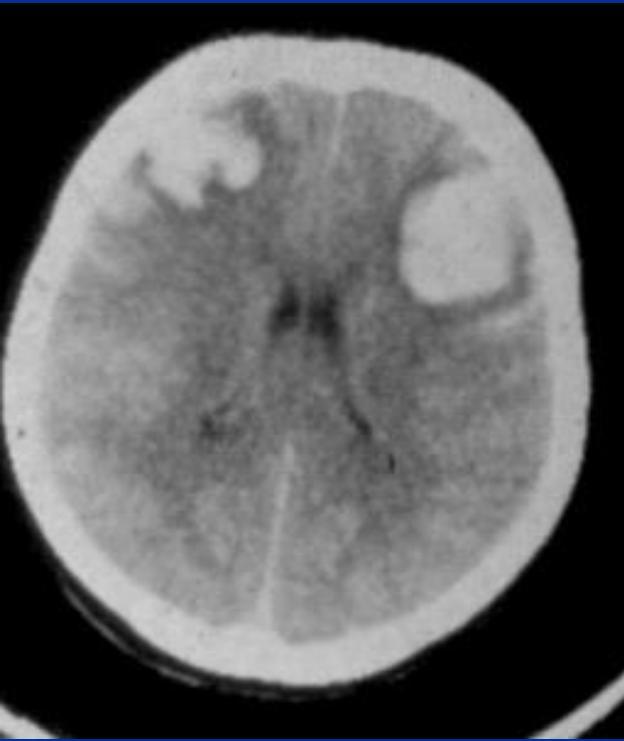
# Cerebral haemorrhagic contusion



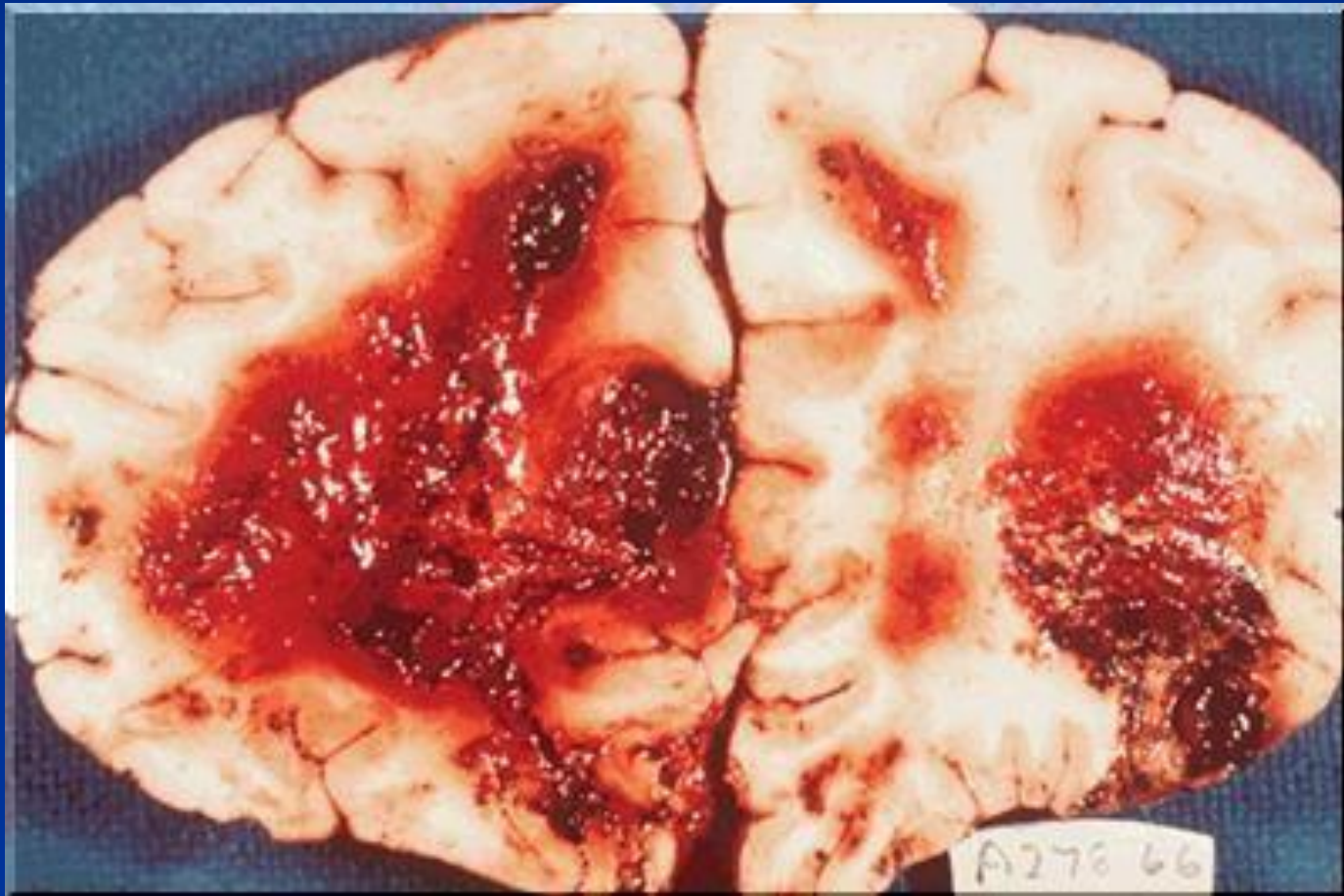
Contre-coup – Most contusions represent the brain coming to a sudden stop against the inner surface of the skull (the opposite side)



# Cerebral haemorrhagic contusion



# Cerebral haemorrhagic contusion



# Treatment

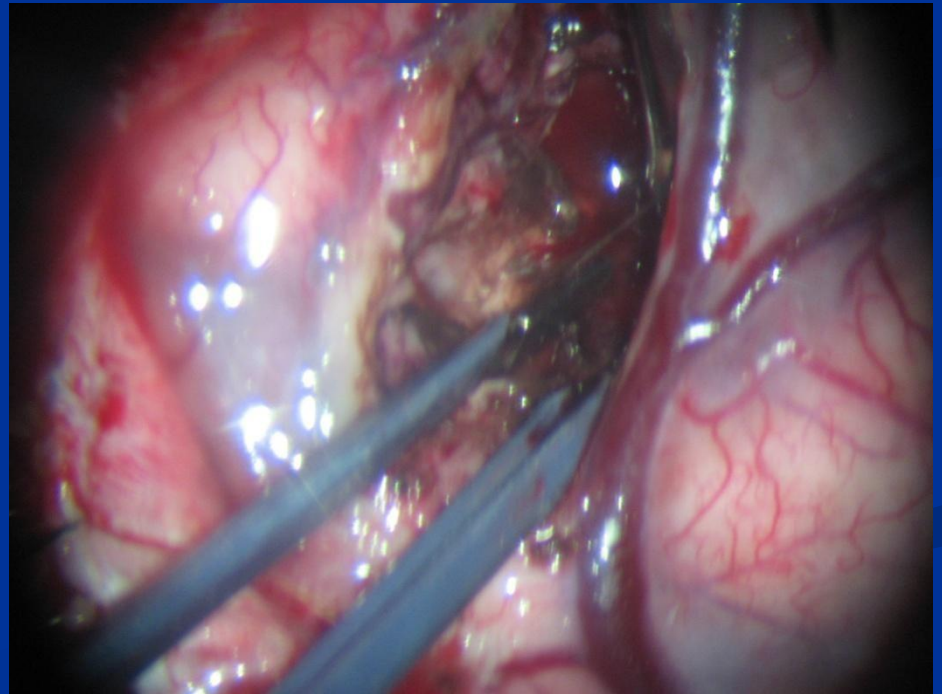
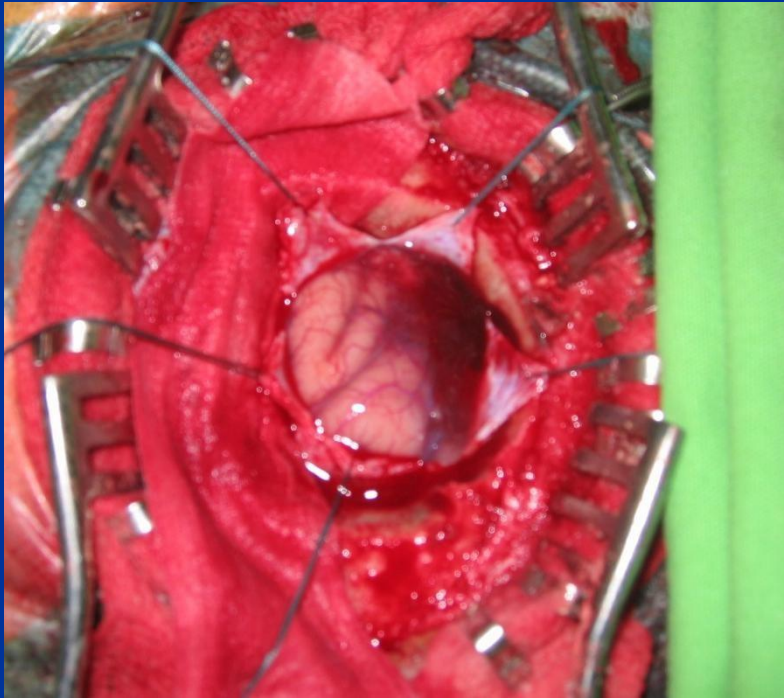
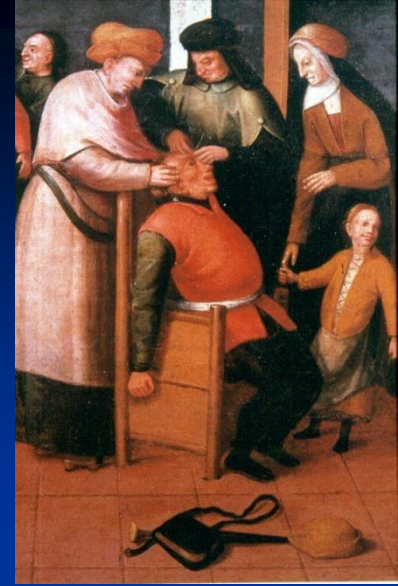
Conservative: monitoring and controlling ICP (decrease of oedema, dehydration)

Operation: craniotomy and haematoma evacuation

Rarely: punctio and drainage

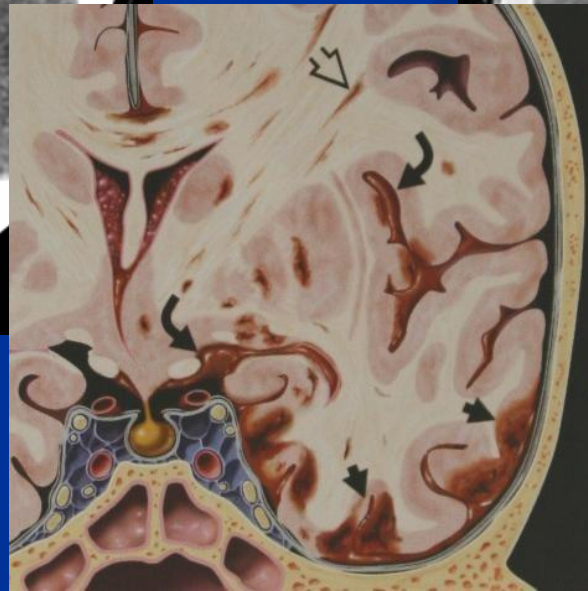
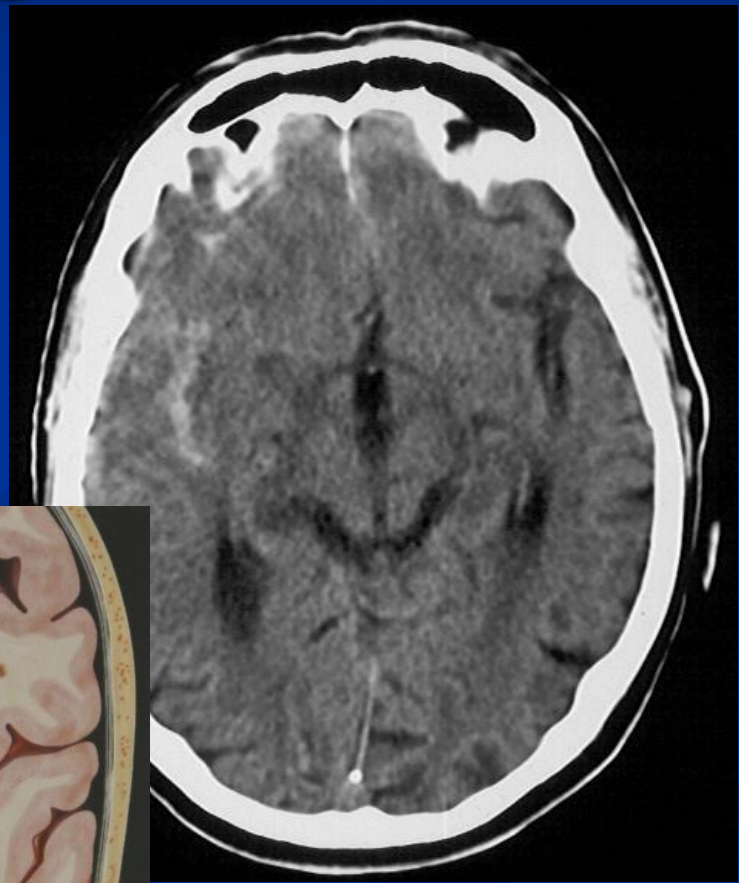
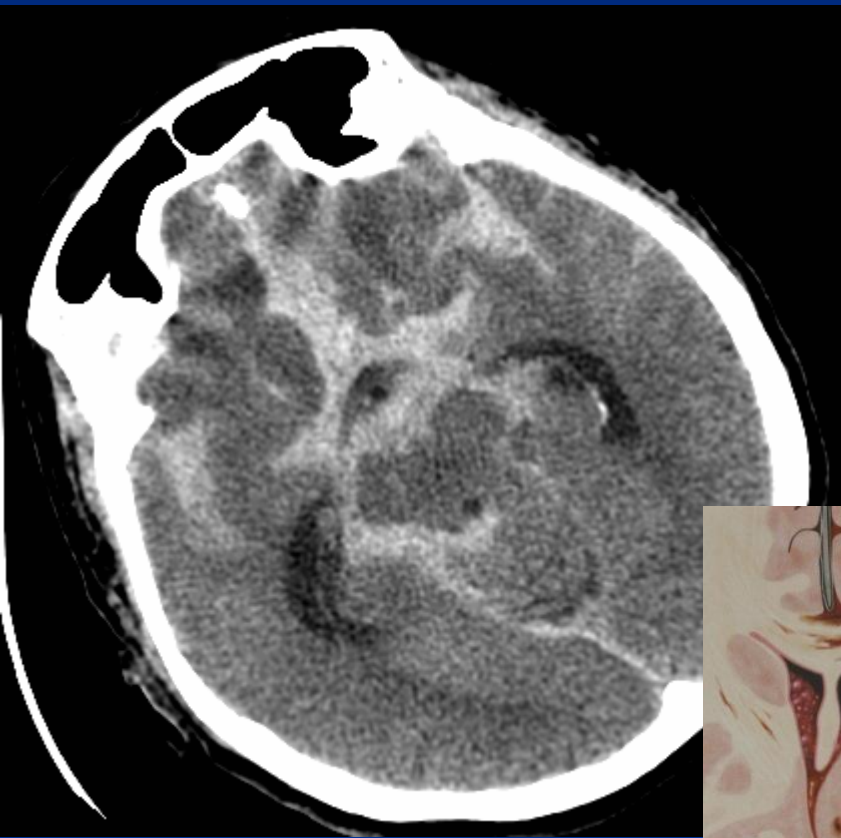


# Haematoma evacuatio





# Traumatic subarachnoid haemorrhage



About treatment:

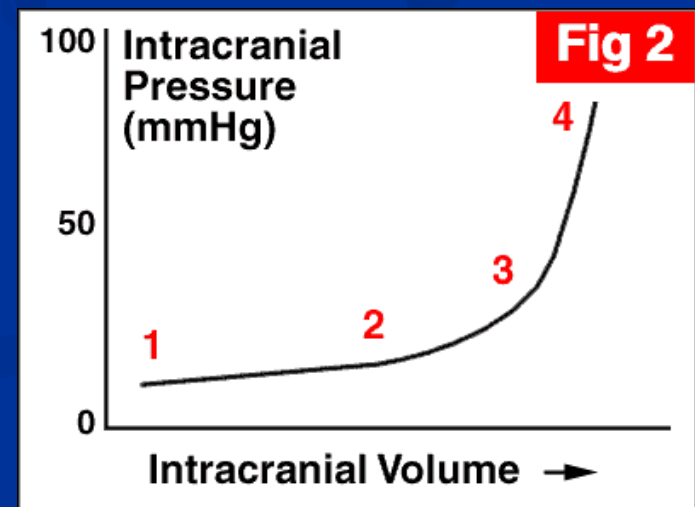
Prevention and treatment of intracranial hypertension

Maintenance of adequate and stable cerebral perfusion pressure

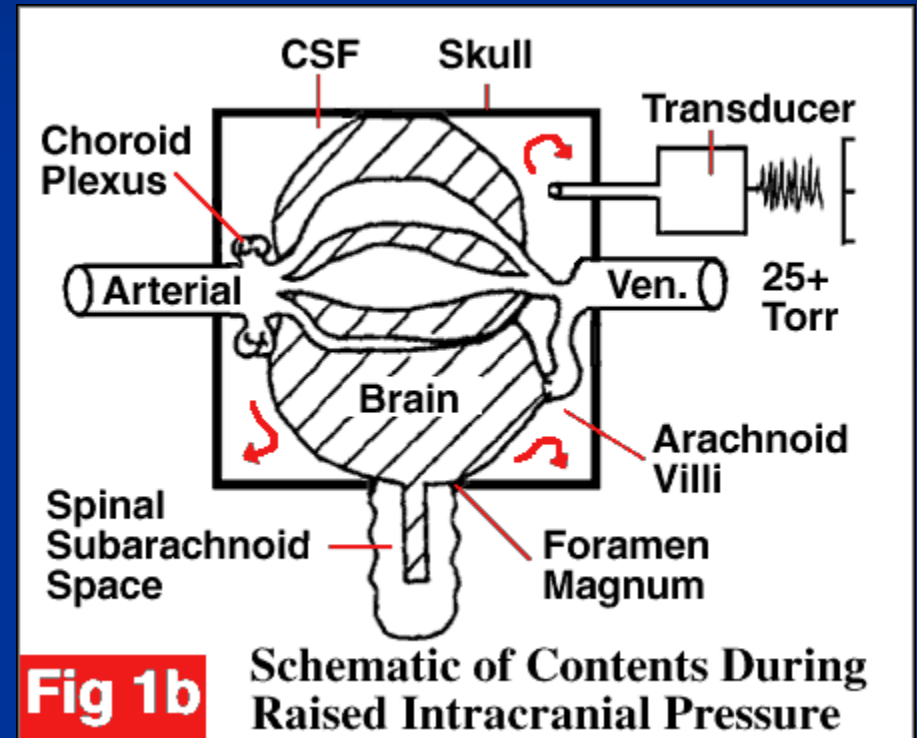
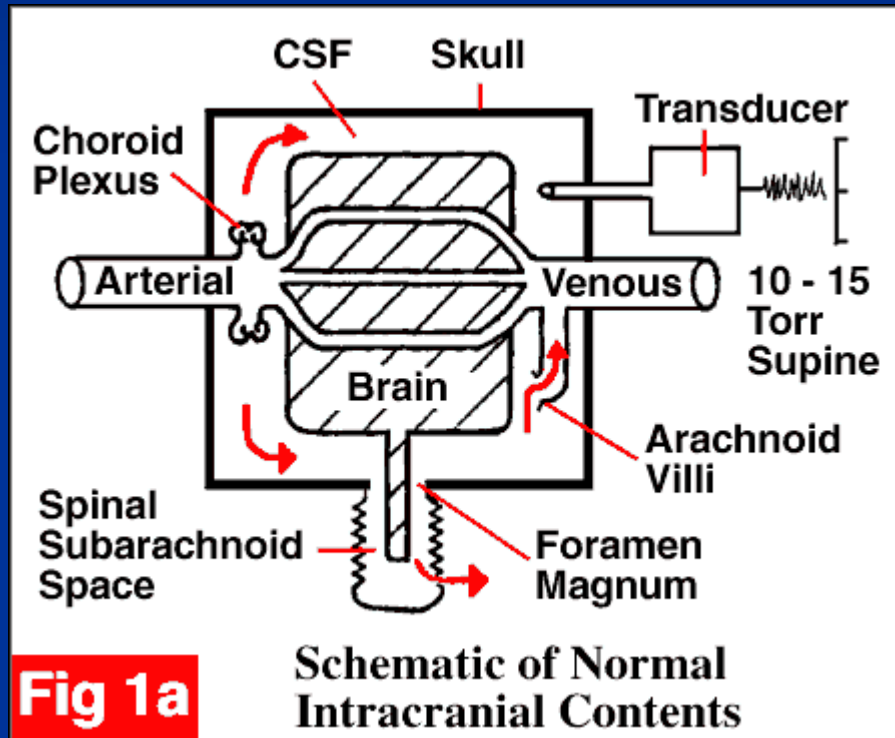
Avoidance of systemic secondary brain insults

Optimization of cerebral haemodynamics and oxygenation

Norm. ICP < 10 mmHg



# Intracranial volume

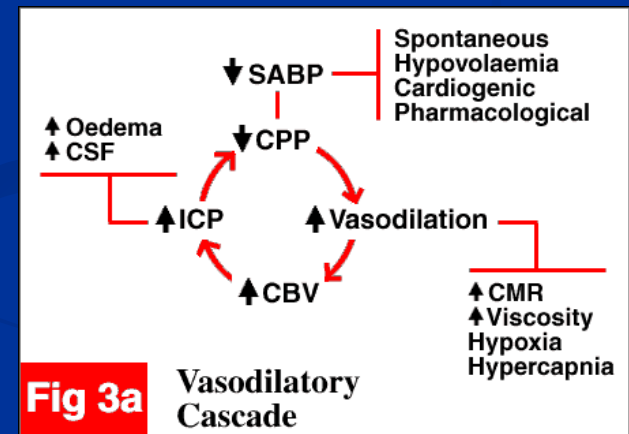


brain (80%), blood (12%) and liquor (8%).

# ICP and CPP

Cranial perfusion pressure = mean arterial pressure – intracranial pressure  
(**CPP = MAP - ICP**)

Normal CPP= 80Hgmm  
( mild oedema: (GCS 13-15)  
ICP=20 or more Hgmm,  
severe oedema,somatosus-  
soporosus (GCS <8)  
Normal ICP= 5-13Hgmm

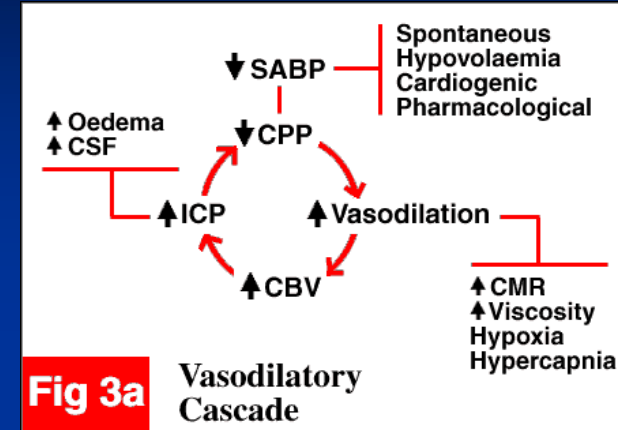




# Pathophysiology

„circulus vitiosus“:

- trauma
- disturbed of microvascularis regulatio ischaemia, hypoxia
- oedema - (cytotoxic - vasogen) ICP increase
- disturbed of brain tissue perfusion , ischaemia



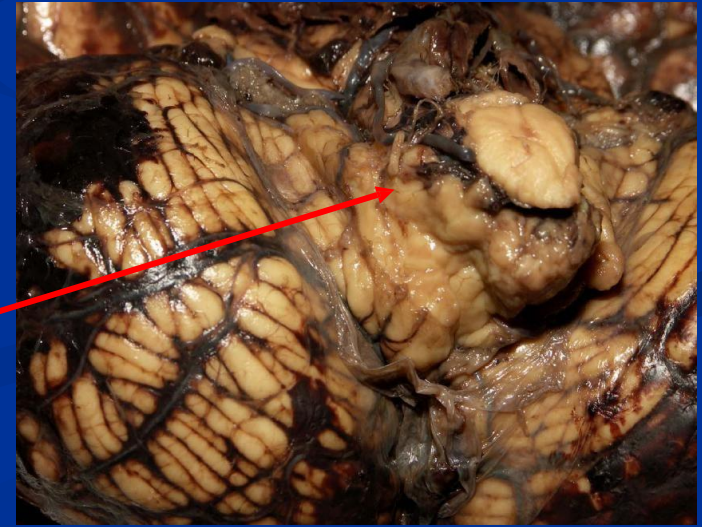
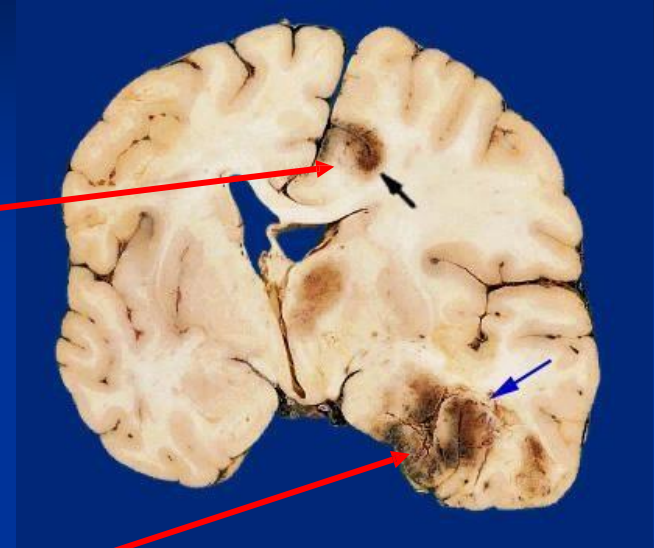
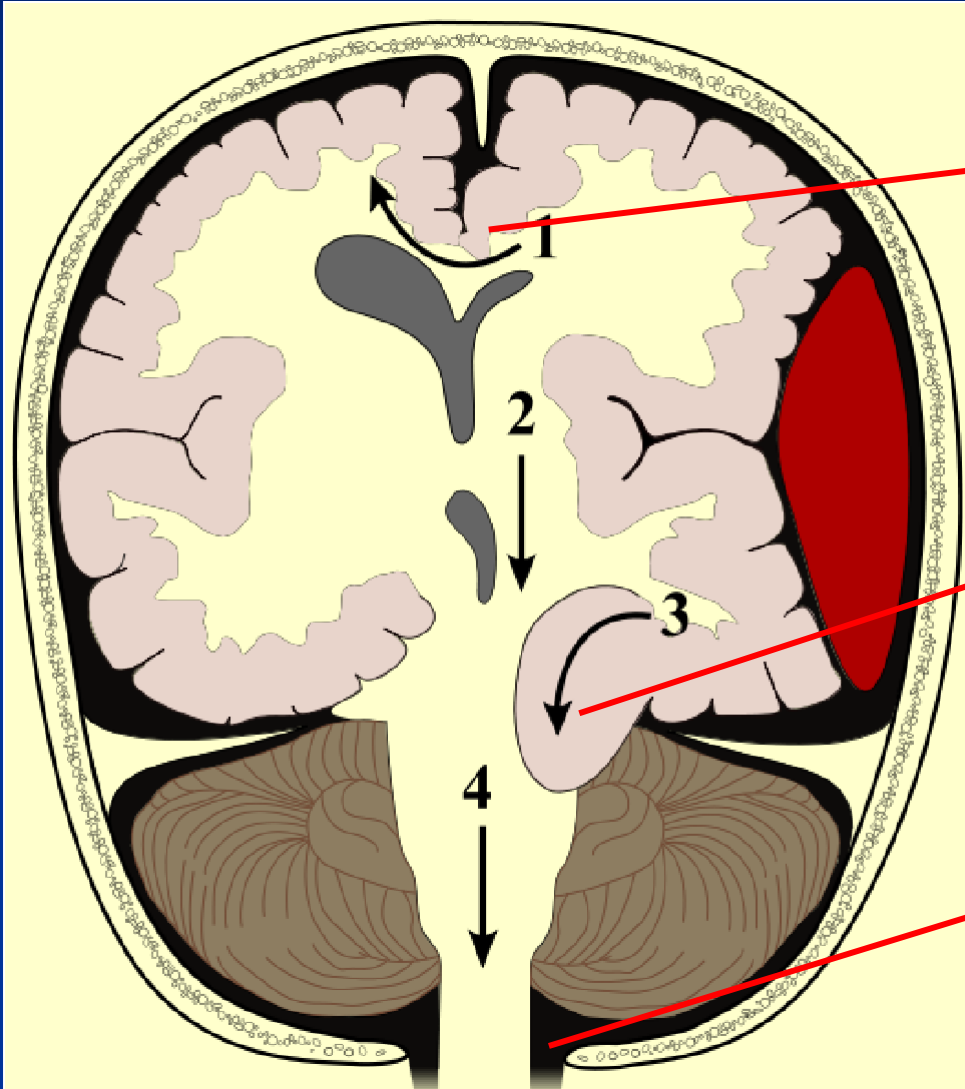
# Therapy of oedema

- Hyperosmotic (Mannitol)
- hyperventillation (PCO<sub>2</sub> decrease – vasoconstrictio)
- liquor drainage (ventricular punctio)
- barbiturat therapy (decrease of brain tissue metabolism)
- diureticum (Furosemid)

# Symptoms of increase ICP

- Headache
- Vomiting
- Papilla oedema, visual disturbances
- Pupilla difference
- Loss of consciousness
- ”Brainstem” herniation: bulbus deviation, Wernicke-Mann (flexio-extensio)
- Disturbe of cardiorespiratory system
- Cushing-reflex: increase of blood pressure  
+bradycardia

# Brain herniation





# Treatment

subdural, epidural, i.c. haematoma evacuation

- ICP < 20 Hgmm and CPP > 60 Hgmm  
Diuretic therapy, osmotherapy, liquor drainage, intubation, hyperventilation
- If in spite of osmotherapy ICP > 20 Hgmm  
barbiturat coma
- If in spite of using osmotherapy and barbiturat  
ICP > 20 Hgmm

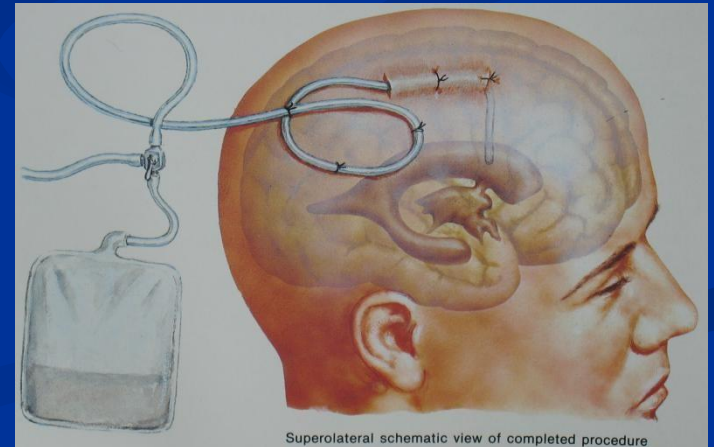
**Decompressiv craniectomy**

# ICP monitoring

Under GCS 8

Type of ICP monitoring

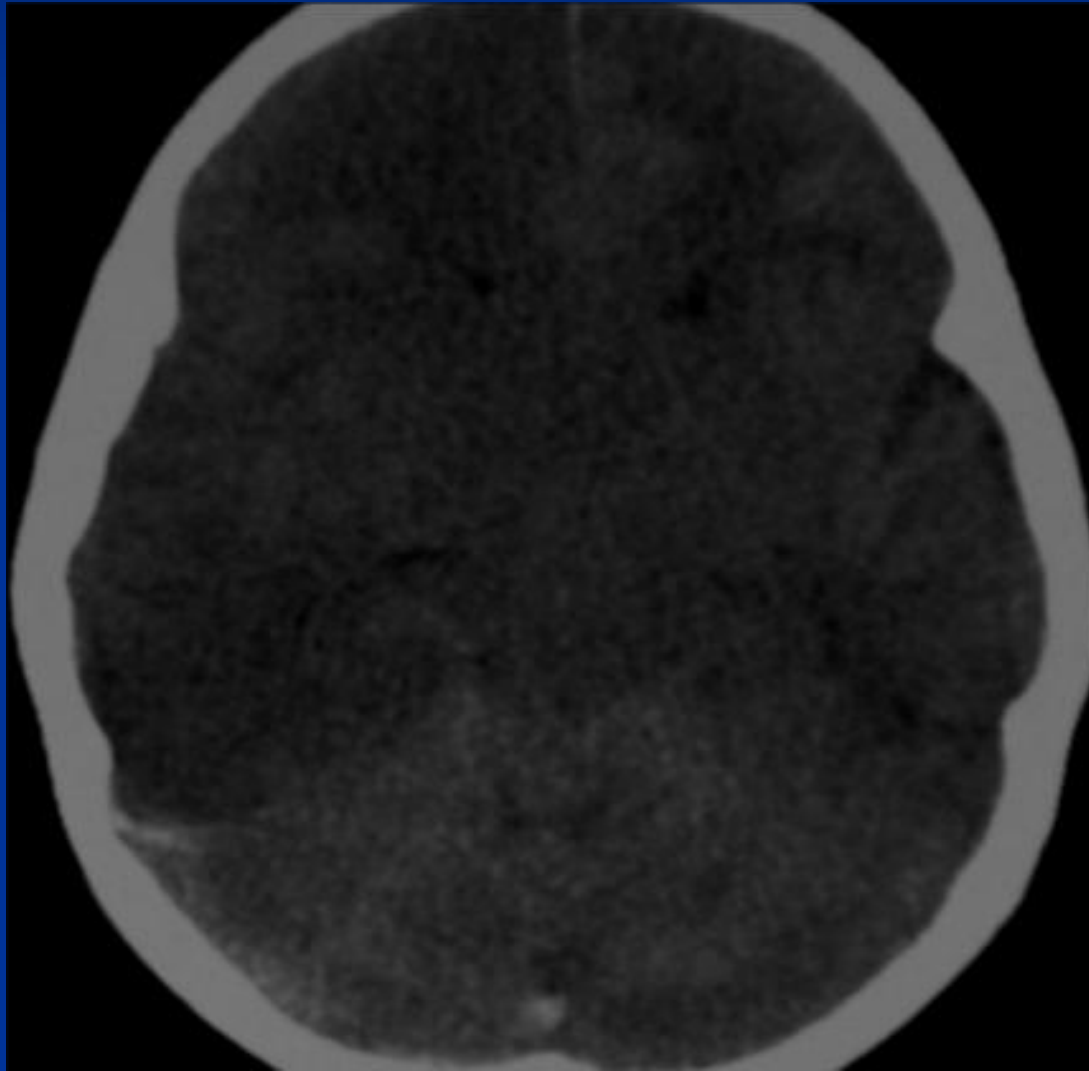
- Epidurális
- Intraparenchymális
- Intraventriculáris  
(with liquordrainage)



# Intensive care unit

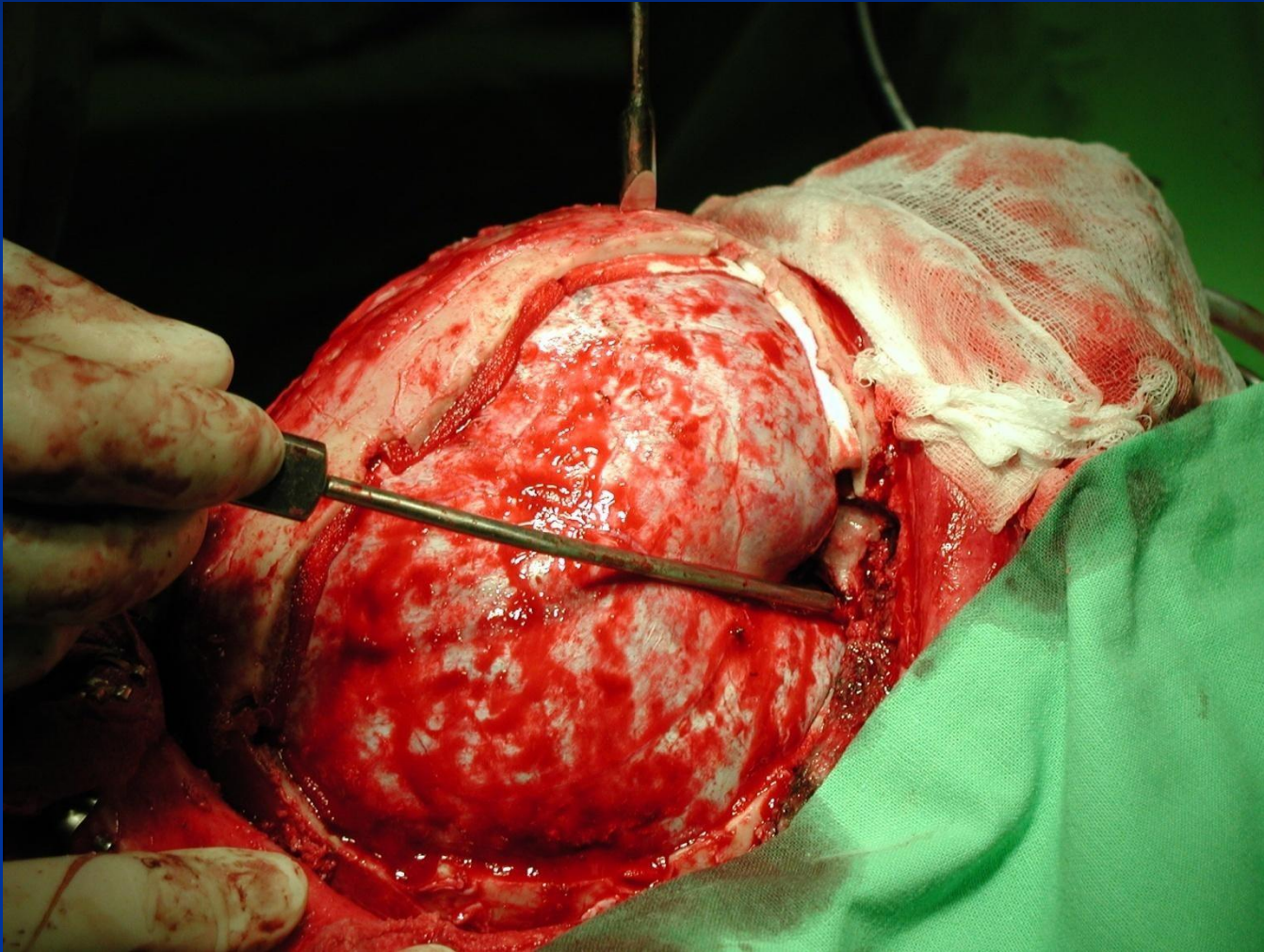


# Diffuse brain oedema

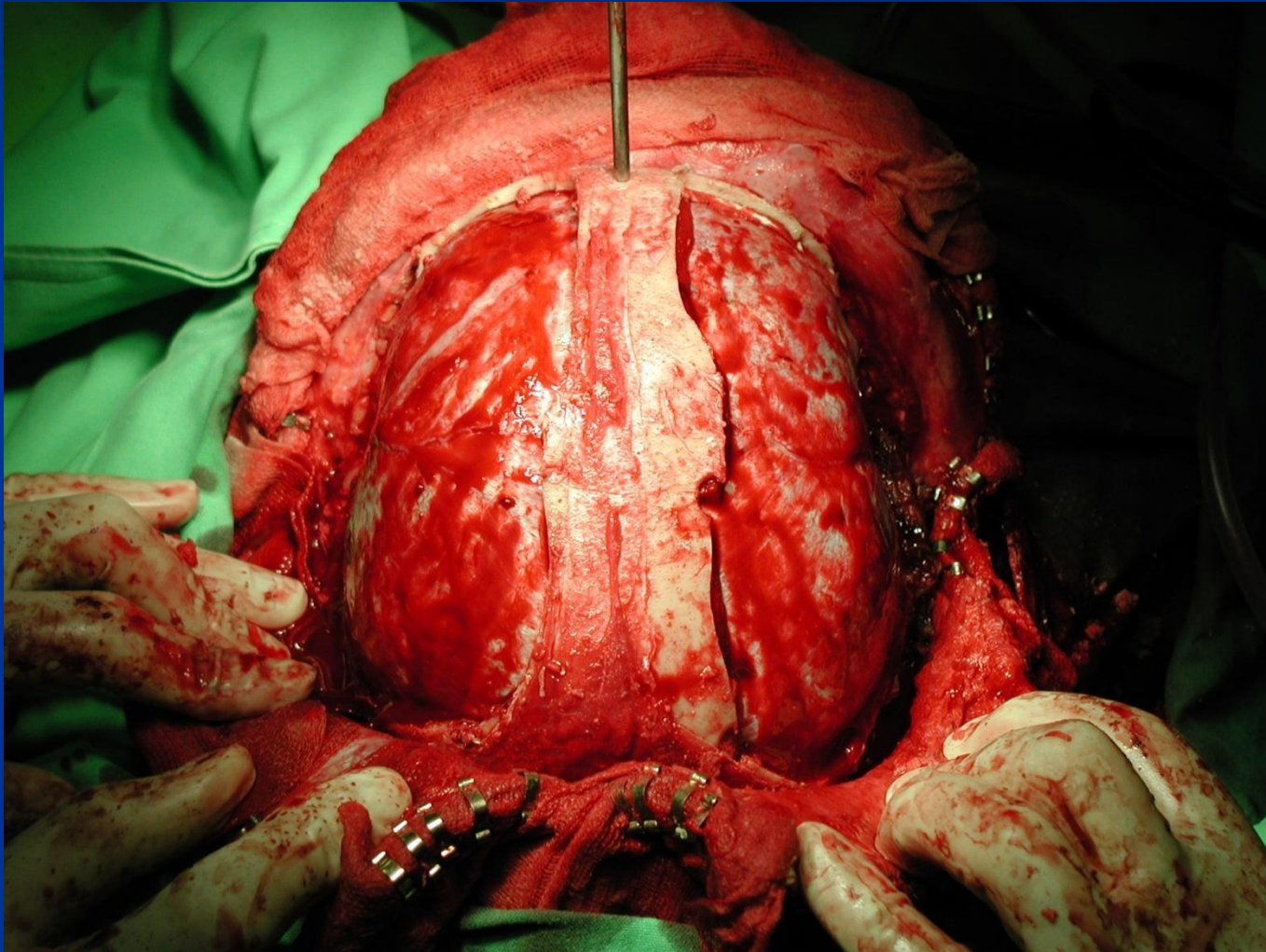




# Decompressive craniectomy

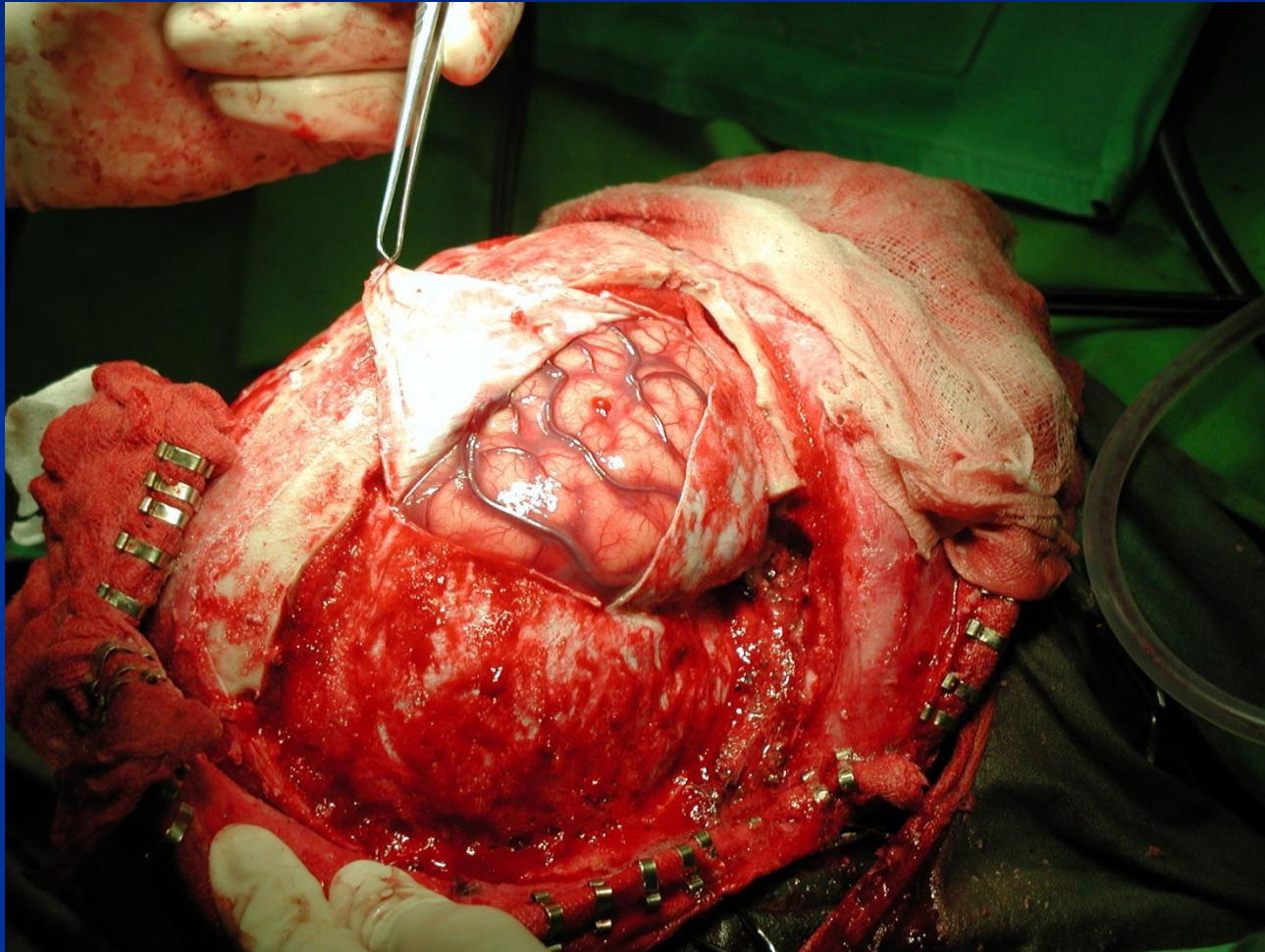


# Decompressive craniectomy

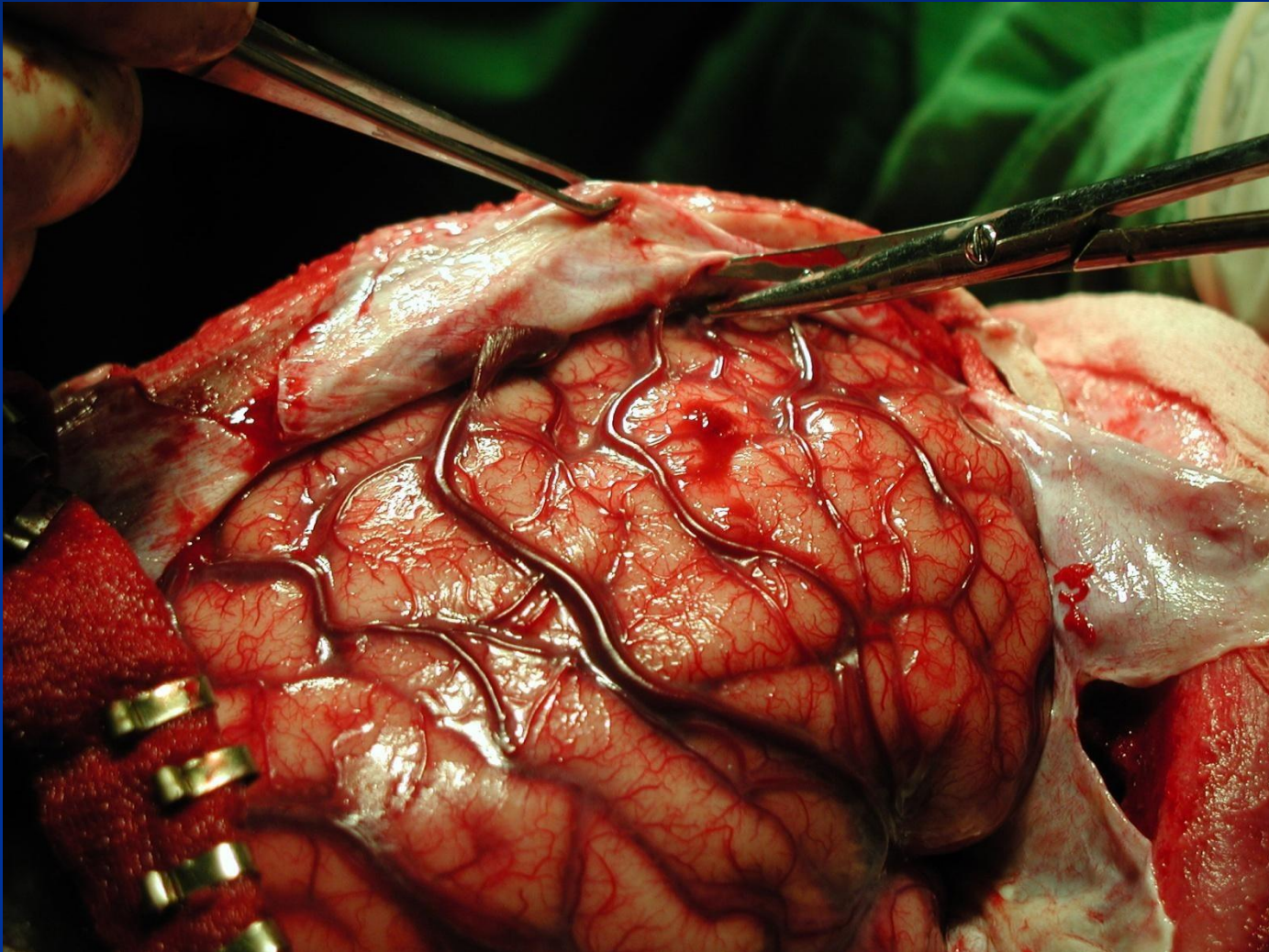




# Decompressive craniectomy

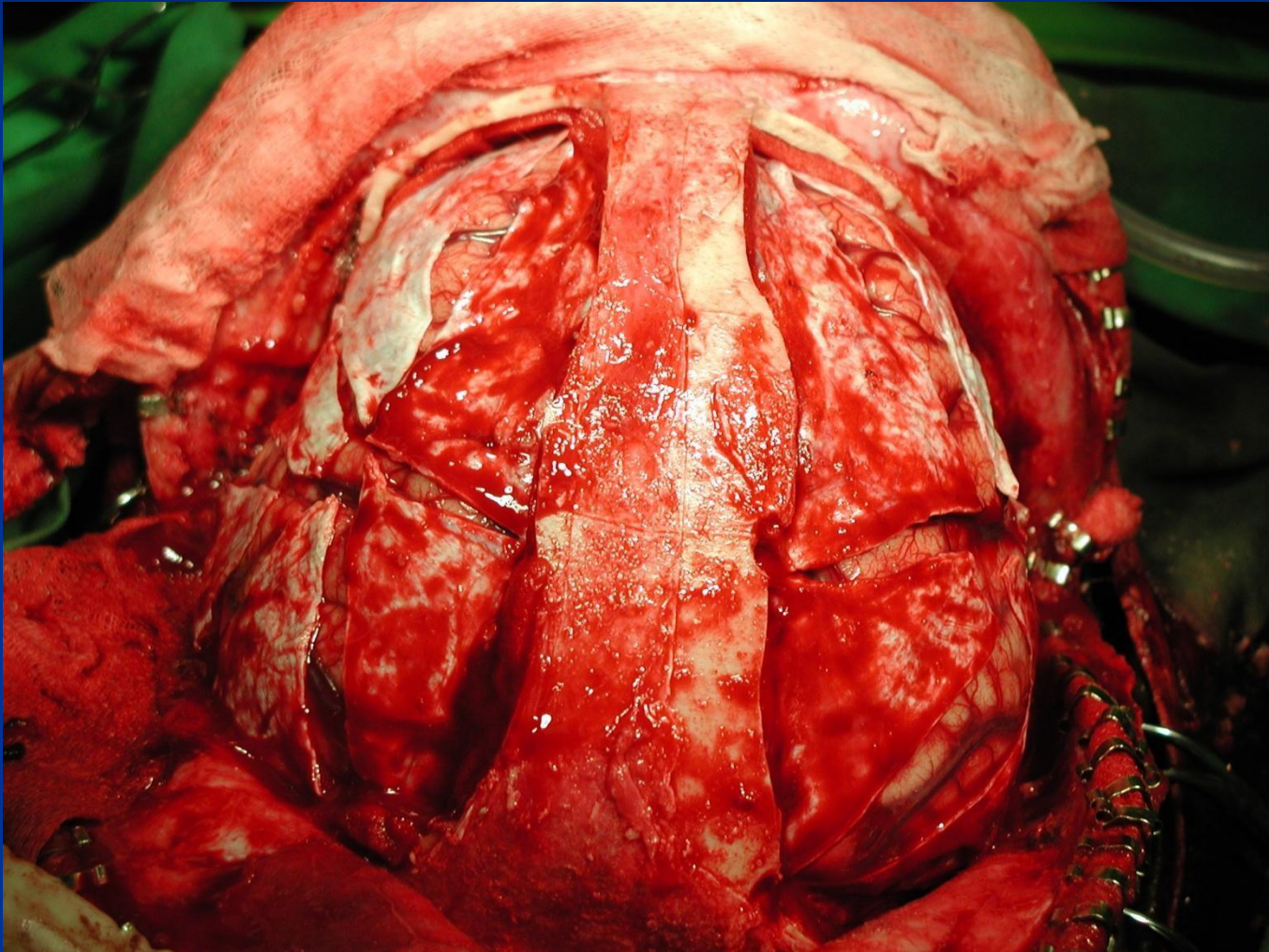


# Decompressive craniectomy





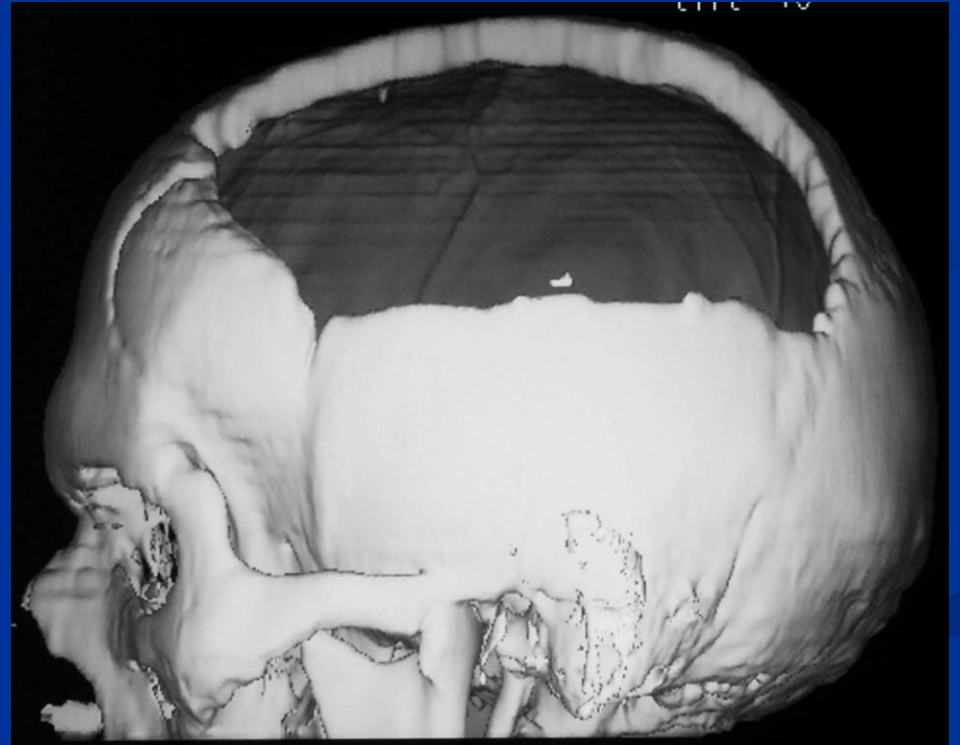
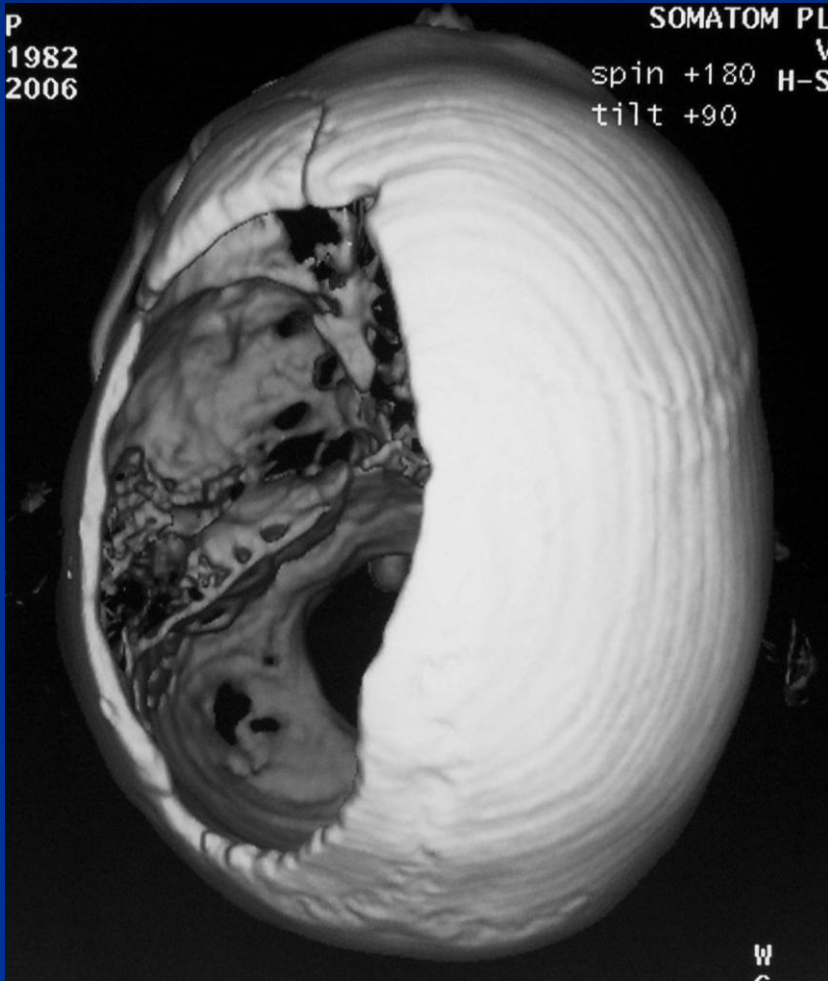
# Decompressive craniectomy



# Decompressive craniectomy

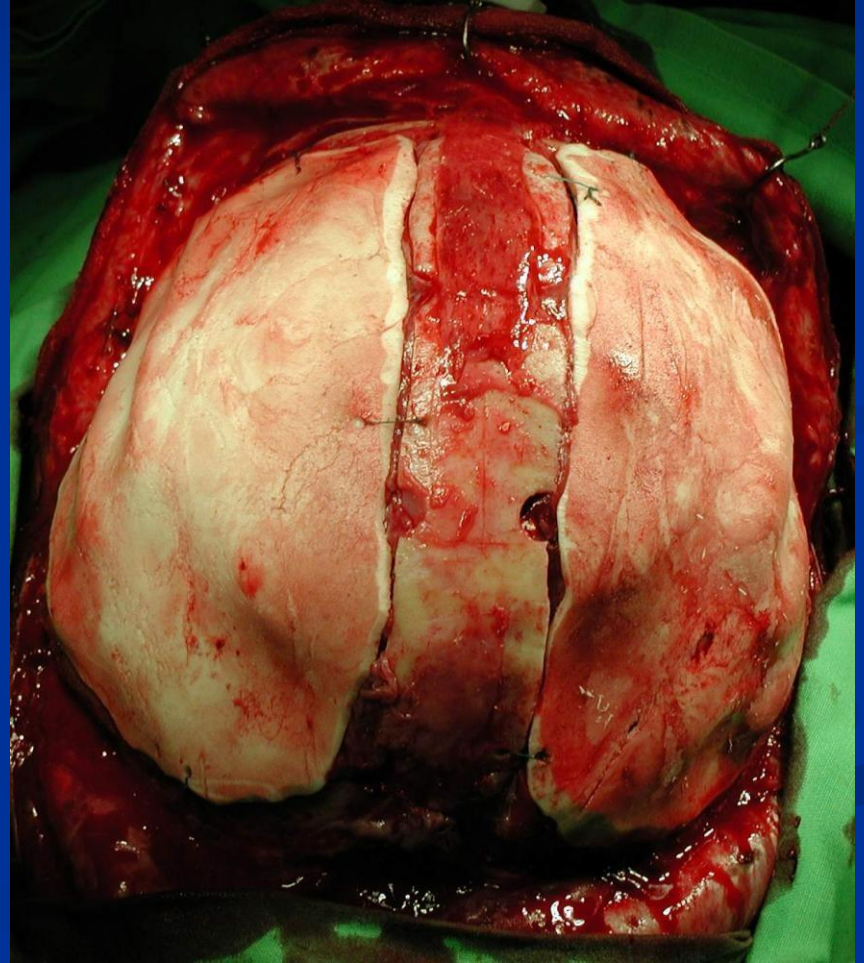


# Decompressive craniectomy





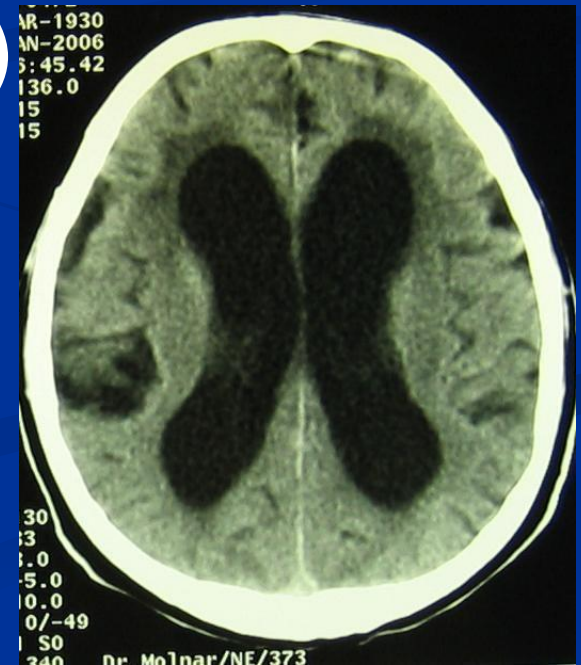
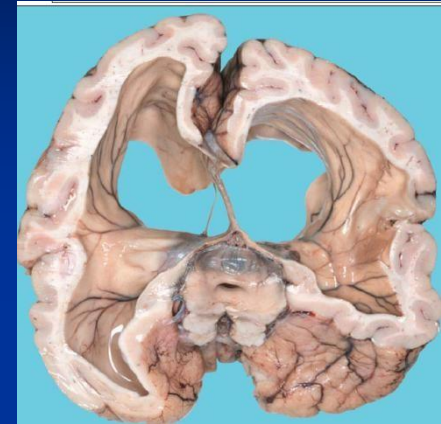
# Cranioplasty





# Posttraumatic hydrocephalus

- headache
- dementia
- ataxia (frontalis dysbasia)
- incontinentia urinae
  
- Treatment: Ventriculo-peritoneal shunt



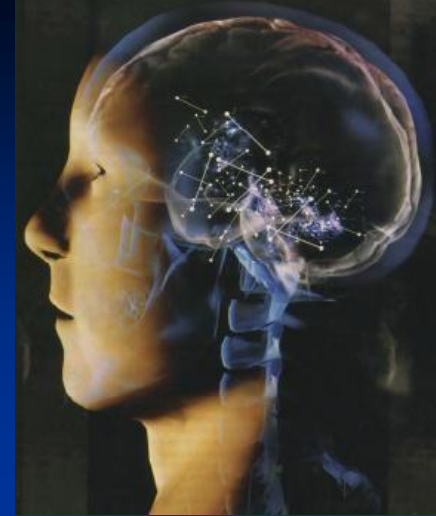
# Posttraumatic epilepsy



- <5% cases without loss of consciousness,
- 10-11% impressed bone fracture
- 10-14% loss of consciousness >24h
- 20-30% haemorrhagic contusion
- 45% IC haematoma

Treatment: carbamazepin, hydantoin

# The aim:



management of  
secondary brain injury to  
decrease of mortality



**KEEP  
CALM  
AND**

**THANK YOU FOR  
YOUR ATTENTION**

