



Pain Modulation

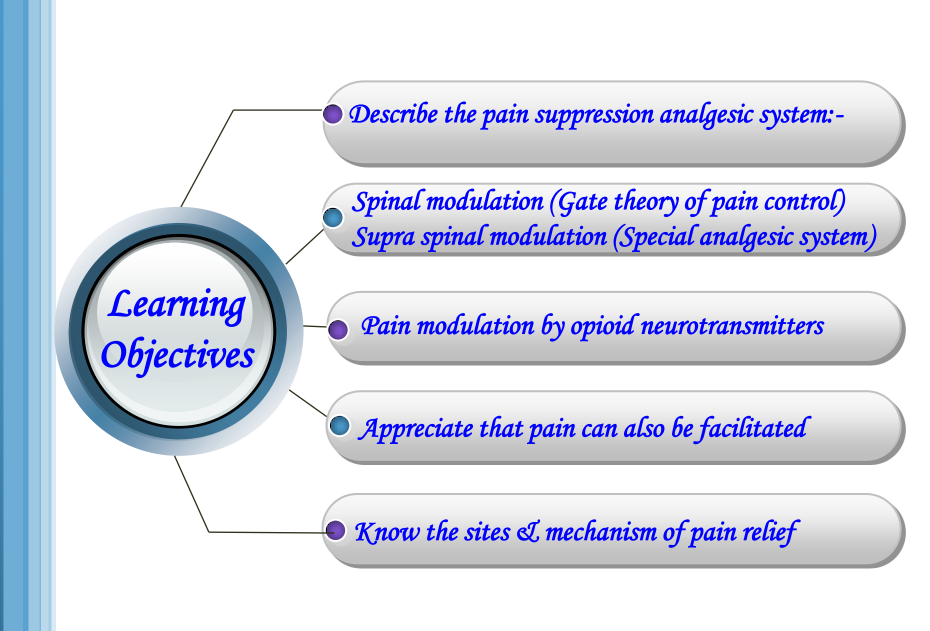
Dr. Hayam Gad

MBBS, MSc, PhD

Associate Professor Of

Physiology

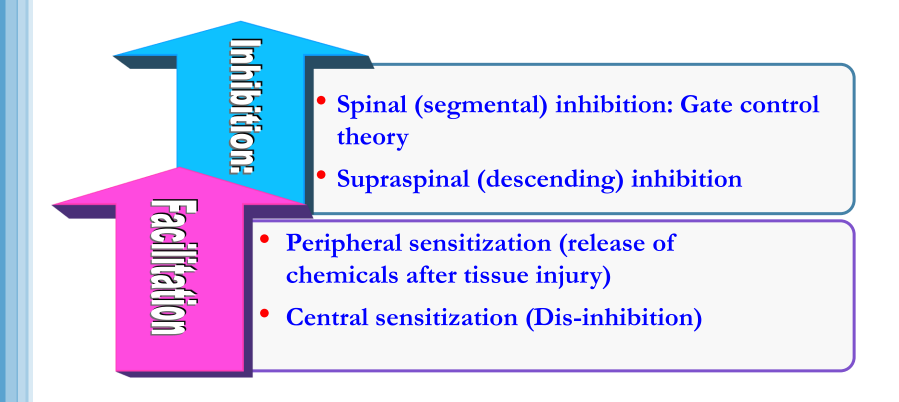
College of Medicine, KSU



What is Pain Modulation

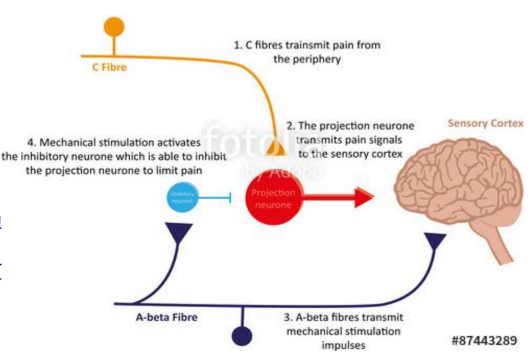
It means pain perception variability (the degree to which a person reacts to pain)

i.e. A decrease or an increase in the sensation of pain caused by inhibition or facilitation of pain signals



The gate theory of pain control

Special neurons in the the dorsal horn of spinal cord (SGR) form the gate through which pain impulses must pass to reach brain.



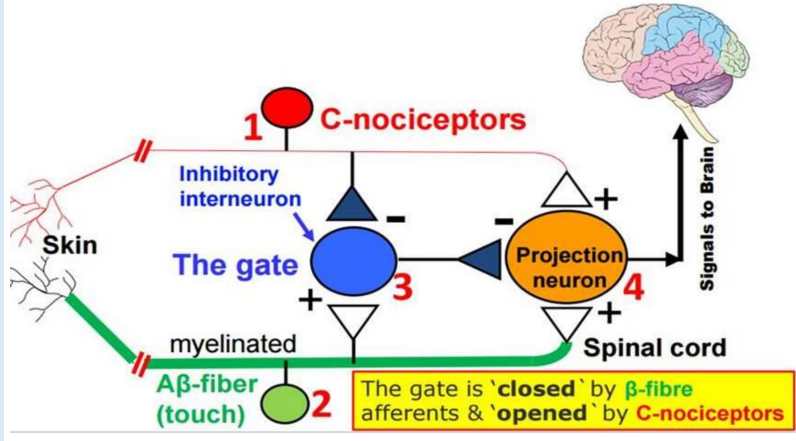
Four variables control this gate:

Type C- fibres (slow pain)

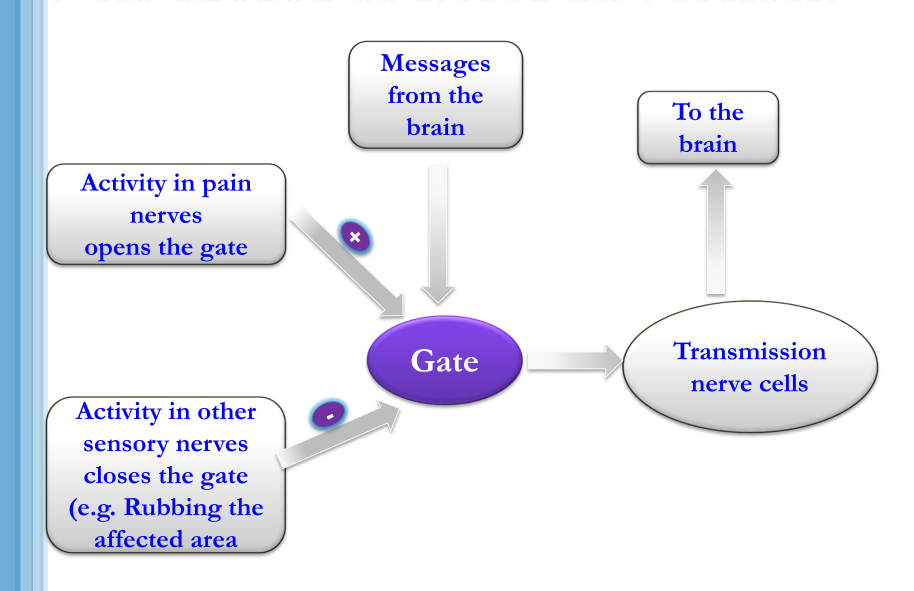
Type A-β fibres (light touch)

Inhibitory interneurons

Projection neuron

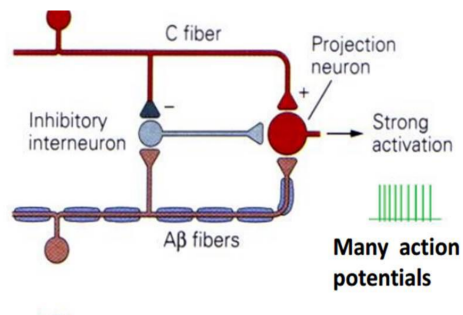


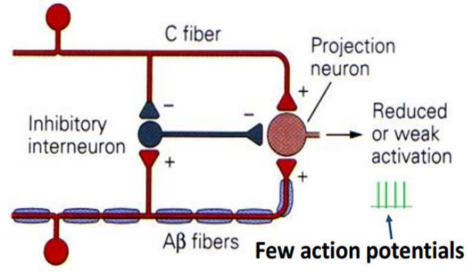
Gate opened or closed by 3 factors:

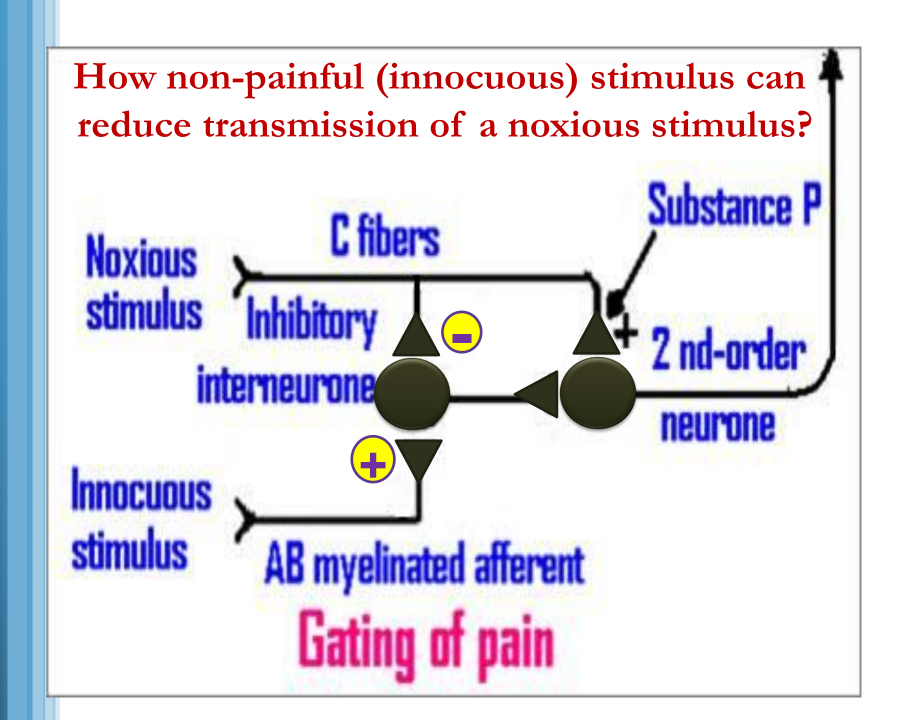


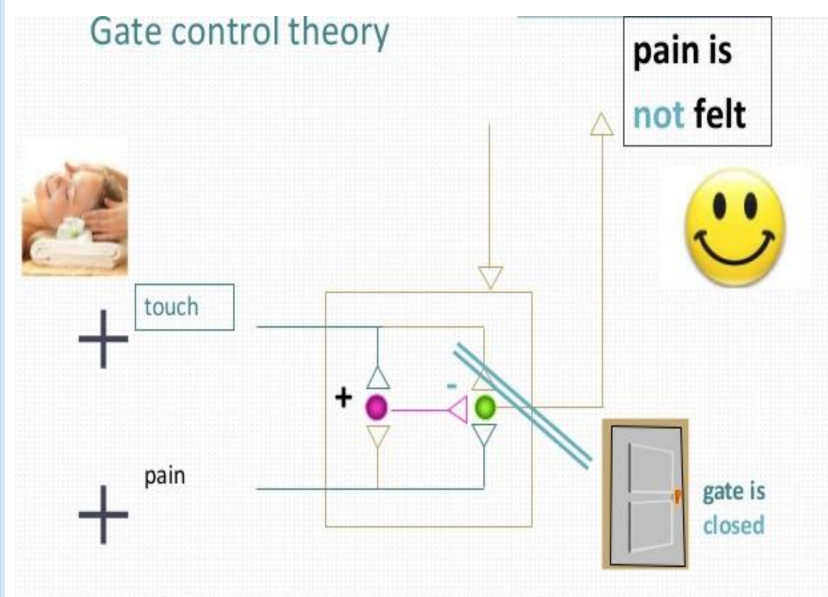
The gate theory of pain control (Cont.)

- Projection neuron receives input from both C-fibers and Aβ fibers.
- Impulses coming along type C pain fibers cause the release of substance P from these fibers and inhibits the inhibitory interneuron (open the gate).
- While impulses coming along Aβ fibers tend to keep the gate closed by activating the inhibitory interneuron.

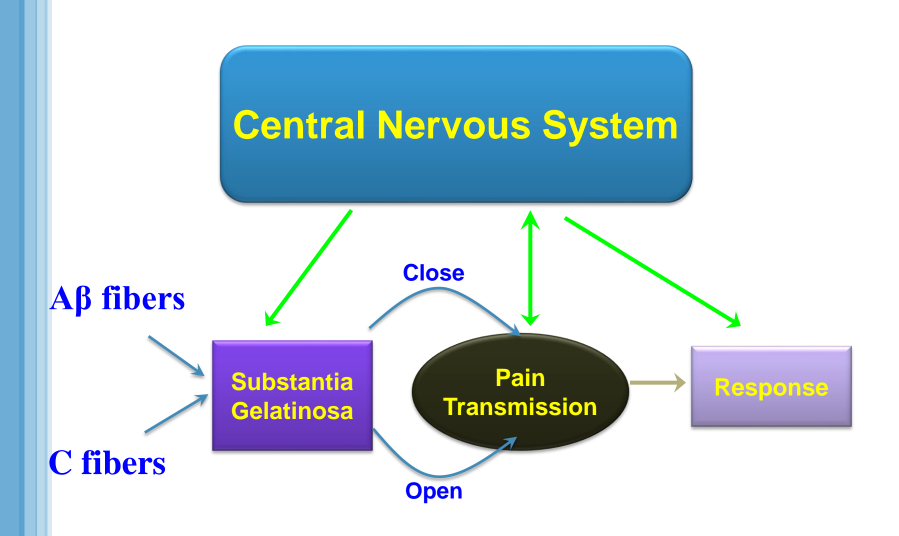








When pain and touch fibres are stimulated together, gate will be closed & pain is not felt



- The gate theory explains the pain relief by:
 - Skin rubbing
 - Shaking the painful part
 - Trans Cutaneous Electrical Nerve Stimulation (TENS)
 - Acupuncture
- All are supposed to stimulate mechanoreceptors that activate neurons of dorsal column, the collaterals relieve pain.





What is the Central Control Trigger

- •Specialised nerve impulses arise in the brain itself and travel down the spinal cord to influence the gate.
- "It can send both inhibitory and excitatory messages to the gate sensitising it to either C or A-β fibres.
- The inhibitory neurons make a pain blocking agent called encephalin.
- ■Encephalin is an opiate substance which can block substance P, the neurotransmitter from the C fibers, and this keeps the gate closed.

Supra spinal modulation

(Special pain control analgesic system)

This is a specific system that blocks pain transmission in CNS. Its major constituents are:

The periventricular & periaqueductal gray areas

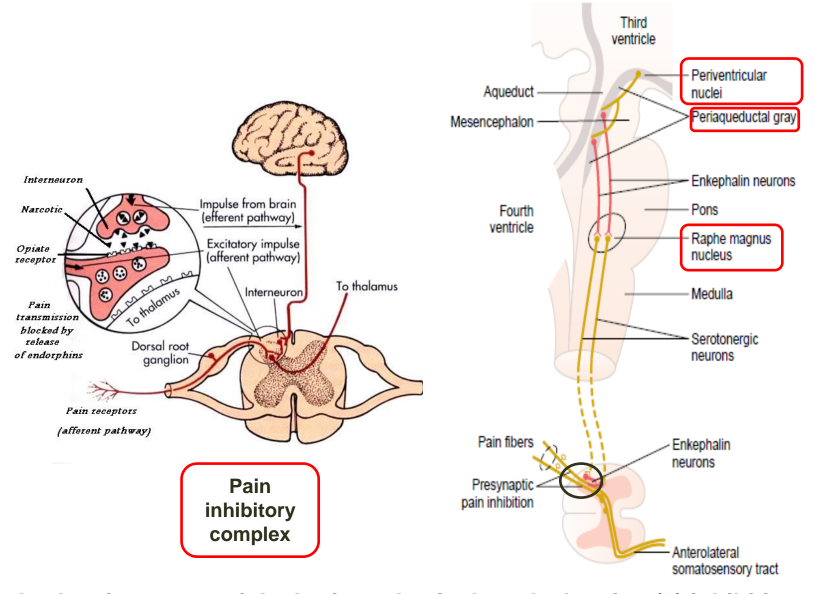
In the mesencephalon and upper pons surround portions of the third and fourth ventricles and the aqueduct of Sylvius

Raphe magnum nucleus (RMN)

A thin midline nucleus located in the lower pons and upper medulla.

Pain inhibitory complex

Multiple short neurons, terminate on central endings of pain conducting afferent fibers In dorsal horn of SC, release encephalin cause pre & postsynaptic inhibition of pain transmission



Analgesia system of the brain and spinal cord, showing (1) inhibition of incoming pain signals at the cord level and (2) presence of *enkephalin-secreting neurons* that suppress pain signals in both the cord and the brain stem.

Analgesia occurs as follows:



Periventricular
nucleus
projects to
PAG.



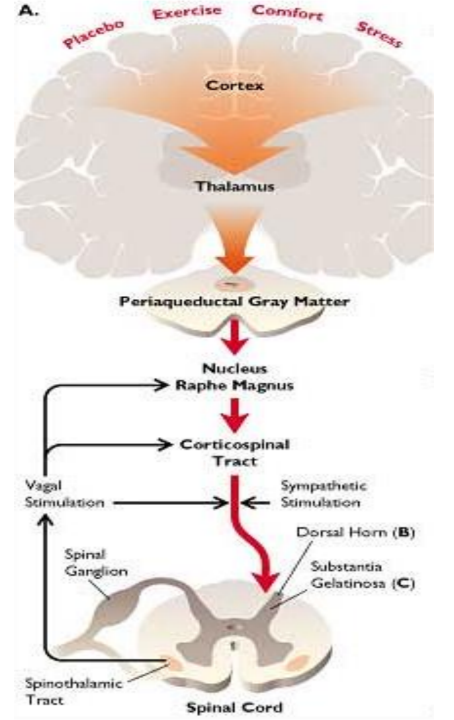
PAG
projects
neurons
containing
aspartate &
glutamate,
stimulate
(RMN)



RMN
projects
serotoninergic
neurons to
dorsal
horn.



They
block
pain
signals by
activating
PIC.

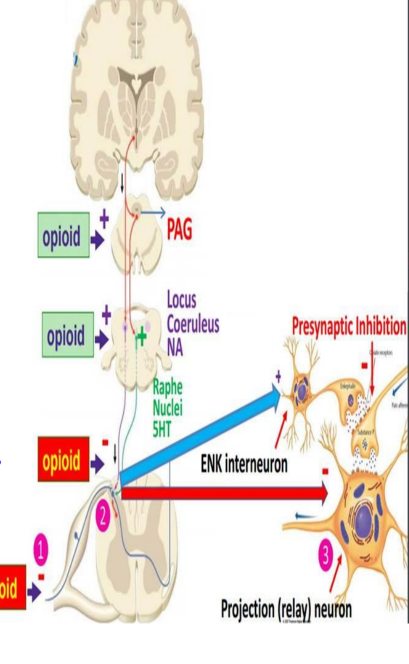


Opioid Peptides and Pain Modulation

- They are natural analgesic substances (morphine-like substances) present in body.
- They act by binding to opiate receptors in analgesic system and dorsal horn of SC on central ending of pain conducting pain fibers.

Site of release of opioid peptides:

- Endorphin: In PAG. They inhibit GABAnergic interneurons that normally suppress the antinociceptor neurons
- Encephalin: In interneurons of lamina II responsible for inhibiting the nocioceptor- spinothalamic neurons
- Dynorphin: In hypothalamus, PAG, reticular formation, and dorsal horn.
- Endogenous morphine: In terminals forming synapses with neuron involved in pain modulating pathways.



Mechanism of actions of Opioid peptides on pain transmission

They exerts their analgesic effects by acting at various sites in peripheral & CNS

Direct effect

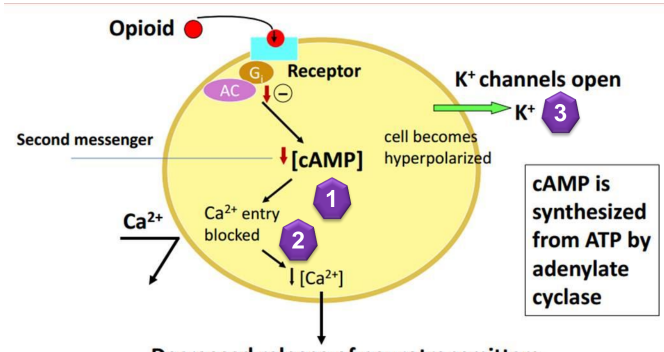
- Inhibiting discharge of nociceptor neurons.
- Inhibiting release of substance P from central terminal of nociceptor neurons
- Cause inhibition of dorsal horn spinothalamic neuron.

Indirect effect

- Activating the descending inhibitory pathway by exciting PAG neurons
- Activating neurons in the brain stem which suppress pain transmission directly or indirectly via activation of encephalinergic containing inhibitory interneurons

Mechanism

Cellular actions of Opioid peptides

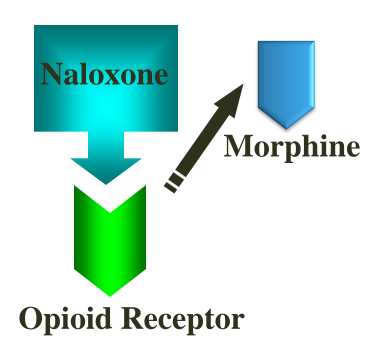


Decreased release of neurotransmitters

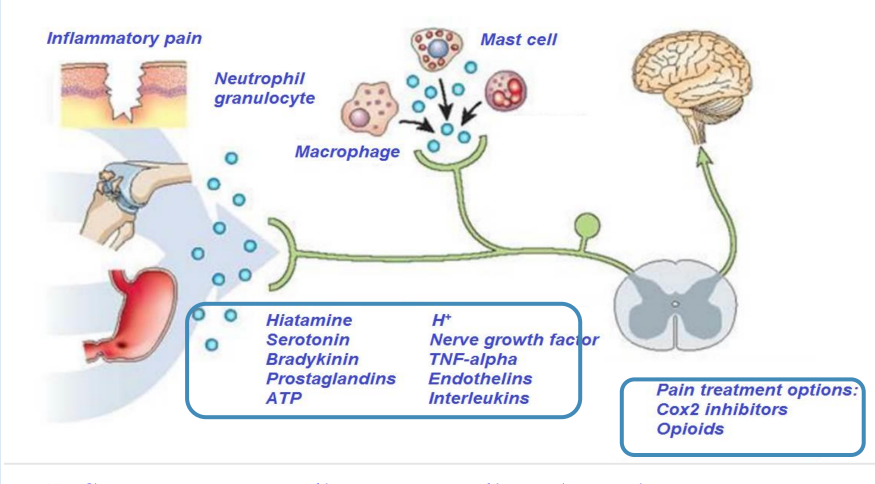
- Reduction of cAMP synthesis by inhibiting Adenyl cyclase
- Inhibition of transmitter release by inhibiting opening of Ca⁺⁺ channels
- Hyperpolarization by facilitating opening of voltage gated K⁺ channels

Opioid Antagonist: Naloxone

- Used to reverse opioid overdose
- Displaces receptorbound opioids
- Good for overcoming respiratory and CV depression

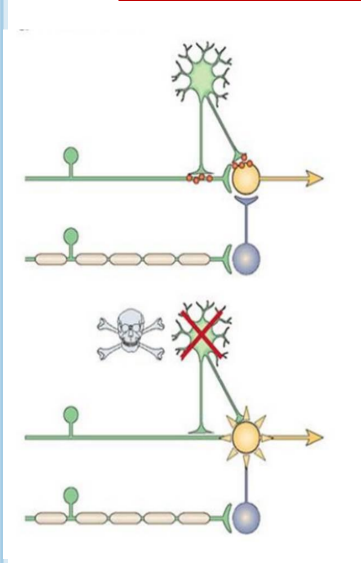


Pain Facilitation: Peripheral Sensitization



 Inflammatory mediators can directly activate nociceptors or cause their sensitization (decrease threshold)

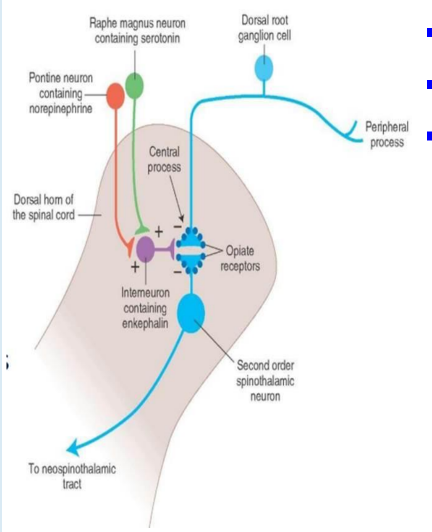
Pain Facilitation: Dis-inhibition



 Pain transmission is controlled by inhibitory interneurons

Loss of these inhibitory
 interneurons after excessive
 release of glutamate results in
 increased excitability of
 projection neurons and thus
 enhanced pain sensation

Neurotransmitters for Pain Modulation



- Serotonin
- Noradrenaline
- · Encephalin
 - The serotonergic and noradrenergic neurons are crucial in the supraspinal modulation
 - Destroying these neurons with neurotoxins blocks the their analgesic actions

Terms frequently used

Hyperalgesia

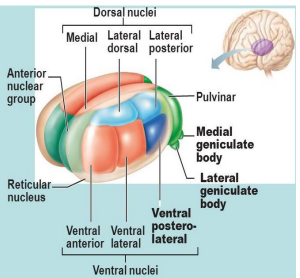
Excessive Pain (e.g due to sun burn)

Muscular Pain
Less blood flow in the muscles (ischemia)

Allodynia
Pain caused by any
other
sensation e.g. touch

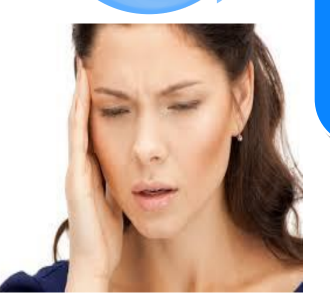
Causalgia
Burning pain





- It is a neurological condition that results from a brain stroke affecting the thalamus.
- Cause: Obstruction of the thalmogeniculate branch of the posterior cerebral artery.
- Affects posterior thalamic nuclei.

Trigeminal neuralgia



- It is excruciating intermittent pain by stimulation of trigger area in the face.
- e.g. Washing of face, combing hair, blast of air on face.
- It results from compression of trigeminal nerve root by blood vessels.

Stress induced analgesia



- Pain suppression response that occurs during or following exposure to a stressful or fearful stimulus.
- It's a well known phenomenon seen when the soldier is wounded in battle field but feels no pain until the battle is over.

the cause is not known may be it is similar to gate control hypothesis.

Phantom pain sensations Perceptions that an individual experiences relating to a limb or an organ that is not physically part of the body.

Post-Amputation Pain

Phantom Limb Pain

Neuropathic pain (NP)

- Pain caused by a primary lesion or dysfunction in the nervous system.
- Classification:
 - Central NP-Damage of CNS
 - Peripheral NP- Damage to PNS
- Resistant to the current analgesic therapy.
- Can persist for years.
- Clinical symptoms: Hyperalgesia, allodyni and spontaneous pain
- Examples: post herpetic neuralgia, diabetic neuropathy and after chemotherapy.



Mechanism of pain relief

Block production of inflammatory mediators .e.g. Aspirin & nonsteroidal anti-inflammatories.

Exogenously administration of opioid like drugs.

Sympathectomy can be useful.

Electrical stimulation of the dorsal column.

Selective activation of large diameter afferent fibers by transcutaneous electrical nerve stimulation.

Stimulation of brainstem sites or administration of drugs which can modify serotoninergic or adrenergic neurons e.g. antidepressants.

