

PHGY 212 - Physiology

SENSORY PHYSIOLOGY

Somatic Senses

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Somatosensory Modalities

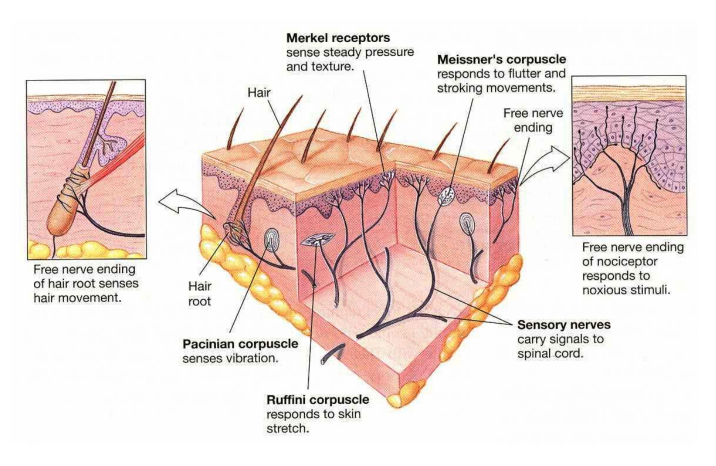
There are four somatosensory modalities:

Touch-Pressure
Proprioception
Temperature
Nociception

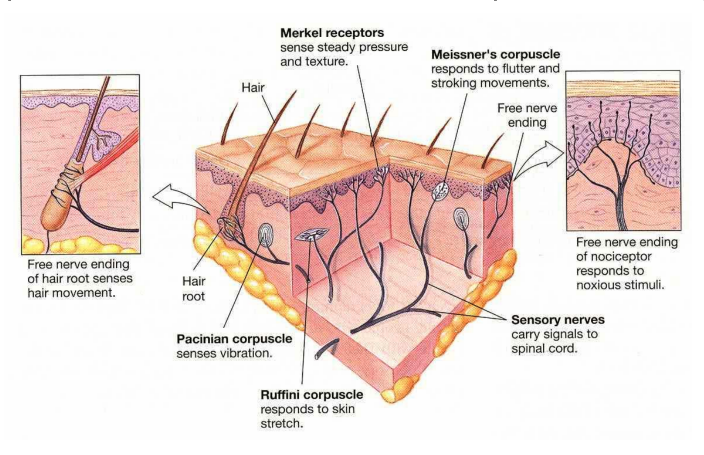
Somatosensory Receptors

Free nerve endings (touch-pressure, temperature, pain)

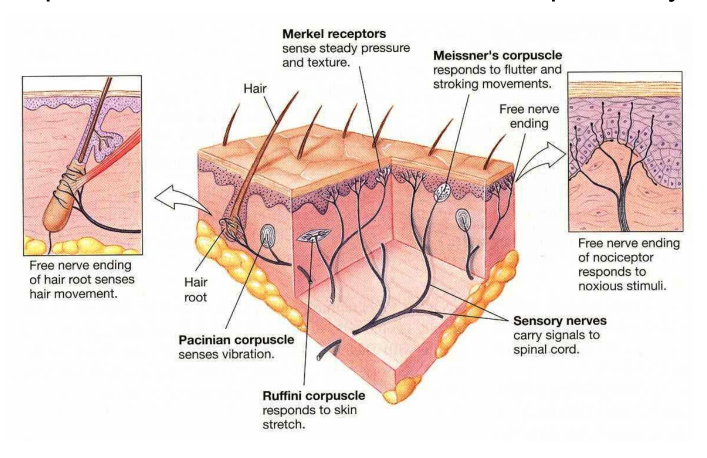
Around hair roots and under skin surface



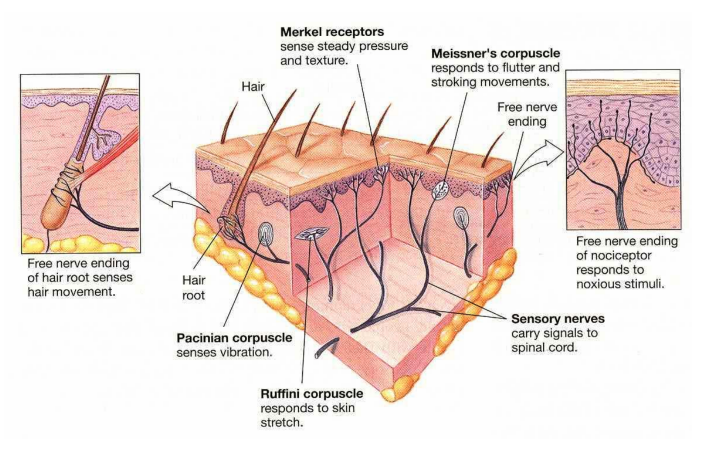
Meissner's corpuscle (flutter) Encapsulated in connective tissue in superficial skin layers



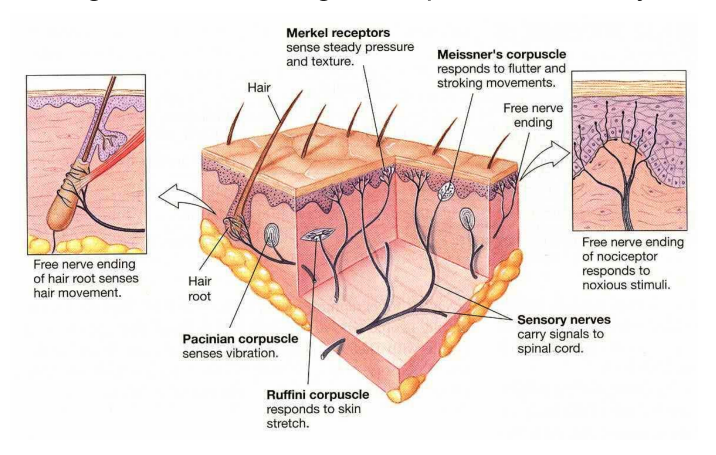
Pacinian corpuscle (vibration) Encapsulated in connective tissue in deep skin layers



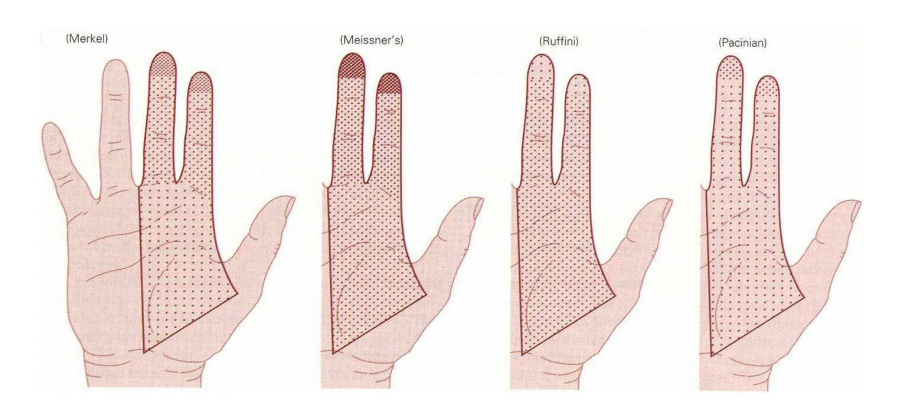
Ruffini corpuscle (skin stretch) Enlarged nerve endings in deep skin layers



Merkel receptors (steady pressure on skin) Enlarged nerve endings in superficial skin layers



Touch-pressure receptors are distributed neither identically nor uniformly.



Primary Sensory Fibers

Aβ **fibers**: large, myelinated, fast fibers (30 – 70 m/s) fine touch; pressure; proprioception

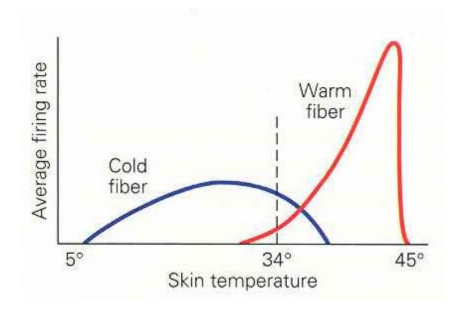
Aδ **fibers**: small, myelinated, slow fibers (12 – 30 m/s) crude touch; cold; fast & sharp pain

C fibers: small, unmyelinated, very slow fibers (0.5 – 2 m/s) temperature; slow & dull pain

Sensing Temperature

Thermoreceptors are free nerve endings with small (1-mm) receptive fields scattered across the body.

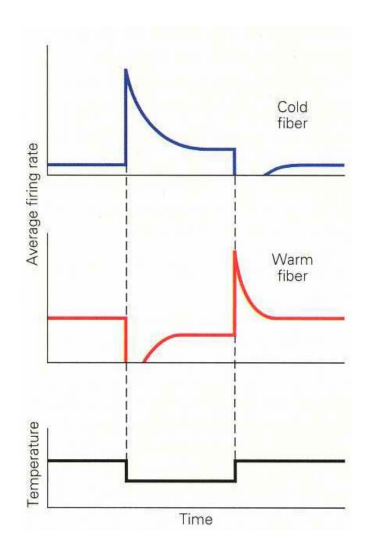
Skin thermoreceptors play a role in temperature regulation, which is controlled by centers in hypothalamus.



Sensing Temperature

Thermoreceptors are sensitive to changes in temperature, not to absolute temperature.

Thermoreceptors adapt only between 20° and 40° C. Stimuli outside this range activate *nocireceptors* because of the high probability of tissue damage.



Sensing Pain

Nociceptors are free nerve endings sensitive to a variety of molecules released with tissue injury.

Chemical mediators include:

- 1) K⁺, histamine, bradykinin & prostaglandins from site of injury;
- 2) ATP & 5-HT (serotonin) from platelets activated by injury;
- 3) Substance P from the primary sensory neurons.

Sensing Pain

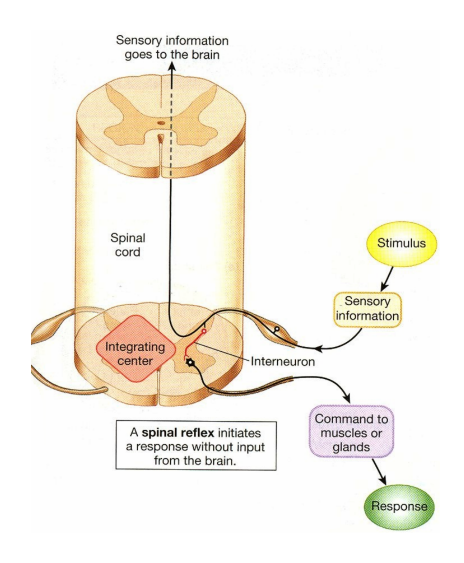
Fast & sharp pain: transmitted via A δ fibers from the activation of *thermal nociceptors* (>45° or <5° C) or *mechanical nociceptors* (intense pressure).

Slow & dull pain: transmitted via C fibers from the activation of *polymodal nociceptors* (high-intensity mechanical, thermal or chemical stimuli).

Spinal Reflexes

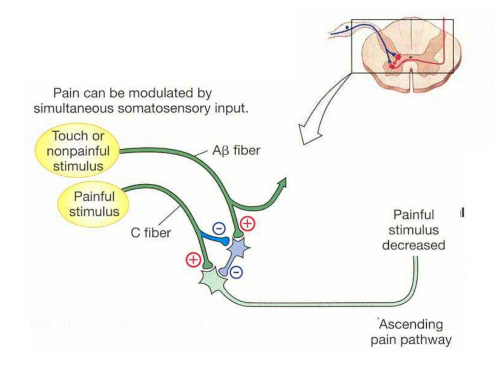
Painful stimuli are carried by ascending pathways to the cortex, where they become conscious sensation.

Not all nociceptive responses rely on cortical circuits. Subconscious withdrawal reflexes can occur within the spinal cord. These are called **spinal reflexes**.



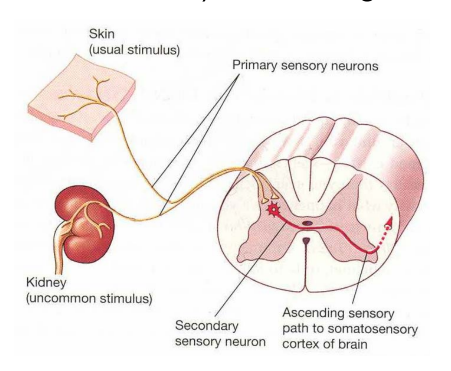
Gating Theory of Pain Modulation

Our perception of pain is subject to modulation. In the gating theory of pain modulation, inhibition of the ascending pain pathway can be enhanced by the activation of non-nociceptive somatic A β fibers.



Referred Pain

Visceral pain is often poorly localized and often felt in somatic areas distant from the painful stimulus. This **referred pain** is due to the *convergence* of nociceptive fibers (of same **dermatome**) onto a single ascending tract.



Analgesia

Analgesic drugs range from aspirin to opiates.

Aspirin inhibits the synthesis of *prostaglandins* and thus slows the transmission of pain signals from the site of injury.

Opiates (endogenous opioids: endorphins & enkephalins) act directly on opioid receptors in the brain, which activate descending pathways that inhibit incoming pain signals.

Reading

Silverthorn (2nd edition)

pages 291 – 294

Silverthorn (1st edition)

pages 273 – 276