

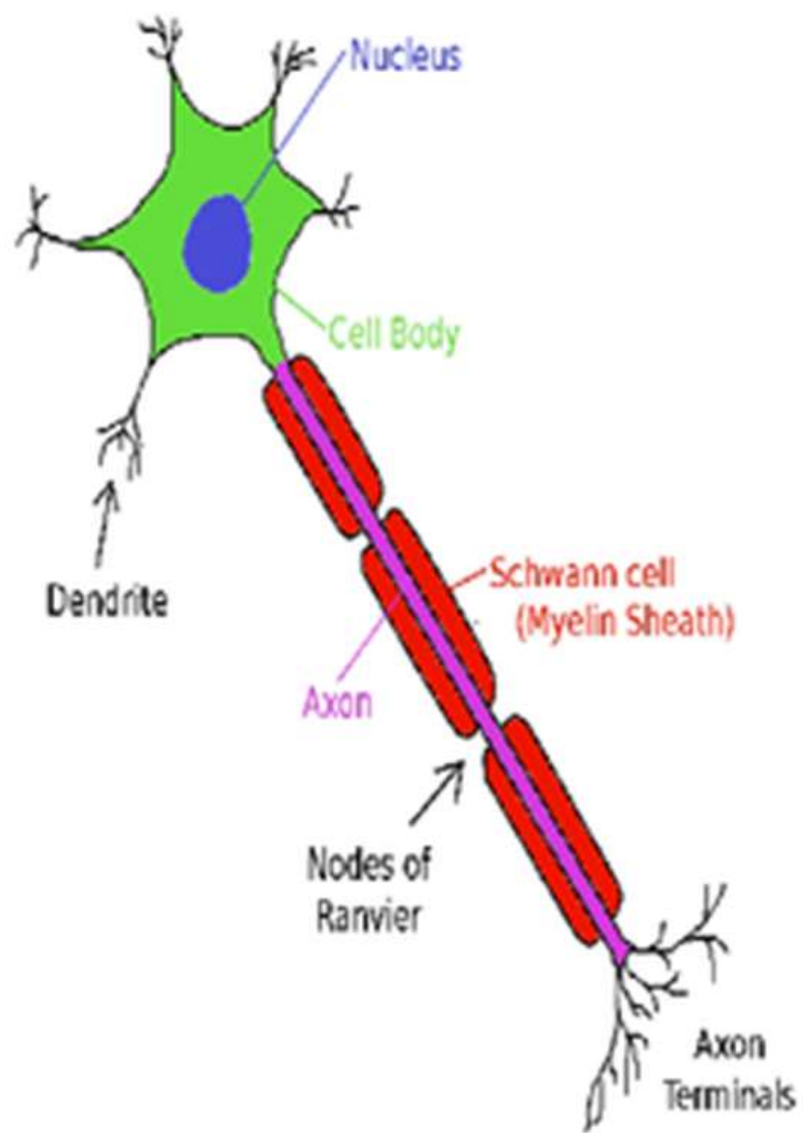
# **The Nervous System**

# Anatomy

# Introduction

- The nervous system consists from CNS and PNS
- CNS consists from brain and spinal cord
- PNS consists from somatic and autonomic nerves

- ▶ The neuron is the functional unit of the nervous system. Each neuron has a cell body and axon terminating at a synapse, supported by astrocytes and microglial cells.
- ▶ **Astrocytes** provide the structural framework for the neuron, control their biochemical environment and form the **blood–brain barrier**.
- ▶ **Microglial** cells are blood-derived mononuclear macrophages **with immune and scavenging functions**.
- ▶ In the CNS, myelin is produced by oligodendrocytes. In the PNS, myelin is produced by Schwann cells.



# Meninges

- Brain and spinal cord are covered with three membranous layers called the meninges: dura mater next to the bone, arachnoid and pia mater next to the nervous tissue.

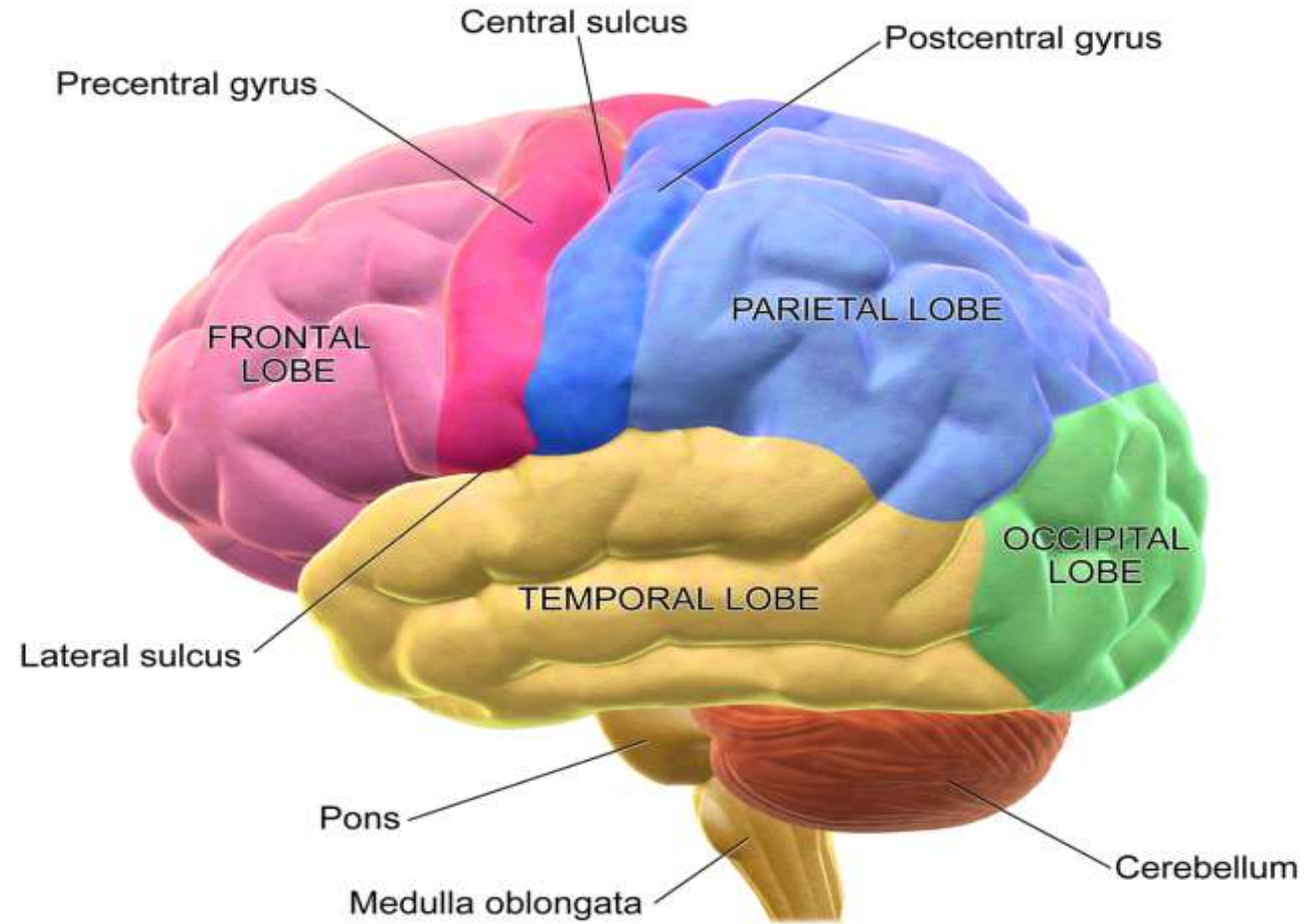
# CSF

- The subarachnoid space between the arachnoid and pia is filled with cerebrospinal fluid (CSF) produced by the choroid plexuses.
- The total volume of CSF is between 140 and 270 mL and there is a turnover of the entire volume 3–4 times a day
- Rate of production 700 mL per day

# Brain

- Two cerebral hemispheres, each with four lobes (frontal, parietal, temporal and occipital), the brainstem and the cerebellum.
- The brainstem comprises the midbrain, pons and medulla.
- The cerebellum has two hemispheres and a central vermis attached to the brainstem by three pairs of cerebellar peduncles

# Lateral View of the Brain

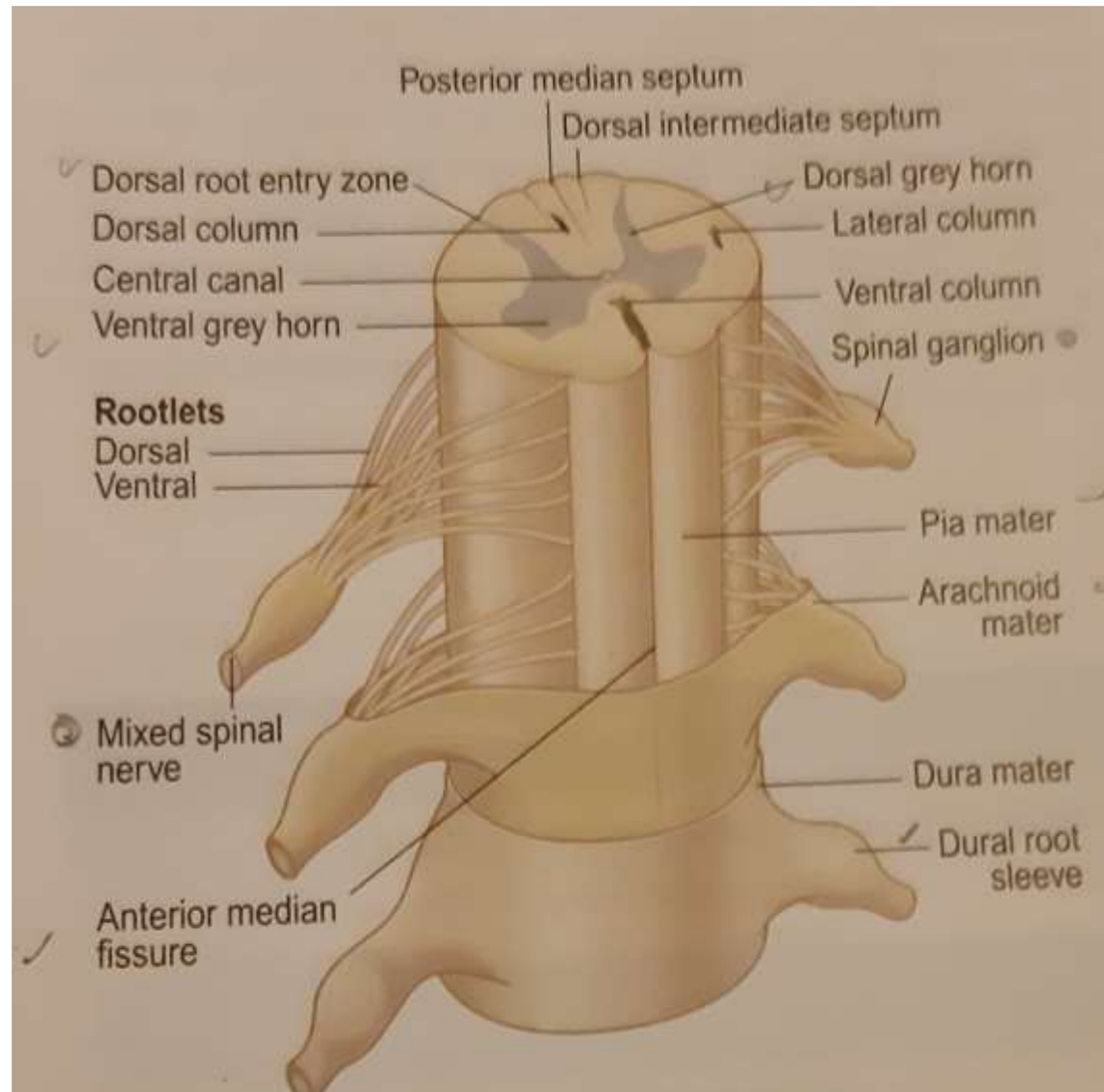


# Spinal cord

- The spinal cord is the main pathway for information connecting the brain and PNS.
- It contains the ventral grey horn and dorsal grey horn

- Ventral roots consist of efferent fibers that arise from motor neurons found in ventral grey horns.
- The dorsal roots are afferent fibers, receiving sensory information from organs to be transmitted to brain through sensory neurons found in dorsal grey horn.

# Nerve root and meninges



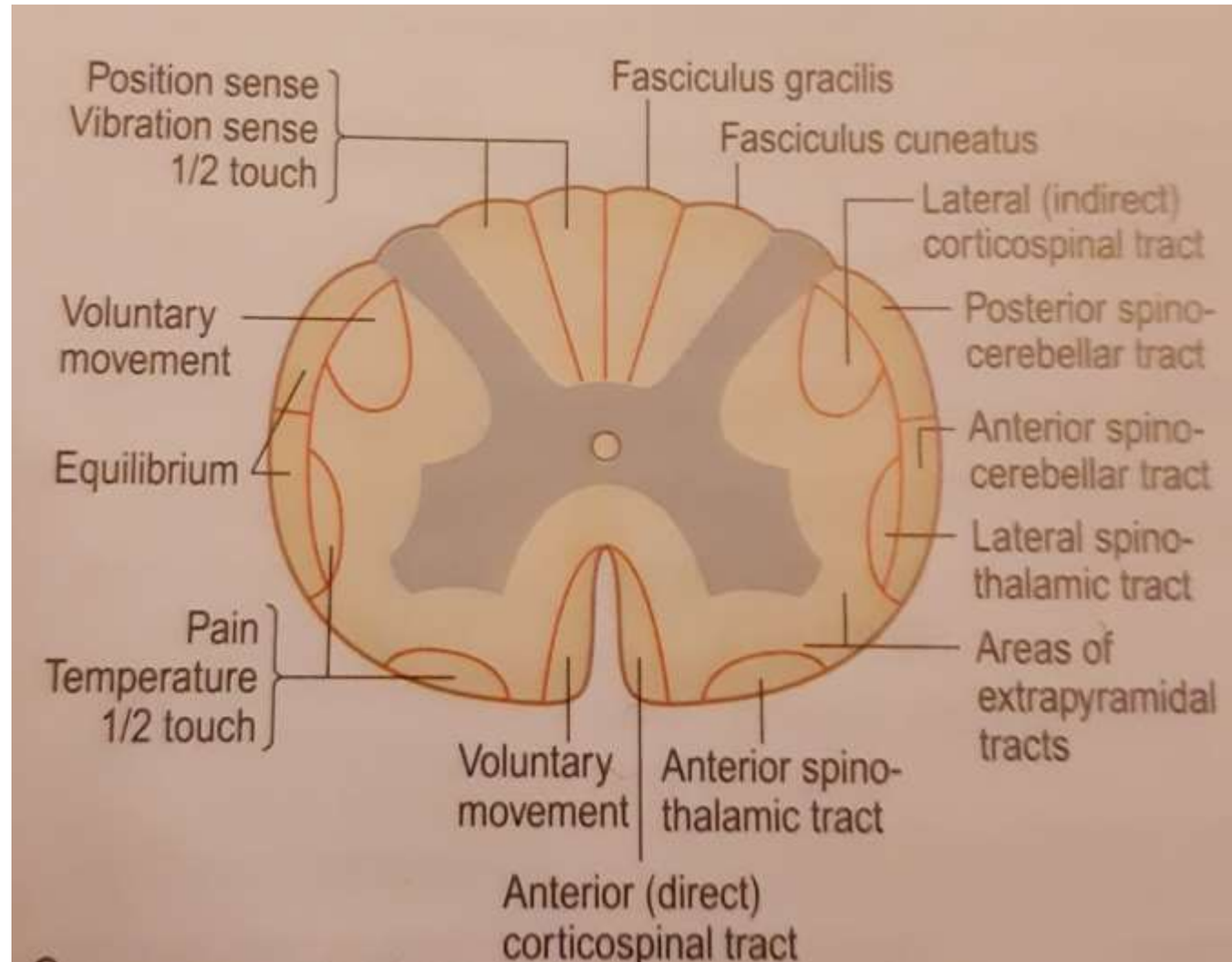
# Spinal cord

- The spinal cord contains multiple tracts
  1. Dorsal column (fasciculus gracilis and fasciculus cuneatus) : responsible for proprioception , vibration and half touch
  2. Anterior and lateral corticospinal tracts: voluntary movements

# Spinal cord

3. Anterior and lateral spinothalamic tracts: pain , tempertaure and half touch
4. Spinocerebellar tract: equilibrium

# Cross-section spinal cord



# Peripheral nervous system

- Peripheral nerves may have myelinated or unmyelinated axons
- Contains somatic and autonomic nerves
- Somatic nerves: consist of sensory and motor nerves

# History

- The history is a the key for diagnosis as physical exam maybe normal or unhelpful.
- In cases of amnesia or loss of consciousness we need additional witness history.
- We should clarify exactly what the patient means by any neurological symptom.
- Ask the patient what they fear might be wrong.

- For any neurological symptom ask about onset , duration , pattern , exacerbating , relieving factors and associated symptoms.

# Symptoms

# 1- Headache

- May be either primary or secondary.
- Use SOCRATES to analyze it
- Isolated headache with a truly abrupt onset may represent a potentially serious cause such as subarachnoid haemorrhage or cerebral vein thrombosis, whereas recurrent headache is much more likely to be benign primary headache.

# Primary (idiopathic) causes

1. migraine
2. tension headache
3. trigeminal autonomic cephalalgias (including cluster headache)
4. primary stabbing headache
5. cough, exertional or sex headache
6. primary thunderclap headache

# Migraine

- Recurrent attacks of moderate to severe headaches
- Mostly unilateral
- Evolve over 30 minutes to 2 hours , lasting up to 72 hours, with weeks to months of symptoms free.
- Classified into classic (with aura) and common (without aura)

# Migraine

- Associated with nausea/vomiting, photophobia/phonophobia
- During the attack the patient prefers to be in a dark room .

# Cluster headaches

- Abrupt onset , Attacks last up to 2 hours
- Recurrent attacks 1-4 times within 24 hours , lasting weeks to months , with months to years of remission
- Awake the patient from sleep
- Orbital/retro-orbital; always same side during cluster, may switch sides between clusters

# Cluster headache

- Autonomic features, including conjunctival injection, tearing, nasal stuffiness, ptosis, miosis, agitation
- During the attack the patient keep pacing around the room in an agitated state, or even head banging

# Stabbing headache

- Abrupt onset
- Last very briefly , seconds or less
- Anywhere
- Common in patients with migraine

# Secondary (or symptomatic) headaches

- **Meningitis:** associated with neck stiffness, fever, rash, signs of raised intracranial pressure and false localizing signs, meningism
- **Subarachnoid hemorrhage:** Abrupt onset, maybe fatal at onset , associated with nausea/ vomiting, reduced consciousness, false localising signs, III nerve palsy
- **Temporal arteritis:** usually occur in patients more than 55 yrs, presents with jaw pain on chewing, visual symptoms , and tender temporal arteries, elevated ESR and CRP

## 2- Disturbances of consciousness

### **\*\*\*Causes:**

- Postural hypotension
- Neurocardiogenic syncope (vasovagal)
- Hypersensitive carotid sinus syndrome (pressure over carotid sinus may lead to reflex bradycardia and syncope)
- Cardiac syncope due to arrhythmias or mechanical obstruction of cardiac output

## ✓ Syncope

- Most common cause of transient loss of consciousness
- Due to inadequate cerebral perfusion
- Maybe due to vasovagal(reflex) or cardiac syncope

# Vasovagal syncope

- ▶ It occurs due to stimulation of parasympathetic system due to pain , emotion or illness or in people forced to stand in warm environment. Leads to vasodilation and bradycardia
- ▶ Often preceded by light-headedness, vision dimming, tinnitus, and nausea
- ▶ Lasts 1-2 minutes
- ▶ It causes pale or grey skin

- Maybe associated with myoclonic jerks
- If kept flat, recovery is rapid

# Cardiac syncope

- Syncope with no previous alarm or trigger or exercise
- Causes: hypertrophic cardiomyopathy, severe aortic stenosis or arrhythmia.

## ✓ **Postural hypotension**

Could be due to

1- drugs (levodopa or anti hypertensive drugs) or

2- autonomic diseases such as DM

3- in people more than 65 years

4- hypovolemia

# How to ask about syncope ??

- Ask about witness
- any preceding symptoms (palpitation, chest pain, lightheadedness, nausea, tinnitus, sweating and visual disturbance)
- Duration of loss of consciousness
- Appearance of the patient while unconscious
- Any injuries sustained.
- Time to recovery to full consciousness and normal cognition

# 3- Epileptic Seizures

- paroxysmal electrical discharges from either the whole brain (generalized) or part of the brain (focal).
- The history from the **patient** and **witnesses** can help distinguish epilepsy from syncope
- Usually triggered by sleep deprivation or alcohol or drugs
- Types
  1. Generalized: tonic–clonic seizure is the most common form
  2. Focal (partial)

# Tonic clonic seizure

- Tonic phase: typically follows a **stereotyped** pattern with early loss of consciousness associated with body stiffening
- clonic phase: rhythmical jerking crescendoing and subsiding over 0.5 – 2 minutes
- postictal phase: period of unresponsiveness often with heavy breathing, the patient appears to be deeply sleep and finally confusion as the patient awakes.

# How to differentiate epilepsy from vasovagal??

- Trigger: alcohol , sleep deprivation or drugs
- Prodrome: May have focal (aura) prodrome
- Convulsion and loss of consciousness lasts for 1-2 minues
- Full recovery occurs over 30 minutes

- Cyanosed skin
- Lateral tongue biting, headache, generalized myalgia, back pain (vertebral compression fractures), shoulder fracture/ dislocation may occur

# Focal seizure

- Simple (consciousness is preserved) or complex (impaired consciousness)
- Characterized by whichever part of the brain is involved
- frontal lobe seizures: focal motor seizure
- temporal lobe seizures characterised by autonomic and/or psychic symptoms, often associated with automatisms such as lip smacking or swallowing.

# Functional dissociative attacks (non-epileptic or psychogenic attacks or pseudoseizures)

- difficult to distinguish from epileptic seizures, clues to differentiate psychogenic seizures:
  1. occurring multiple times in a day
  2. may last considerably longer
  3. symptoms waxing and waning

4. asynchronous movements

5. pelvic thrusts , side-to-side rather than flexion/extension movements

6. absence of postictal confusion

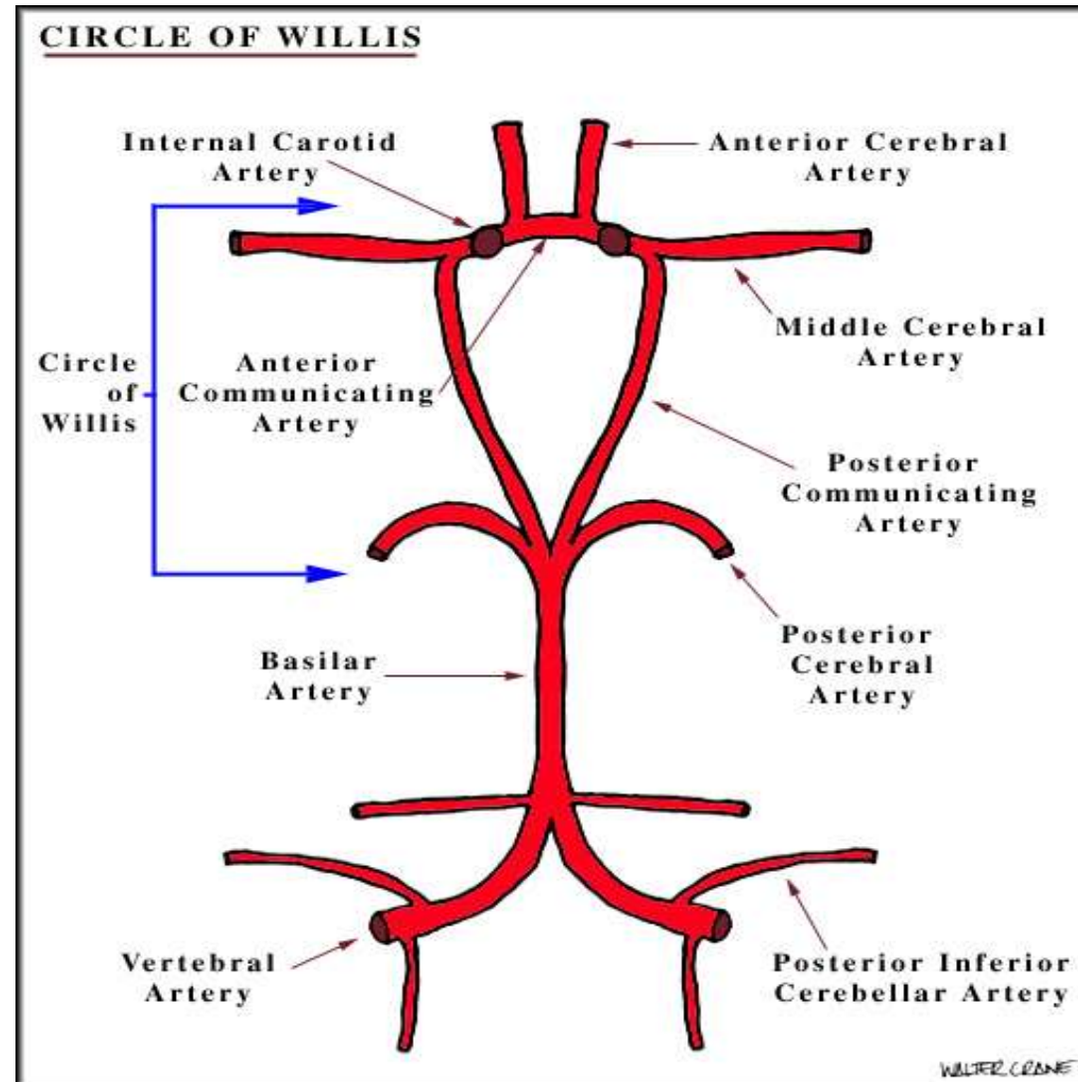
- The widespread availability of videophones allows witnesses to capture such events

# 4- Stroke symptoms

- STROKE is a focal neurological deficit of rapid onset that is due to a vascular cause , maybe ischemic or hemorrhagic
- A transient ischaemic attack (TIA) is the same but symptoms resolve **within 1 hour**.
- TIAs are an important risk factor for impending stroke and demand urgent assessment and treatment.

- Symptoms are dictated by the vascular territory involved
- **Much of the cerebral hemispheres** are supplied by the **anterior circulation** (the anterior and middle cerebral arteries which are derived from the internal carotid artery) , while **the occipital lobes and brainstem** are supplied by the **posterior circulation** (posterior cerebral artery which is derived from vertebrobasilar circulation)

# Arterial blood supply of brain



# Ischemic and hemorrhagic stroke

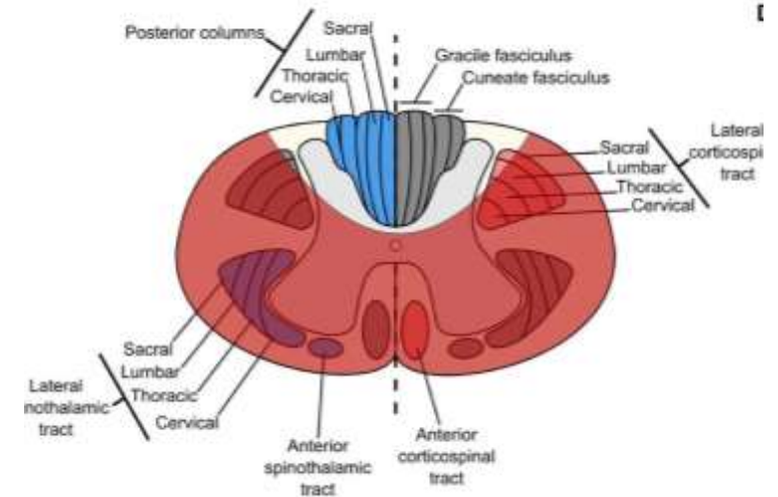
- 80% of strokes are ischemic
- Hemorrhagic stroke is much more frequent in Asian populations.
- Factors in the history or examination that increase the likelihood of haemorrhage include: use of anticoagulation, headache, vomiting, seizures and early reduced consciousness.

- **We have to do brain CT without contrast to differentiate between them**
- Isolated vertigo, amnesia or TLOC are rarely, if ever, due to stroke

# Spinal strokes

Spinal strokes are very rare; patients typically present with abrupt onset, depending on the level of spinal cord affected.

The anterior spinal artery syndrome is most common and causes loss of motor function and pain/temperature sensation, with relative sparing of joint position and vibration sensation below the level of the lesion (sparing dorsal column)



# Clinical classification of stroke

- **Total anterior circulation syndrome (TACS)**  
Hemiparesis plus  
hemianopia and  
higher cortical deficit (e.g. dysphasia or visuospatial loss)
- **Partial anterior circulation syndrome (PACS)**
  - 1-Two of the three components of a TACS
  - 2-OR isolated higher cortical deficit
  - 3- OR motor/sensory deficit more restricted than LACS

- Posterior circulation syndrome (POCS): one of the following:

1- Ipsilateral cranial nerve palsy with contralateral motor and/or sensory deficit

2-bilateral motor and/or sensory deficit

3-disorder of conjugate eye movement

4-cerebellar dysfunction without ipsilateral long-tract deficits

5-isolated homonymous visual field defect

- Lacunar syndrome (LACS):

ONE OF:

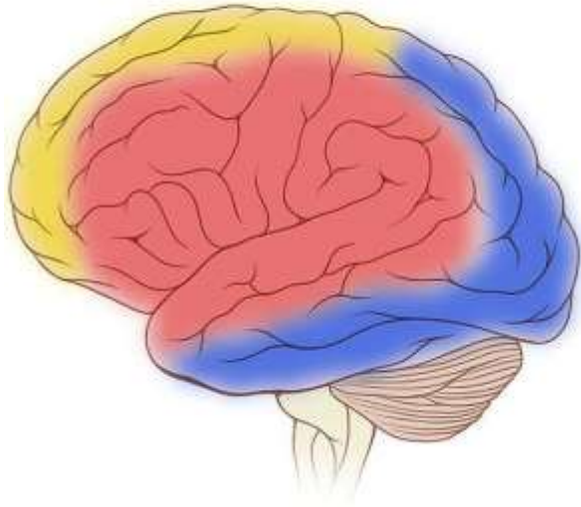
1- Pure motor > 2 out of 3 of face, arm, leg

2- Pure sensory > 2 out of 3 of face, arm, leg

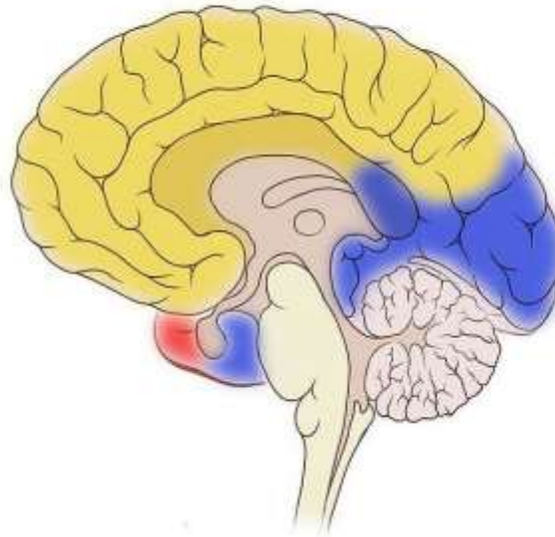
3- Pure sensorimotor > 2 out of 3 of face, arm, leg

4- ataxic hemiparesis

Lateral Brain



Medial Brain



-  Anterior Cerebral Artery
-  Middle Cerebral Artery
-  Posterior Cerebral Artery

# 5-Dizziness and Vertigo

## ✓ Dizziness :

- Recurrent dizzy spells affect approximately 30% of those over 65 years.

- Causes

1-postural hypotension

2-cerebrovascular disease

3-cardiac arrhythmia

4-hyperventilation induced by anxiety and panic.

## ✓ Vertigo:

- the illusion of movement
- specifically indicates a problem in the vestibular apparatus (most common) or the brain (central)

# Peripheral causes of vertigo

- **Benign paroxysmal positional vertigo (BPPV):** recurrent episodes of vertigo lasting a few seconds , attacks increased when sleeping on the affected side or with movement
- **Meniere disease:** vertigo lasting minutes or hours, associated with hearing loss, tinnitus, nausea and vomiting

# Central causes of vertigo

- Migrainous vertigo (with or without headache)
- Stroke
- Multiple sclerosis
- **TIAs do not cause isolated vertigo.**

# Functional/psychogenic/hysterical/ somatisation/conversion disorder

- Not due to a true neurological disease
- Presentations include blindness, tremor, weakness and collapsing attacks, and patients will often describe numerous other symptoms, with fatigue, lethargy, pain, anxiety and other mood disorders commonly associated.

# Continue

- Clues include

- 1- symptoms not compatible with disease (such as retained awareness of convulsing during non-epileptic attacks, or being able to walk normally backwards but not forwards)
- 2- considerable variability in symptoms (such as intermittent recovery of a hemiparesis)

3- **multiple symptoms** with numerous visits to other specialties and **multiple unremarkable investigations**, leading to numerous different diagnoses

- Beware of labeling symptoms as functional simply because they appear odd.
- Most functional neurological disorders follow recognizable patterns, so be cautious when the pattern is atypical.

# Past medical history

- History of previous visual loss (optic neuritis) in someone presenting with numbness suggests multiple sclerosis.
- Birth history and development may be significant, as in epilepsy.
- If considering a vascular cause of neurological symptoms, ask about important risk factors, such as other vascular disease, hypertension, family history and smoking.

# Drug history

- **Always enquire about drugs, including prescribed, over-the counter, complementary and recreational/illegal ones**
- phenytoin toxicity causes ataxia
- excessive intake of simple analgesia causing medication overuse headache; use of cocaine provoke convulsions.

# Family history

- Parental consanguinity is common, increasing the risk of autosomal recessive conditions
- Single-gene defects: such as myotonic dystrophy or Huntington's disease.
- Polygenic influences, as in multiple sclerosis or migraine.

- Charcot–Marie–Tooth disease may be autosomal dominant, autosomal recessive or X-linked.
- **Mitochondria** uniquely have their **own DNA**, and abnormalities in this DNA can cause a range of disorders that manifest in many different systems (such as **diabetes, short stature** and **deafness**), and may cause common neurological syndromes such as **migraine** or **epilepsy**.

- Some diseases, such as Parkinson's or motor neuron disease, may be either due to single-gene disorders or sporadic

# Social history

- How are patients coping with their symptoms? Are they able to work and drive?
- What are their support circumstances, and are these adequate?
- Ask about **alcohol** as it affects CNS (ataxia, seizures, dementia) and PNS (neuropathy)

- Ask about **diet**
  - **Vitamin deficiency** may occur in alcoholism or dietary exclusion
  - Vegetarians may be susceptible to vitamin B12 deficiency (subacute combined degeneration of the spinal cord)
- Ask about **recreational drugs**
  - **nitrous oxide inhalation** causes subacute combined degeneration of the cord due to dysfunction of the vitamin B12 pathway
  - **smoking** contributes to vascular and malignant disease.

- A **travel** history may give clues to the underlying diagnosis such as:
  - Lyme disease (facial palsy)
  - neurocysticercosis: parasitic infection (brain lesions and epilepsy)
  - malaria (coma)
- Always consider **sexually transmitted or blood-borne infection**, such as human immunodeficiency virus (HIV) or syphilis, as both can cause a wide range of neurological symptoms and are treatable

# Occupational history

- lead exposure :motor neuropathy
- manganese causes Parkinsonism.
- Some neurological diagnoses may adversely affect occupation, such as epilepsy in anyone who needs to drive or operate dangerous machinery

- For patients with cognitive disorders, particularly dementias, it may be necessary to patients to stop working.

**Thank you**