

# SPASTICITY AND INCREASED MUSCLE TONE

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Note: Pure corticospinal tract lesion cause hypotonia instead of spasticity The reason is that pure pyramidal tract lesion is very very rare, and spasticity is due to loss of inhibitory control of extrapyramidal tracts.

# **OBJECTIVES**

At the end of this lecture you should be able to

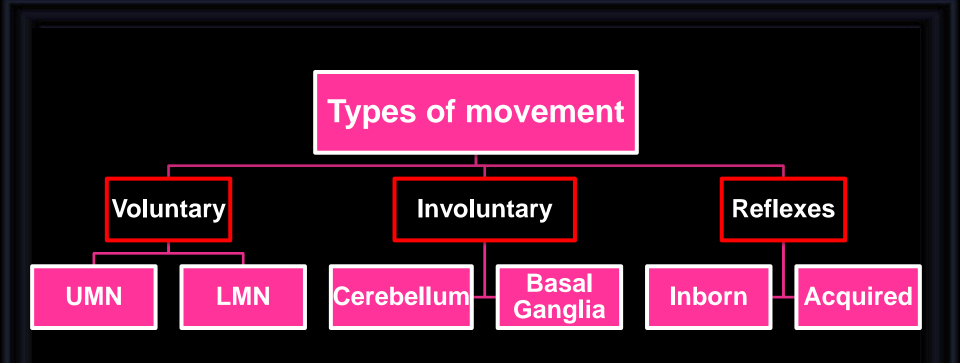
- Define spasticity ,rigidity and dystonia
- Describe the neurophysiology of spasticity
- Describe the features of upper and lower motor neuron lesions
- Describe the causes of spaticity
- Enumarate the effects of spasticity
- List the treatments to reduce spasticity.



## **MOTOR SYSTEM LAYOUT**

Motor = movement
For movement we need
nervous system and muscles.
Components of motor system are:

- Motor cortex
- Upper and lower motor neurons
- Cerebellum
- Basal ganglia
- Spinal cord
- Muscles



#### **INITIATOR OF MOTOR ACTIVITY**

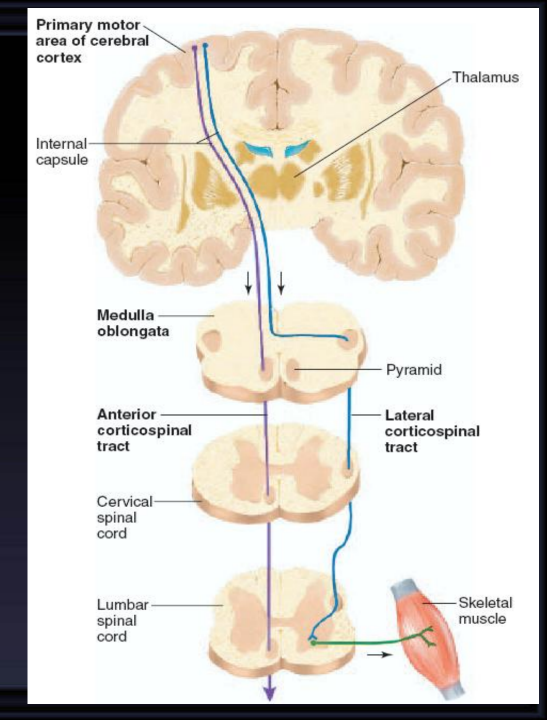
**Corticospinal tract (Pyramidal tract)** 

#### **REGULATOR OF MOTOR ACTIVITY**

Cerebellum (Extrapyramidal tracts)

#### PYRAMIDAL TRACTS

- Originating Cells: Giant pyramidal cells
- \*80% to 90% of the pyramidal fibers then cross in medulla to the opposite side and descend into the lateral corticospinal tracts
- Anterior or ventral corticospinal tract is formed by uncrossed fibers are primarily concerned with the control of fine movements that require dexterity.



# MUSCLE TONE → ↓

The resistance of a muscle to stretch is often referred to as its tone or tonus.

If the motor nerve to a muscle is severed, the muscle offers very little resistance and is said to be flaccid.

A hypertonic (spastic) muscle is one in which the resistance to stretch is high because of hyperactive stretch reflexes

# COMAPRISON BETWEEN UPPER & LOWER MOTOR NEURON LESIONS

#### <u>UMN LESION</u>

- Paralysis affect movements
- Wasting not pronounced.
- Spasticity Muscles hypertonic (Clasp Knife).
- Tendon reflexes increased.
- Superficial reflexes diminished
- Babinski's sign +ve,

#### **LMN LESION**

- Individual muscle or group of muscles are affected.
- Wasting pronounced.
- Flaccidity. Muscles hypotonic.
- Tendon reflexes diminished or absent.
- Superficial reflexes often unaltered.

# What is Spasticity?

"Spasticity is a motor disorder characterized by a velocity-dependant increase in tonic stretch reflexes with exaggerated tendon jerk, resulting from hyperexcitability of the stretch reflex"

James Lance (1980)

In UMN lesions Spasticity is of Clasp Knife Type

Rigidity is increased neural activity throughout the range of muscle movement and is not velocity dependent. Rigidity is present in both agonist and antagonist muscles. It is often associated with basal ganglia disease such as Parkinson's disease

### Rigidity in Parkinsonism

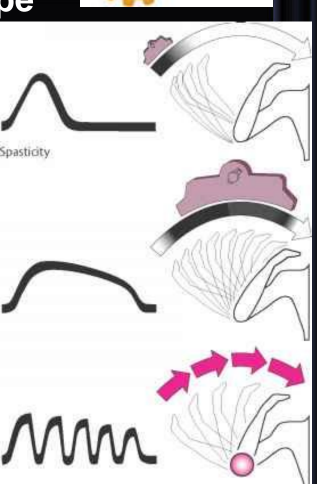
Lead-pipe rigidity. Passive movement of an extremity meets with a constant dead feeling resistance like a lead pipe throughout the range of movement.

Sometimes a series of "catches" takes place during passive motion like a cogwheel

### Cog-wheel rigidity



In cogwheel rigidity one feels that resistance varies rhythmically when applying a passive movement. It is because of an underlying resting tremor associated with rigidity.

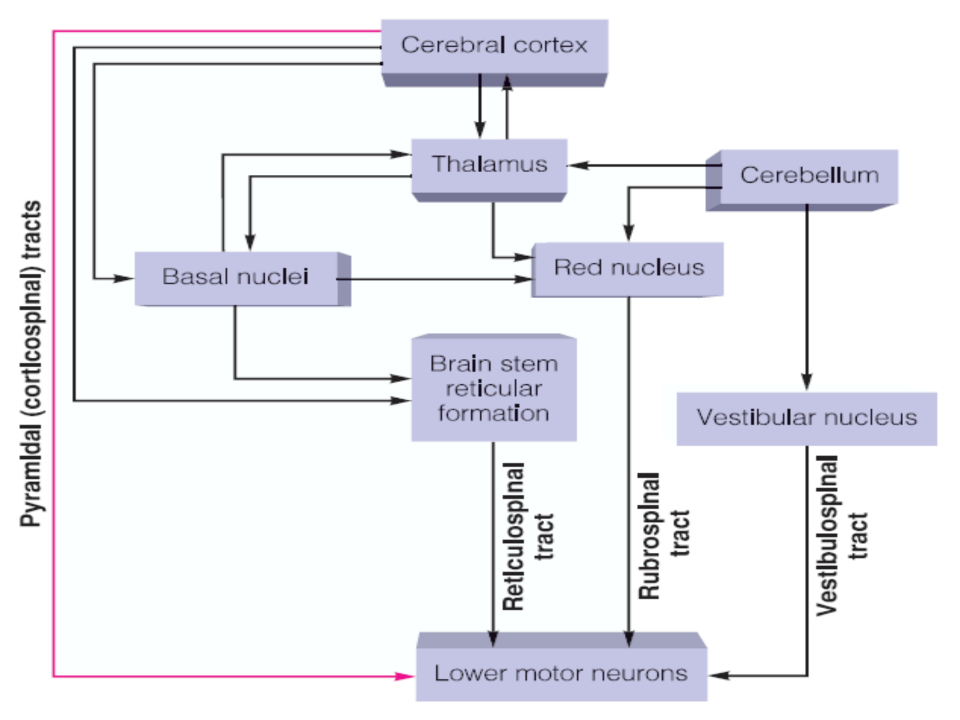


### **CAUSES OF SPASTICITY**

#### A-(UMNS) syndrome include:

- (1) Cerebral palsy
- (2) Stroke
- (3) Spinal cord injury
- (4) Multiple Sclerosis
- (5) Acqiured brain injury (trauma, etc)
- **B-Parkinsonism**
- C- Decerebrate & decorticate rigidity

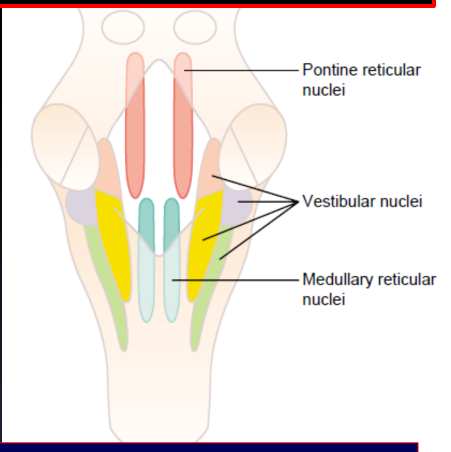
Patients complain of pain, stiffness & inability to relax
Prolonged stiffness leads to bone & joint deformities with disability
and contractures & its consequences



# Support of Body Against Gravity is the role of the Reticular and Vestibular Nuclei

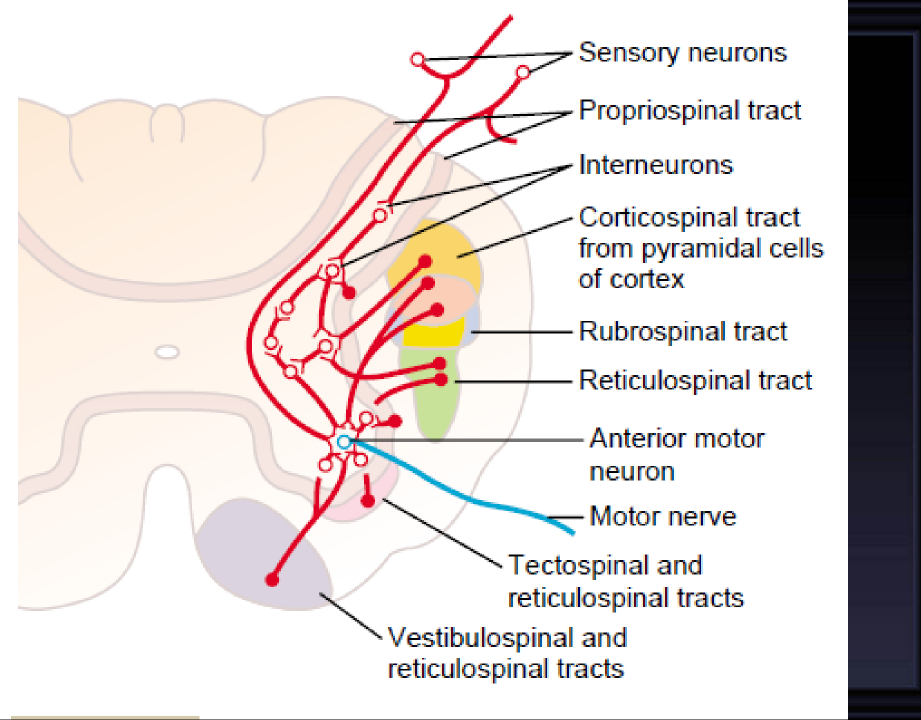
Pontine Reticular System. Transmit excitatory signals downward into the cord through the pontine reticulospinal tract in the anterior column of the cord

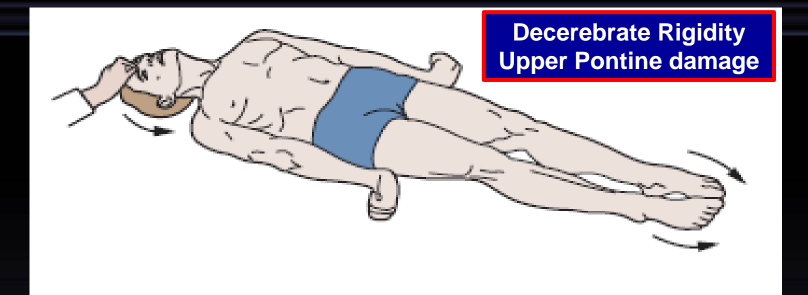
Medullary Reticular System.
Transmit inhibitory signals to the same antigravity anterior motor neurons by way of a different tract, the medullary reticulospinal tract, located in the lateral column of cord



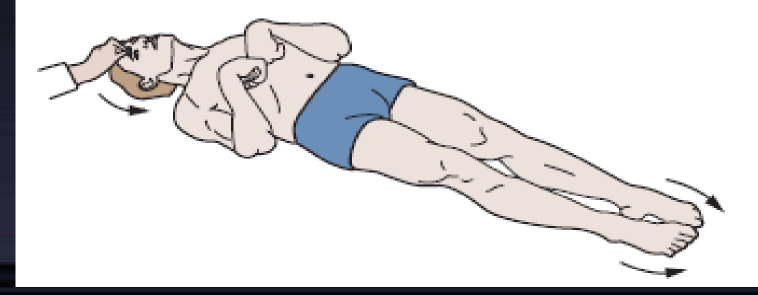
Decerebrate rigidity results from blockage of normally strong input to the medullary reticular nuclei from the cerebral cortex, the red nuclei, and the basal ganglia.

Medullary reticular inhibitor system becomes nonfunctional Pontine excitatory system becomes overactive





# Decorticate Rigidity Upper midbrain damage

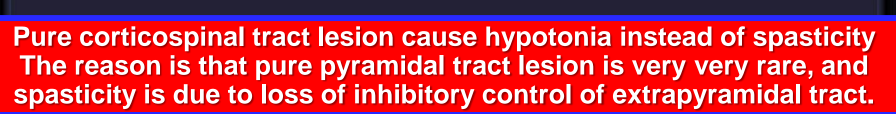


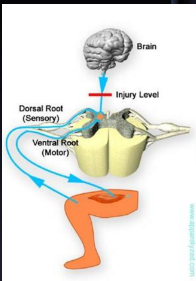
# **Mechanism of Spasticity**

- The absence of an upper motor neuron's inhibitory control on the spinal reflex. Loss of descending inhibition from the brain to BRAIN STEM EXCITATORY CENTERS (pontine RF + vestibular N).
- Vestibulospinal & reticulospinal EXCITATORY signals to muscles to become spastic
- Hyperactive stretch reflexes that are mediated by muscle spindle stretch receptors
- Decreased threshold of the alpha lower motor neurons

# Mechanism of Spasticity

- When Central Nervous System Lesion occurs
  - Hypotoncity
  - Alpha Lower motor neuron
- Days to weeks later
  - Denervation hypersensitivity
    - Alpha lower motor neurons lower threshold
    - An increase in the number of Ach receptors





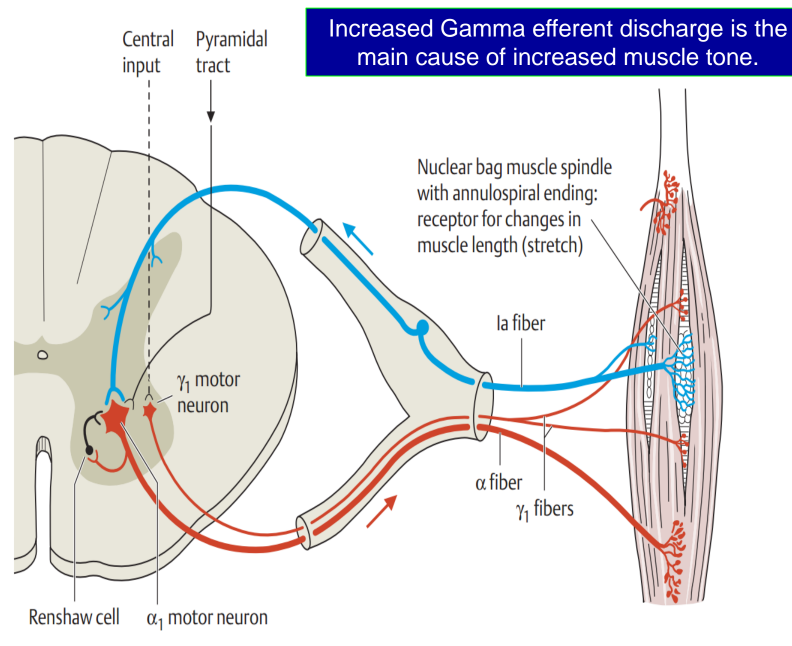
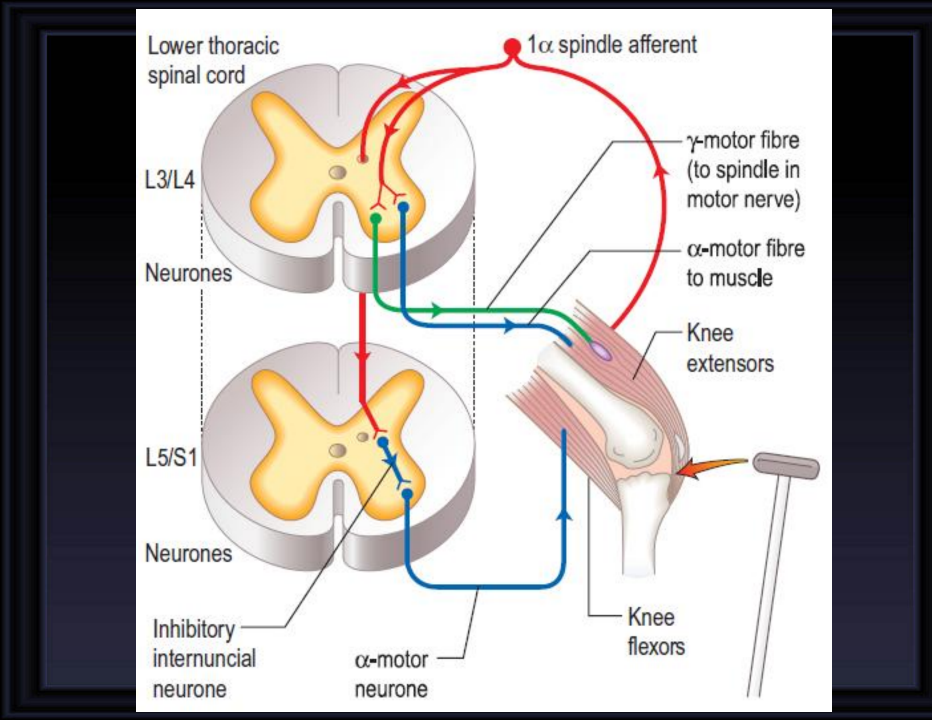


Fig. 2.11 Regulatory circuit for muscle length



#### Table 21.17

# Spinal levels, recording/interpretation of tendon reflexes

Level	Reflex	Symbol for reflex	Meaning
C5-6	Supinator	0	Absent
C5-6	Biceps	+/_	Present with reinforcement
C7	Triceps	+	Normal
L3-4	Knee	++	Brisk, normal
S1	Ankle	+++	Exaggerated (abnormal)
		CL	Clonus

## **Ashworth Scale**

- Physical therapist need clarity to realize how spasticity can have affects on a patients ability to move
- A valid & reliable measure scale for spasticity

#### **Modified Ashworth Scale**

- 0 No increase in muscle tone
- Slight increase in muscle tone, manifested by a catch and release or by minimal resistance at the end range of motion when the part is moved in flexion or extension/abduction or adduction, etc.
- 1+ Slight increase in muscle tone, manifested by a catch, followed by minimal resistance throughout the remainder (less than half) of the ROM
- 2 More marked increase in muscle tone through most of the ROM, but the affected part is easily moved
- 3 Considerable increase in muscle tone, passive movement is difficult
- 4 Affected part is rigid in flexion or extension (abduction, adduction, etc.)

# **Spasticity vs Dystonia**

Increased muscle tone

Resistance to passive muscle stretch at a joint

Velocity and joint-angle dependent

Has to be palpated – ie you need to lay on hands

Other features: clonus, increased reflexes, upgoing plantars

Sustained and repititive muscle contractions & abnormal postures

Visible Involuntary movements

Suggests active rather than passive movements

Not usually velocitydependent

Dependent on mental state: alertness, emotion

# Types of dystonia

Excessive movements:

chorea, athetosis, dystonic
tremor, myoclonus, "spasms"

Basal Ganglia and Cerebellum damage

Decreased movements: extrapyramidal rigidity, hypokinesia

Oropharyngeal: mouth, tongue, pharynx, larynx

Axial muscles: cervical, erector spinae (opisthotonus)

Upper Limbs: Elbow extensor rigidity, fisting

Lower limbs: Ankle evertors as well as invertors

# Interventions/Treatments

# **Effective Treatments for reducing Spasticity**

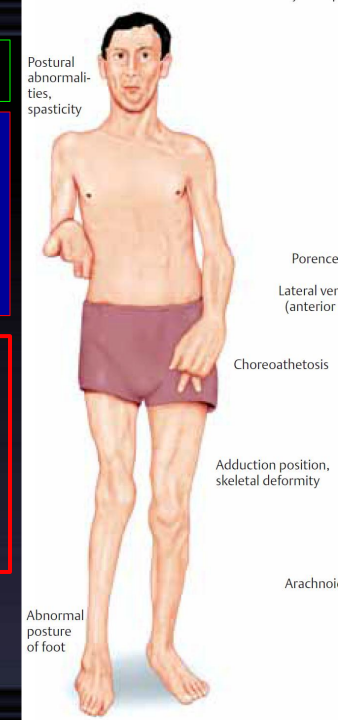
- PT Intervention
- Electrical and Functional Electrical Stimulation
- Casting
- Mental Imagery
- Pharmacological Interventions
- Surgery



### CEREBRAL PALSY (CP)

CP includes disorders apparent at birth or in childhood due to intrauterine or neonatal brain damage mainly motor control centers; deficits are non-progressive

- > hypoxia in utero and/or during parturition
- neonatal cerebral haemorrhage and/or infarction
- > trauma, neonatal or during parturition
- > prolonged seizures status epilepticus
- > Hypoglycaemia
- > kernicterus



#### **MULTIPLE SCLEROSIS**

- Auto-immune demyelinating disease, in which the body's own immune system attacks the myelin sheath of nerves mainly of brain, spinal cord & optic nerve
- Demyelination affects saltatory conduction and cause muscle weakness& wasting.
- Onset usually in young adults (F>M)
- When it causes demyelination of descendindg motor tracts in the brainstem & spinal cord, the patient develops spasticity and other signs of UMN lesions.
- The disease frequently remits and relapses
- Treated by steroids

#### **STROKE**

- Haemorrhagic stroke & Ischaemic stroke
- •Results in paralysis in the opposite half of the body
- A lesion in Corona Radiata on one side can cause Monoplegia in a contralateral limb (UL or LL).
- A lesion in the Internal Capsule on one side may cause Hemiplegia or Hemiparesis on the contralateral side
- Presents with features of upper motor neuron syndrome UMNL.

# **Brown Sequard syndrome**

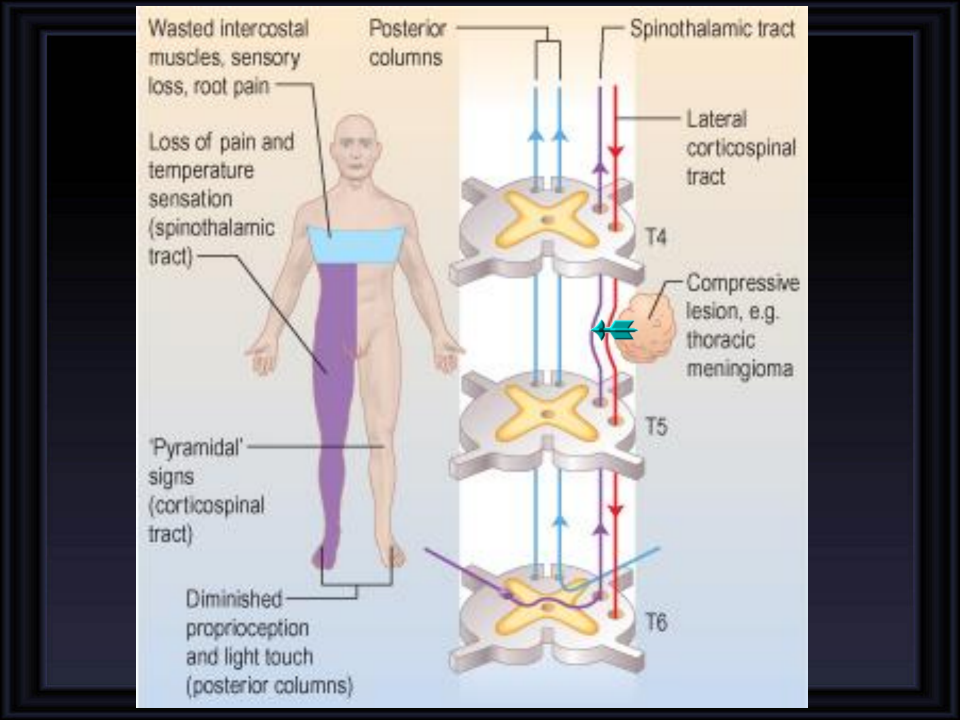
#### **HEMISECTION OF SPINAL CORD**

# **Ipsilateral Loss:**

- Fine touch, Vibration, Proprioception (Dorsal Column)
- Leg Ataxia (Dorsal Spinocerebellar)
- Spastic Paresis below lesion (Lat Corticospinal)
- Flaccid Paralysis (Vent horn destruction)
- Dermatomal Anesthesia (Dorsal Horn destruction)

### **Contralateral Loss:**

- Loss of pain and temp (lat Spinothalamic)
- Loss of crude touch and Pressure (Vent Spinothalamic)
- Minor Contralat Muscle Weakness (Vent Corticospinal)
- Leg Ataxia (Vent Spinocerebellar)



#### Complete transection of spinal cord

- •in lower cervical region →quadriplegia
- •down in the thoracic region → paraplegia (both lower limbs)

#### 3 Stages:-

A/ Spinal shock (2-6 weeks)
B/ Recovery of reflex activity
C/ Paraplegia in extension

Voluntary movements and sensations are permanently lost

A/Spinal shock: Immediately following transection there is loss of all sensations (anaesthesia), reflexes, tone (flacciduty) and voluntary movement below the level of the lesion, due to interruption of all sensory and motor tracts.

Bladder urinary retention with overflow) and Loss of vasomotor tone occurs causes a fall in blood pressure;

# B/Stage of return of reflex activity due to increase in degree of excitability of the spinal cord neurons

- ➤ Gradual rise of arterial blood pressure
- Exaggerated tendon reflexes, spasticity, viceral reflexes return (micturition, defecation & erection reflexes)
- Mass reflex A minor painful stimulus to the skin of the lower limbs causes withdrawal and evoke autonomic reflexes (bladder and rectum emptying, sweating, blood pressure rise)

#### C/ Stage of extensor paraplegia

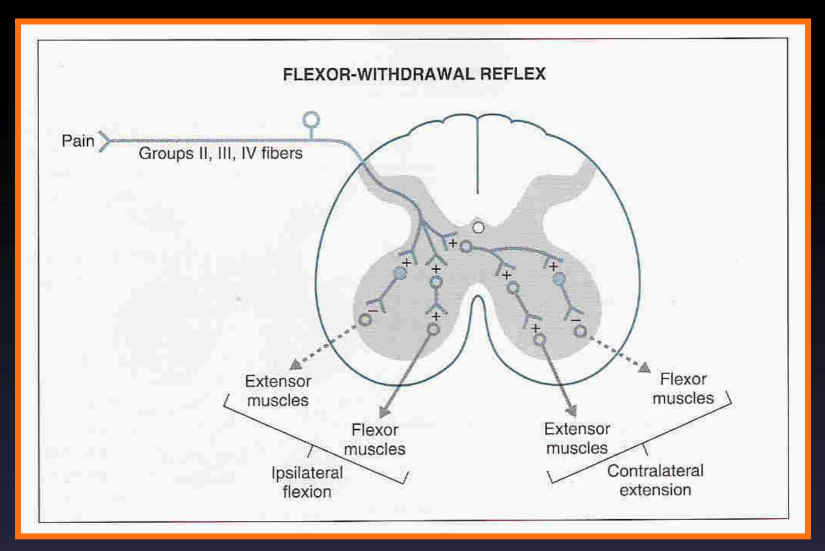
The lower limbs become spastically extended.

- -Extensor reflexes become exaggerated, as shown by tendon jerks and by the appearance of clonus.
- -The positive supportive reaction returns and the patient can stand on his feet with appropriate support.

The flexor withdrawal reflex and crossed extensor reflex returns



# STRETCH REFLEX Homonymous muscle Group la afferent a Motoneuron Synergistic muscles Antagonistic muscles



#### POLYSYNAPTIC REFLEX

