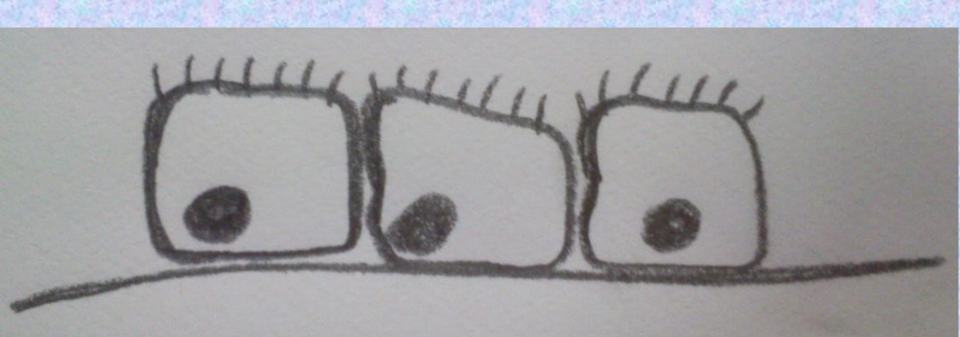
Receptors



Localisation: in membrame, in cytosol, in nuclei, in mitochondria

Receptor transform information from our sense to biological signals = electrical Action Potential

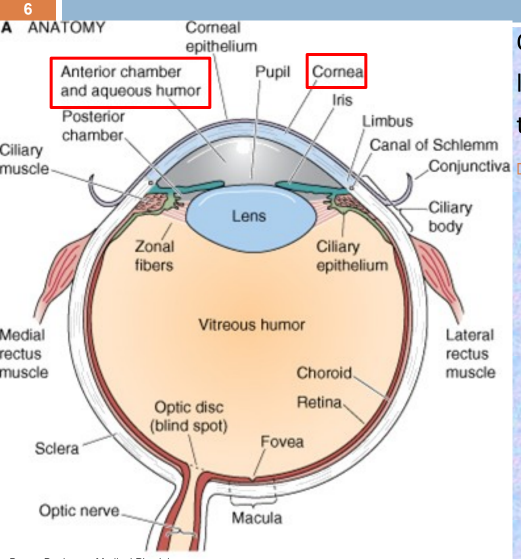
Type of receptors for senses

- Photoreceptors (rod and cone)
- Mechanoreceptors (touch on the skin, sound wave detection, wave detection in inner ear)
- Chemoreceptors (detection of molecules in food)
- Termoreceptors
 - Cold 23-28
 - Warm 38-43
 - Fast chang...0.1 gradius
 - Slow change....bigger difference

Physiology of vision

Functional anatomy of the eye

- Optical
- Neural
- Photoreceptors
 - Rods
 - Cones
- Phototransduction
 - Mechanism
 - Termination
 - Light adaptation
- Colour Vision

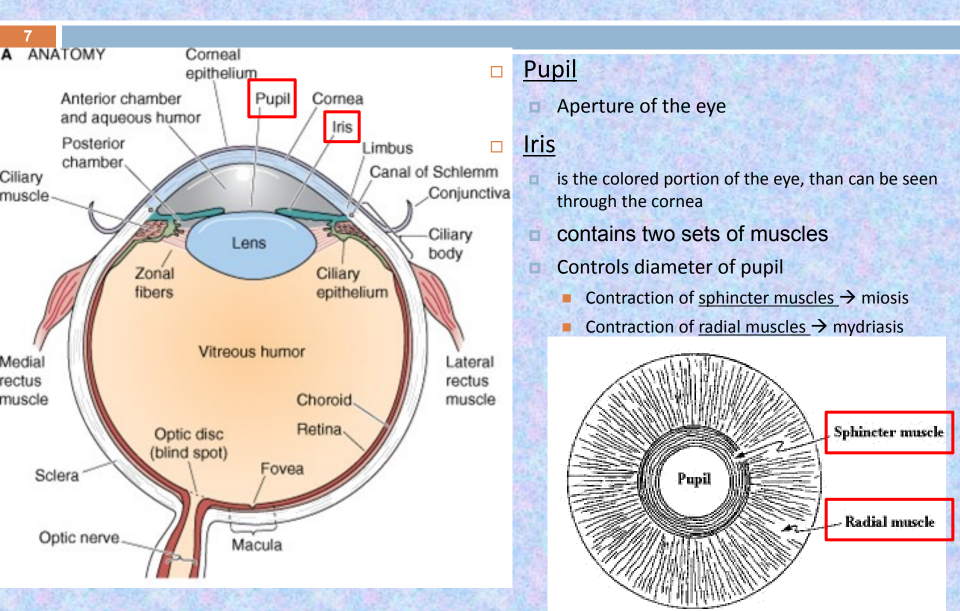


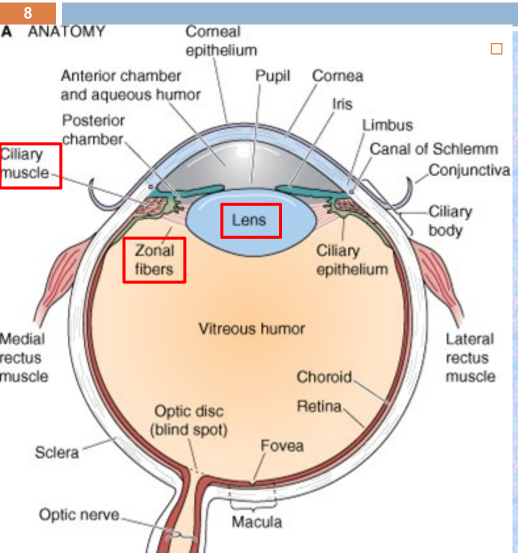
Boron, Boulpaep: Medical Physiology, 2003

Optical portion of eye focuses light thru cornea and lens onto the fovea.

Cornea

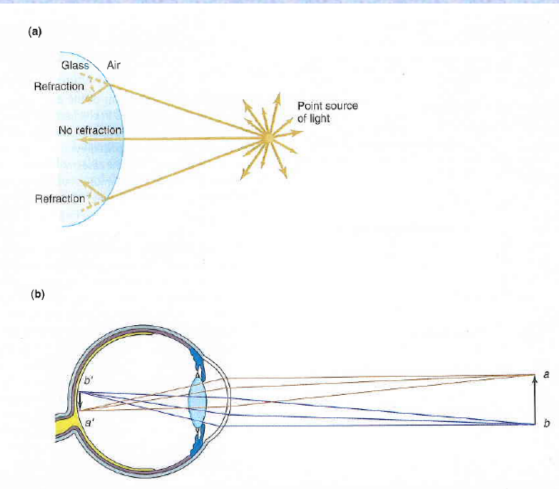
- Thin, transparent epithelium devoid of blood vessels
- Receives nutrients by diffusion from tear fluid
- Major refraktory portion of the eya, has unmyelinated nerve endings sensitive to touch and pressure
- Aqueous humor
- Produced by ciliary epithelial cells. Protein free watery liquid that supplies nutrients to cornea and lens
- Maintains intraocular pressure and gives shape to anterior portion of eye

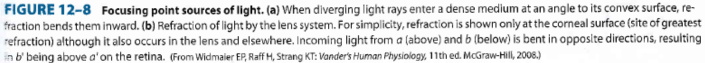




Lens

- Dense, high protein structure that adjusts optical focus
 - Focus adjusted by process called accommodation
 - At rest, <u>zonal fibers</u> suspend lens and keep it flat
 - Focus on objects far away
 - Contraction of <u>ciliary muscles</u> releases tension in zonal fibers
 - Lens becomes rounder
 - Focus on near objects





188 SECTION III Central & Peripheral Neurophysiology

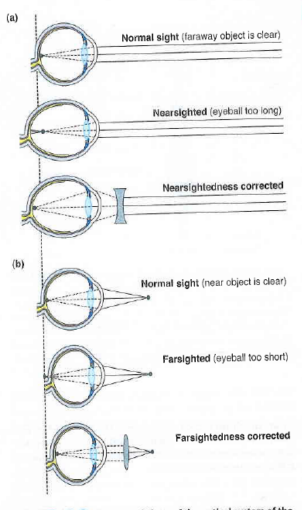
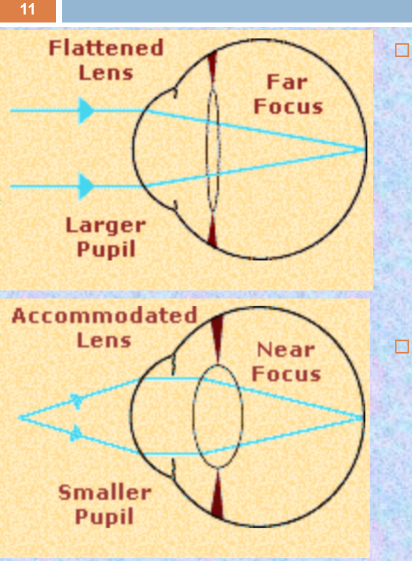


FIGURE 12–9 Common defects of the optical system of the eye. In hyperopia (farsightedness), the eyeball is too short and light rays come to a focus behind the retina. A biconvex lens corrects this by adding to the refractive power of the lens of the eye. In myopia (near-sightedness), the eyeball is too long and light rays focus in front of the retina. Placing a biconcave lens in front of the eye causes the light rays to diverge slightly before striking the eye, so that they are brought to a focus on the retina. (From Widmaier EP, Raff H, Strang KT: Vander's Human

10

Accommodation and associated disorders



Accommodation of the lens is limited and age dependent

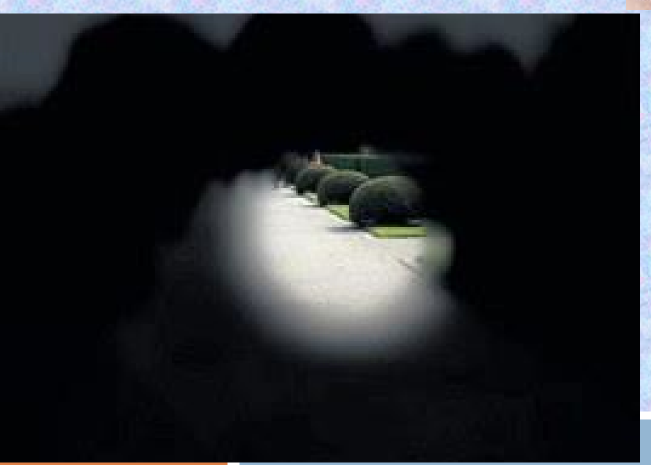
With age, lens becomes stiffer and less compliant.

 Age related loss of accommodation called presbyopia

Accommodation accompanied by adaptive changes in size of pupil



glaucoma

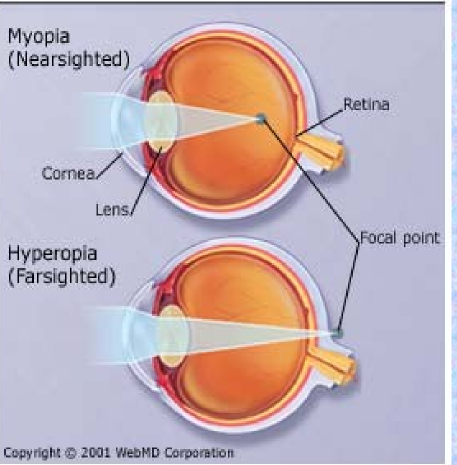




Accommodation and associated disorders

14

Myopia and Hyperopia



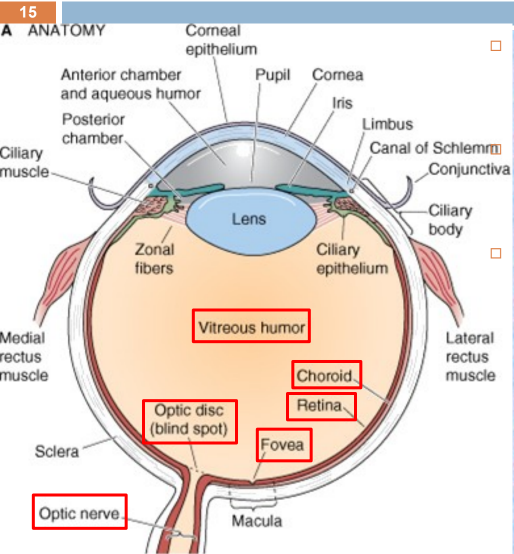
Myopia

- Image focused in front of retina
- Far away objects appear blurry

<u>Hyperopia</u>

- Image focused behind retina
- Close objects appear blurry

Each can be caused by abnormal shape of the eye as well.



Vitreous humor

 Gel of extracellular fluid containing collagen

<u>Choroid</u>

rich in blood vessels and supports the retina

Retina

D

Neural portion that transduces light into electrical signals that pass down the <u>optic nerve</u>

- Optic nerve exits at optic disc. Devoid of photoreceptors: <u>blind spot</u>
- <u>Fovea</u> is point on retina that has maximal visual acuity

RETINA

Its organized on layers Visual receptors+4types of neurons. Many different synaptic transmitters

Pigment epithelium

- Absorps light rays, prevention the reflection of rays back through the retina
 - Contains melanin to absorbs excess light
 - Stores Vitamin A
 - **Photoreceptors**
 - Transduce light energy into electrical energy
 - Rods and cones

Ganglion cells

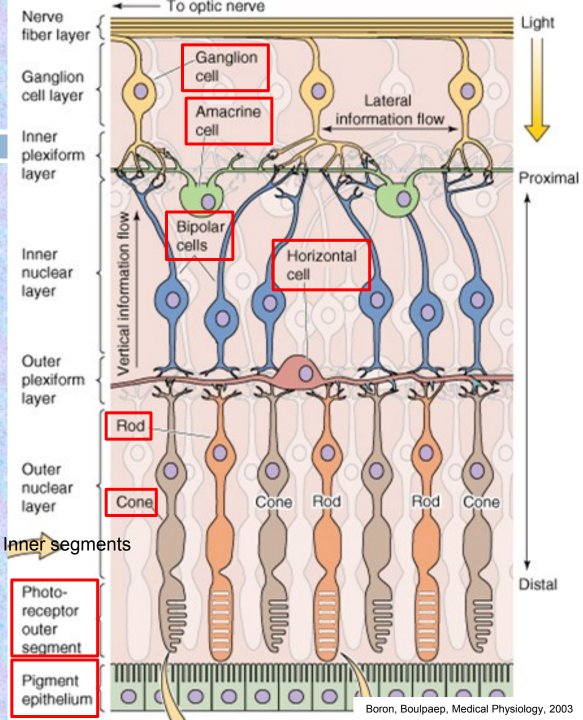
 Output cells of retina project via optic nerve

Bipolar cells – 12 different types occur

Horizontal cells

Amacrine cells - 29types have been described

 The neural elements of retina are bound together by glial cells – Muller cells



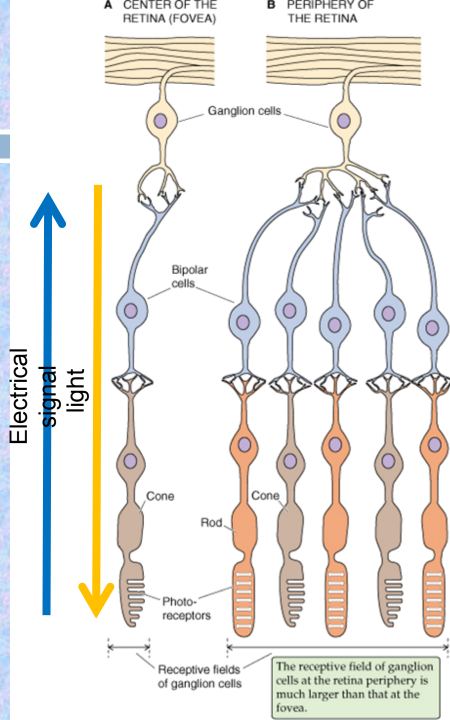
17

Periphery of retina

- □ High degree of convergence → large receptive field
- High sensitivity to light, low spatial resolution

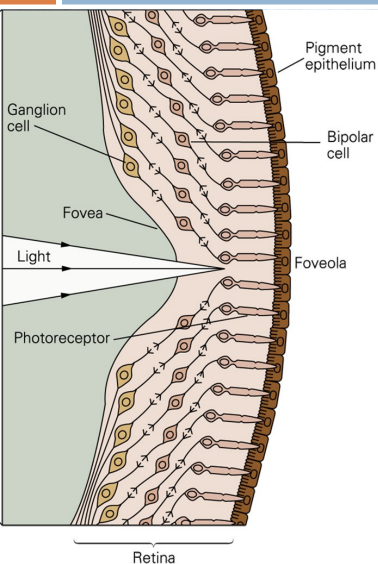
Fovea

- □ Low convergence \rightarrow small receptive fields
- Lower sensitivity to light, high resolution (visual acuity)



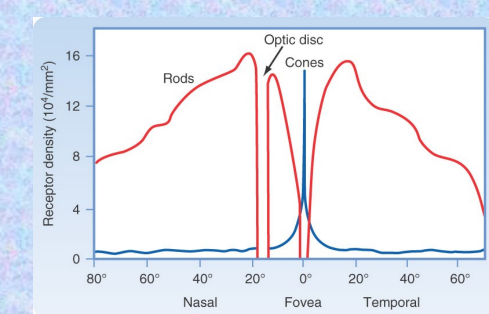
Fovea

18



Visual acuity of fovea enhanced by:

- One to one ratio of photoreceptor to ganglion cell
- Lateral displacement of neurons to minimize scattering of light
- High density of cones

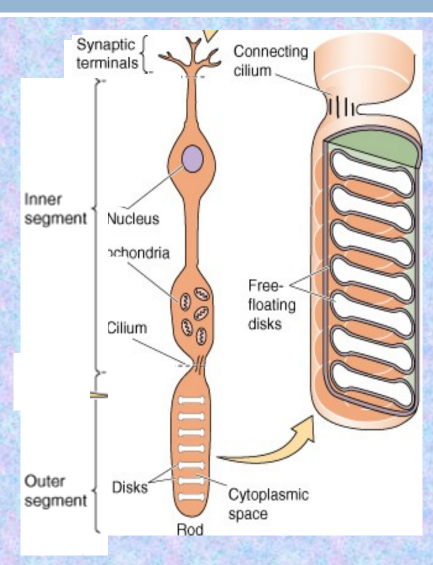


Photoreceptors

19

Rods

- Responsible for monochromatic, dark- adapted vision
- Inner segment contains nucleus and metabolic machinery
 - Produces photopigment
- Outer segments is transduction site
 - Consists of high density of stacks of <u>disk membranes</u>: flattened, membrane bound organelles
 - contain the photopigment rhodopsin

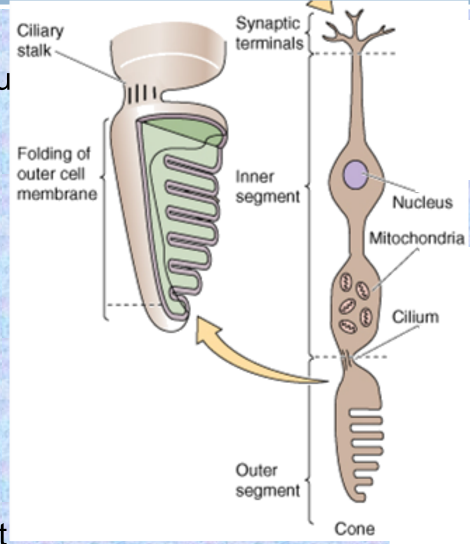


Photoreceptors

20

Cones

- 3 subtypes responsible for colou vision
- Inner segment produces photopigments similar to rhodopsin
- Outer segments is transduction site
 - consist of infolded <u>stack</u>
 <u>membranes</u> that are
 continuous with the outer
 membrane
 - vesicles containing pigment



Phototransduction: Dark current

Partially active guanylyl cyclase keeps cytoplasmic [cGMP] high in the dark

Outer segment contains <u>cGMP-gated</u> <u>cation channels</u>

Influx of Na⁺ and Ca²⁺

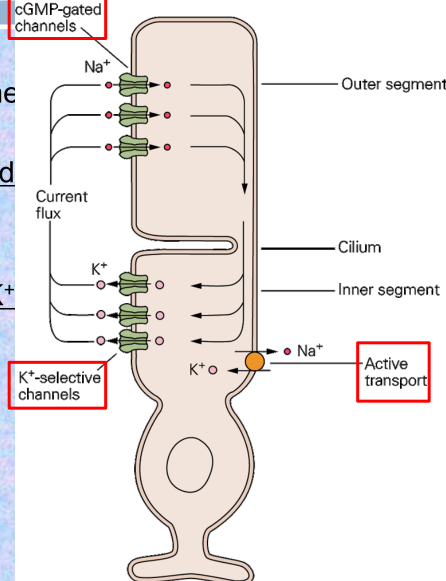
Inner segment contains <u>non-gated K⁺</u> <u>selective channels</u>

K⁺ efflux

21

Resting, or dark V_m is -40 mV

 concentration gradients maintained by Na⁺/K⁺ pump and NCX



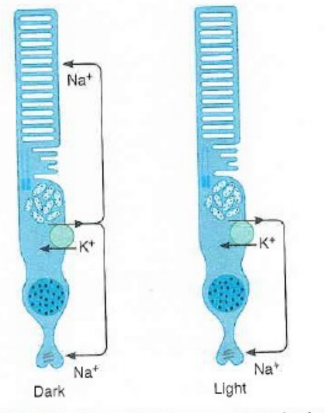


FIGURE 12–12 Effect of light on current flow in visual receptors. In the dark, Na⁺ channels in the outer segment are held open by cGMP. Light leads to increased conversion of cGMP to 5'-GMP, and some of the channels close. This produces hyperpolarization of

and some of the channels close. This produces hyperpolar the synaptic terminal of the photoreceptor.

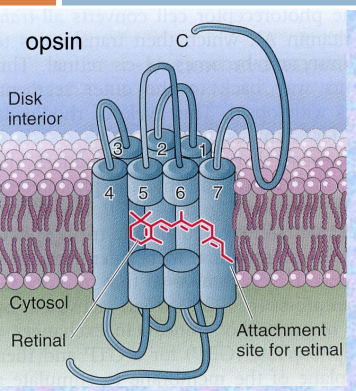
22

Phototransduction

23

- Photoreceptors <u>hyperpolarize</u> in response to light and release <u>less</u> neurotransmitter
- In darkness, the V_m of -40 mV keeps CaV channels in the synaptic terminal open
 - photoreceptors continuously release neurotransmitter
 glutamate
- absorption of light by photopigment &'s [cGMP]
 - cation channels close
 - K⁺ efflux predominates, hyperpolarizes cell (-70mV)
 - CaV channels close, decreased release of glutamate

Phototransduction: mechanism



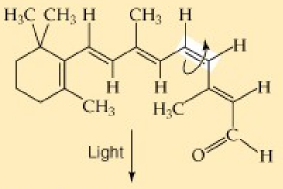
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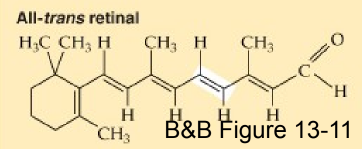
Photopigment rhodopsin is the light receptor in rods

opsin

- G-protein coupled membrane receptor
- Retinal= retinene1
 - Light absorbing compound

 the aldehyde form of retinol or Vitamin A





retinal changes conformation from <u>11-cis</u> to <u>all-trans</u> after absorbing a photon
 isomerization of retinal activates opsin

Outer segment membrane

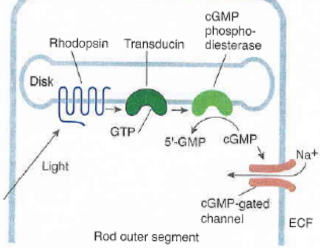


FIGURE 12-14 Initial steps in phototransduction in rods. Light activates rhodopsin, which activates transducin to bind GTP. This activates phosphodiesterase, which catalyzes the conversion of cGMP

to 5'-GMP. The resulting decrease in the cytoplasmic cGMP concentration causes cGMP-gated ion channels to close.

darkness. The amount of rhodopsin in the receptors therefore varies inversely with the incident light level.

CONE PIGMENTS

Primates have three different kinds of cones. These receptors subserve color vision and reepond movimally to light at some

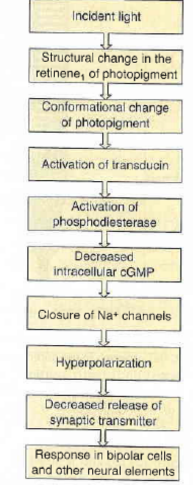
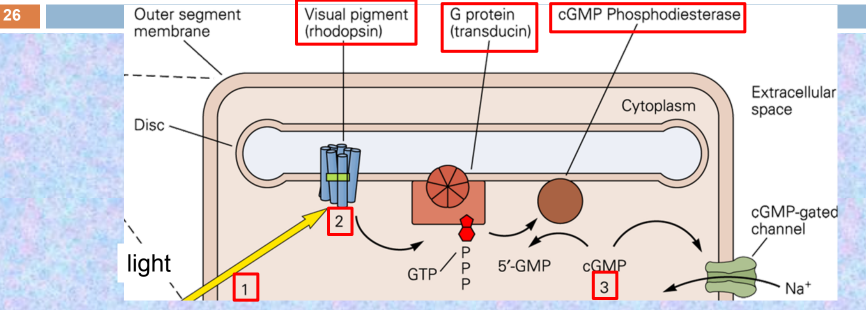


FIGURE 12-15 Sequence of events involved in phototransduction in rods and cones.

Phototransduction: mechanism



- Absorption of a photon isomerizes retinal
 - Converts opsin to metarhodopsin II
- 2. Metarodophsin II activates the G-protein transducin
 - Activates cGMP phosphodiesterase (PDE)
- 3. PDE hydrolyzes cGMP to GMP

a

a

- a) Decreased [cGMP] closes cGMP gated cation channels
- b) Photoreceptor hyperpolarizes, less glutamate released

Phototransduction: termination

- Activated rhodopsin is a target for phosphorylation by <u>rhodopsin kinase</u>
 - Phosphorylated rhodopsin inactivated by cytosolic protein <u>arrestin</u>
- All-trans retinal transported to the pigment epithelium where it is converted back to 11-cis retinal, and recycled back to the rod
- Activated transducin inactivates itself by hydrolyzing GTP to GDP

Phototransduction: light adaptation

Eyes adapt to increased light intensity and remain sensitive to further changes in light

- Optic adaptation:
 - Constriction of pupils to allow in less light
- Photoreceptor adaptation:
 - □ The closure of cGMP gated channel reduces inward flux of Ca²⁺ → decreased [Ca²⁺]_i
 - Ca²⁺ induced inhibition of guanylyl cyclase removed

■ More cGMP made → reopening of some cGMP gated channels → influx of cations → slight depolarization

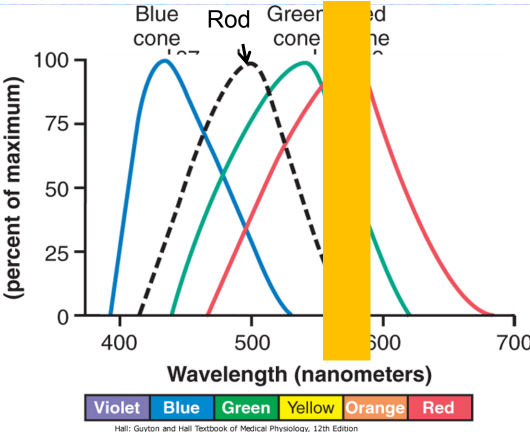
Photoreceptor can once again be stimulated (hyperpolarized) by photons

Colour Vision

29

- 3 types of cones, each conta photopigment with different absorption spectra

 - 420 nm blue
 530 nm green
 560 nm red
 Colour interpreted by ratio of cone stimulation
 - Orange (580nm) light stimulates:
 - Blue cone 0%
 - □ Green cone 42%
 - Red cone 99%
 - 0:42:99 ratio of cone stimulation interpreted by brain as orange



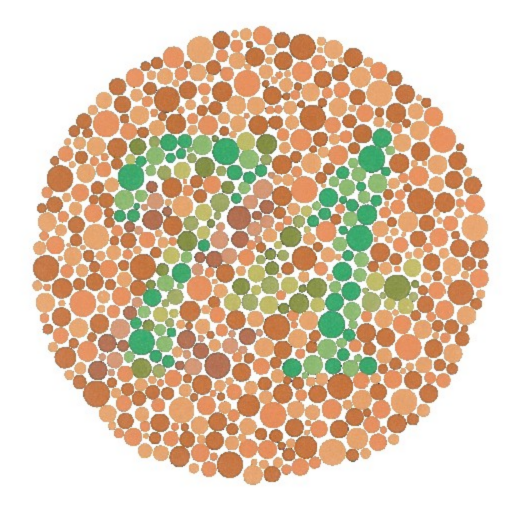
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Guyton Figure 50-8

Colour Vision: Disorders

30

- Malfunction of one group cones leads to colour blindness
- Most common form is red green colour blindness
 - Either red or green con are missing
 - Difficulty distinguishing from green because the colour spectra overlap of cone stimulation is affected → impaired ne interpretation of colours





NORMAL VISION



PROTANOPIA



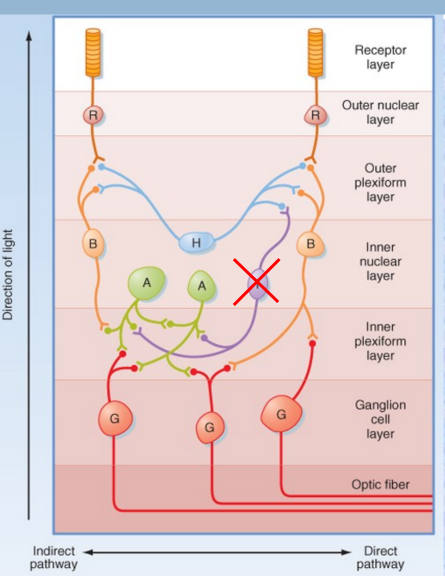




Retinal circuitry: review of cell types

- rods and cones synapse on <u>bipolar</u> <u>cells</u> and <u>horizontal cells</u>
- horizontal cells make lateral inhibitory synapses with surrounding <u>bipolar cells</u> or <u>photoreceptors</u>
- bipolar cells make synaptic connections with ganglion cells and amacrine cells
- amacrine cells transmit signals from <u>bipolar cells</u> to <u>ganglion cells</u> or to other <u>amacrine cells</u>
- ganglion cells transmit action potentials to the <u>brain</u> via the optic nerve
- **B&L** Figure 8-7

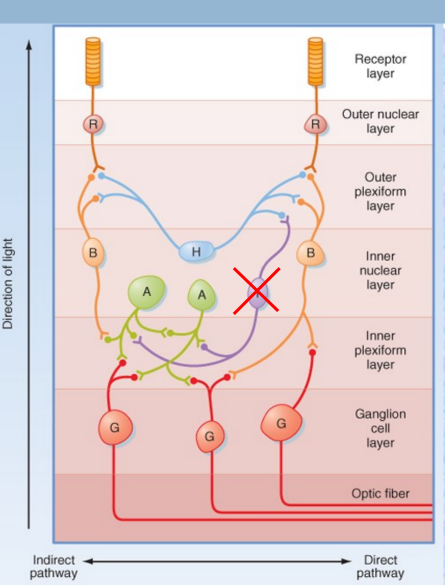
32



Retinal circuitry: key features

2 types of bipolar cells

- <u>On center</u>: hyperpolarized by glutamate
- Off center: depolarized by glutamate
- Bipolar and horizontal cells play a role in lateral inhibition
 - Important for increasing visual contrast
- Set up "surround" arrangement of ganglion cell receptive fields
 B&L Figure 8-7



Receptive fields

- Photoreceptor receptive fields include retinal area that, when stimulated by light, results in hyperpolarization of individual photoreceptor
 - Small and circular
- Ganglion cell receptive field size determined by
 - ganglion cell type
 - degree of convergence of photoreceptors and bipolar cells
 - and field type by retinal circuitry (lateral inhibition)
 - On-center/off-surround
 - Off-center/on-surround

Where in the retina is there is there a high degree of convergence?

Receptive fields

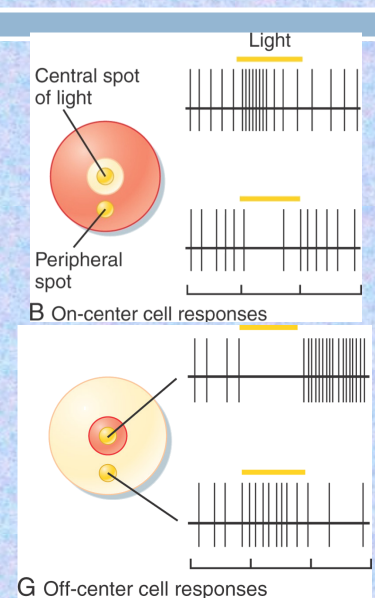
On-center/off-surround

- Light shines on center of ganglion cell receptive field → ganglion cell increases AP firing
- Light on surround region → decreased AP firing

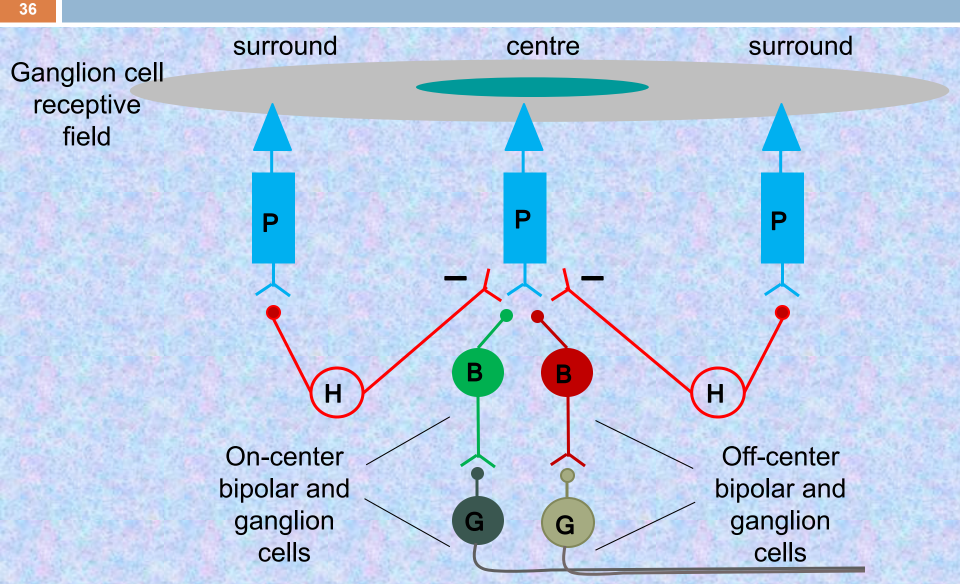
Off-center/on-surround

- Light on center → decreased AP firing
- Light on surround → increased AP firing

B&L Figure 8-8



Neural circuits of retinal receptive fields

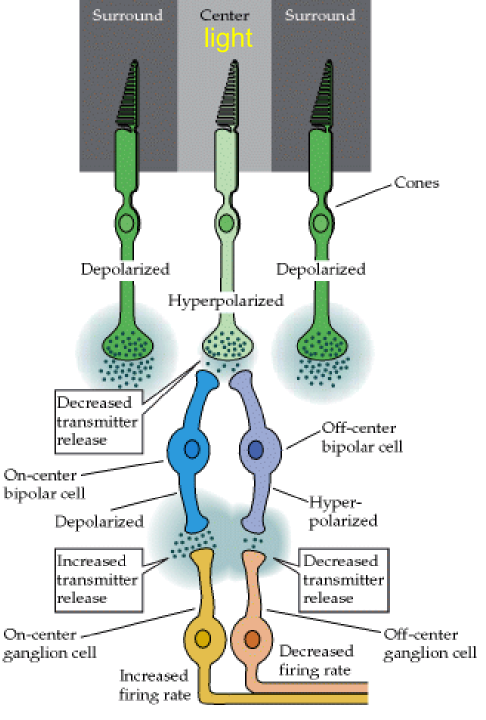


Neural Circuits of R Receptive Fields

37

Light stimulus on center:

- J glu release from central photoreceptor
 - □ ↓ inhibition of on-center bipolar cell → depolarization
 - □ ↑ NT release → <u>on-</u> <u>center ganglion cell</u> <u>excited</u>
 - □ less glu available to excite off-centre bipolar cell → hyperpolarization
 - □ ↓NT release → <u>off-center</u> <u>ganglion cell inhibited</u>

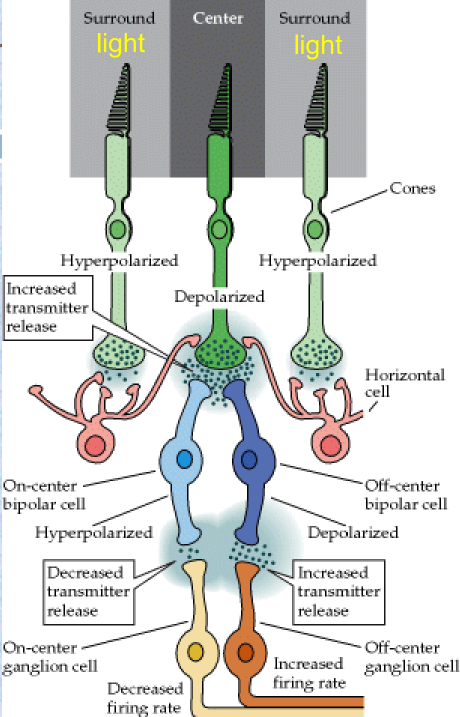


Neural Circuits of Retir Receptive Fields

38

Light stimulus on surround:

- J glu release from surround photoreceptor
 - □ ↓ excitation of horizontal cells →
 ↓ inhibitory NT released
 - □ ↓ inhibition of central photoreceptor → ↑ glu released
 - ↑ glu hyperpolarizes oncenter bipolar cell and depolarizes off-center bipolar cell
 - On-center ganglion cell inhibited, off-center ganglion cell excited



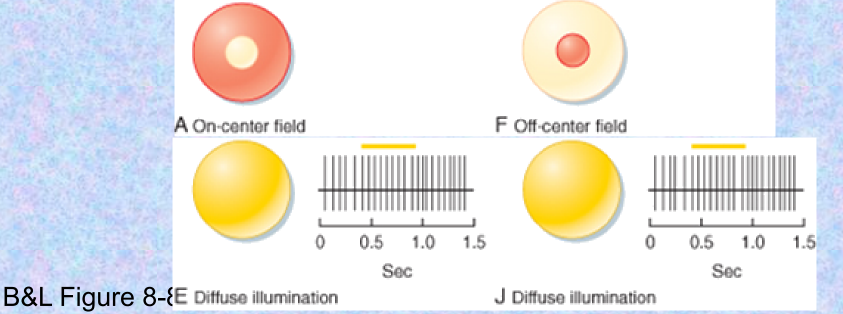
Retinal receptive fields: outcome

39

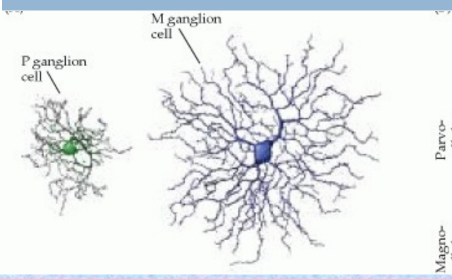
Surround arrangement and lateral inhibition allows ganglion cells to respond best to contrast borders in a visual scene

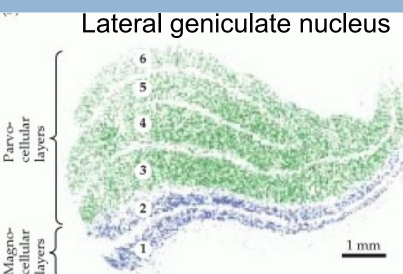
Ex. Reading dark letters against a white background

Respond only weakly to diffuse illumination



Ganglion cell types and projections





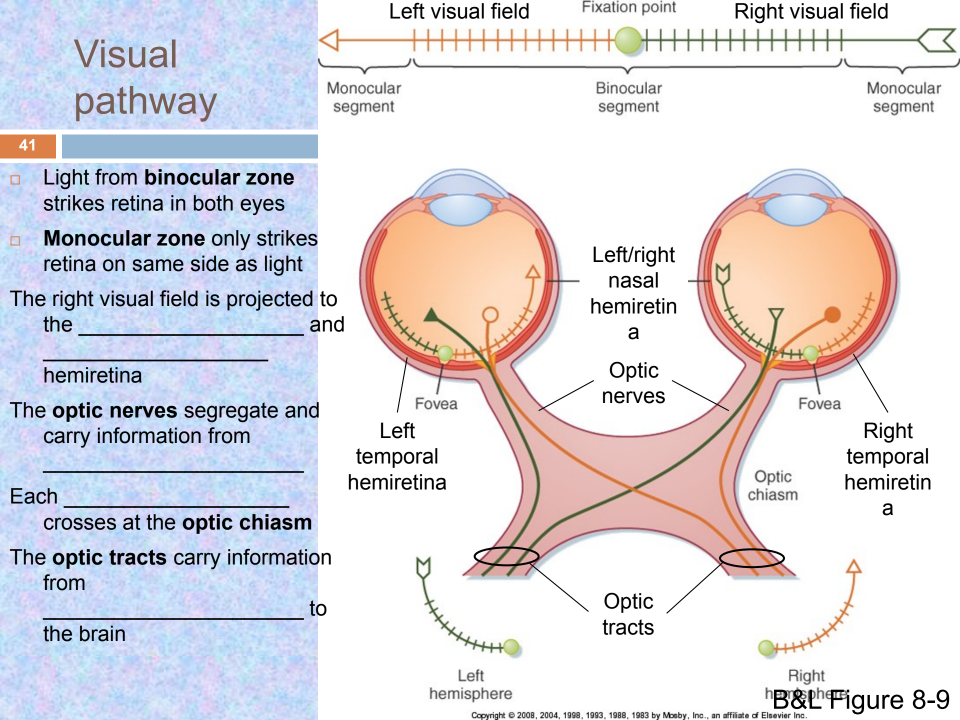
P cells

Π.

- Project to parvocellular layer of LGN
- Tonic firing, small surround receptive fields,
- Important for colour detection, form and detail of visual image

M cells

- Project to magnocellular layer of LGN
- Transient activity, large surround receptive fields
- Convey information about illumination and movement
- W cells
 - Resemble M cells, large diffuse receptive fields
 - Function is less clear



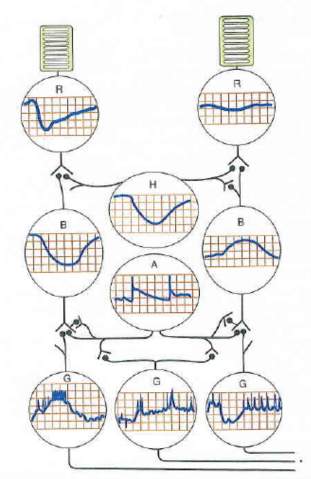
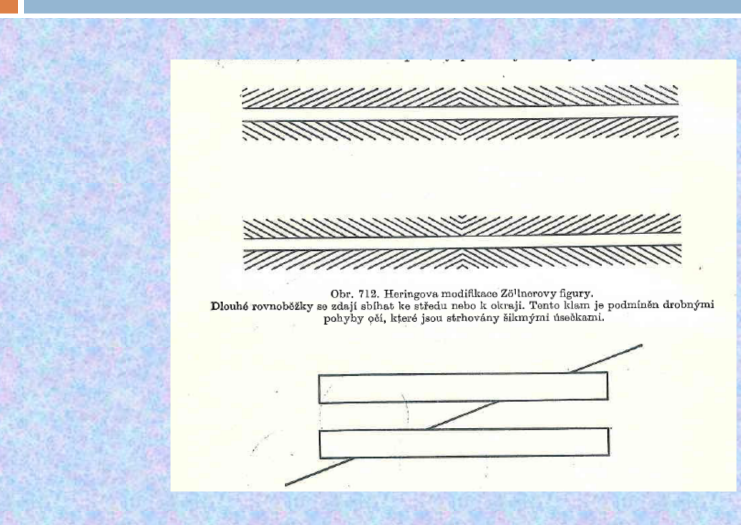
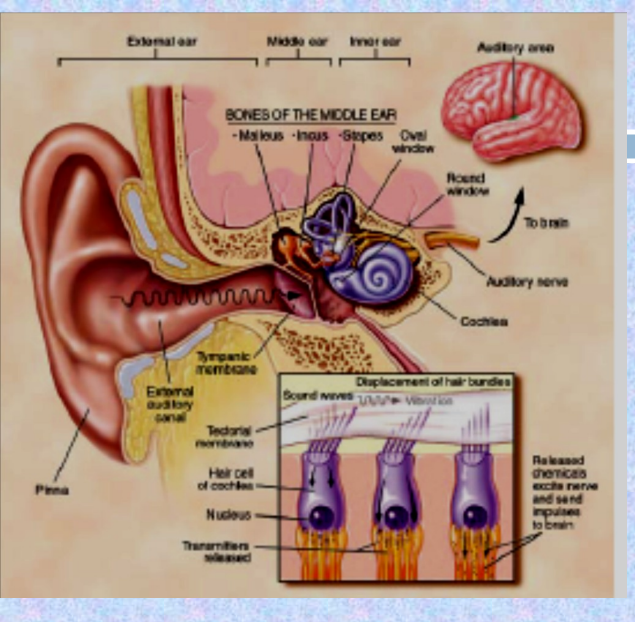


FIGURE 12–11 Intracellularly recorded responses of cells in the retina to light. The synaptic connections of the cells are also indicated. The eye is unique in that the receptor potentials of the photoreceptors and the electrical responses of most of the other neural elements in the retina are local, graded potentials. The rod (R) on the left is receiving a light flash, whereas the rod on the right is receiving steady, low-intensity illumination. The responses of rods and horizontal cells (H) are hyperpolarizing, responses of bipolar cells (B) are either hyperpolarizing or depolarizing, and amacrine (A) cells produce depolarizing potentials and spikes that may act as generator potentials for propagated spikes of ganglion cells (G). (Reproduced with permission from

42

Optical illusion

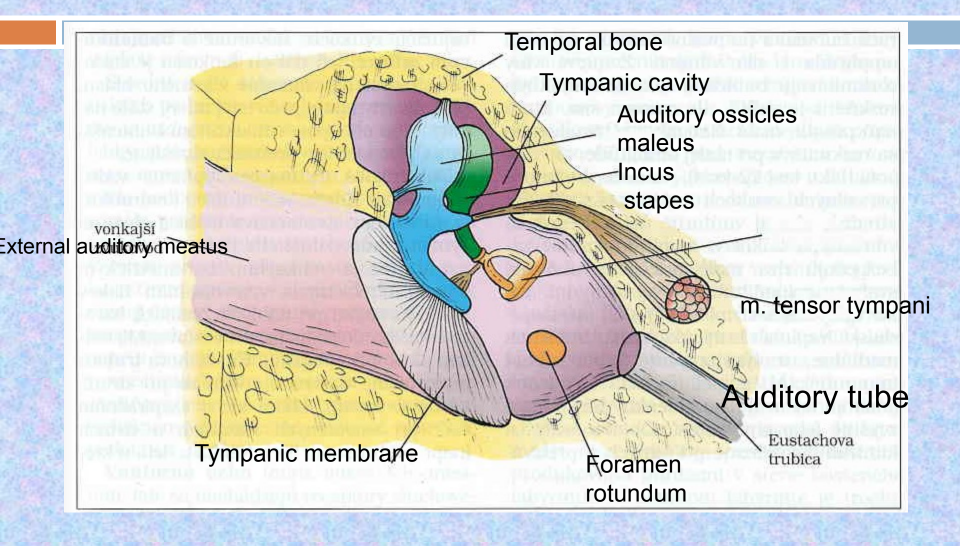




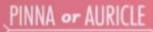
At the **AUDITORY system** is an array of miniature acoustical detectors packed into a space **no larger than a pea.**

These detectors can faithfully transduce vibrations as small as the diameter of an atom, and they can respond a **thousand times faster** than visual photoreceptors

Outer ear / middle ear



Middle ear - transport acustic stimuli by air

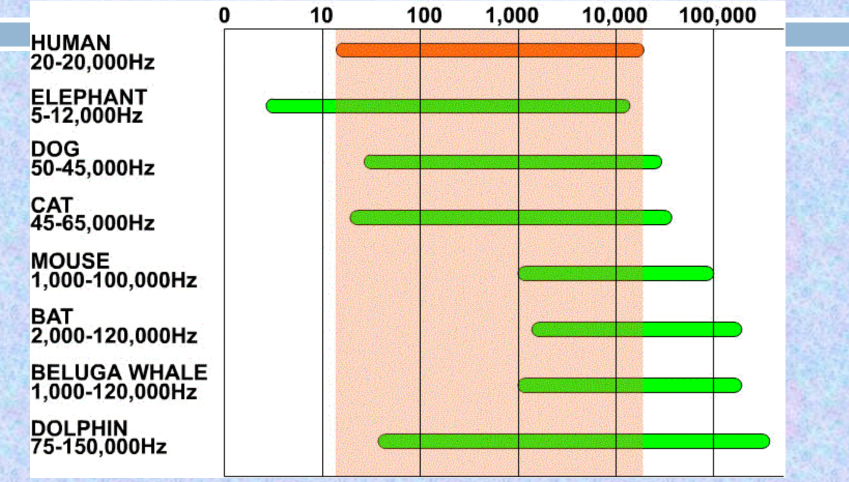


catches sound waves, and passes them along deeper into the ear

EXTERNAL ACOUSTIC MEATUS

auditory canal

The Audible Spectrum



Middle ear: Impedance



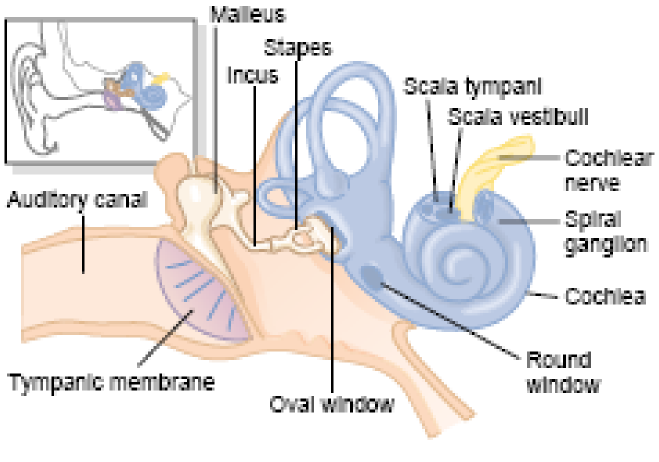
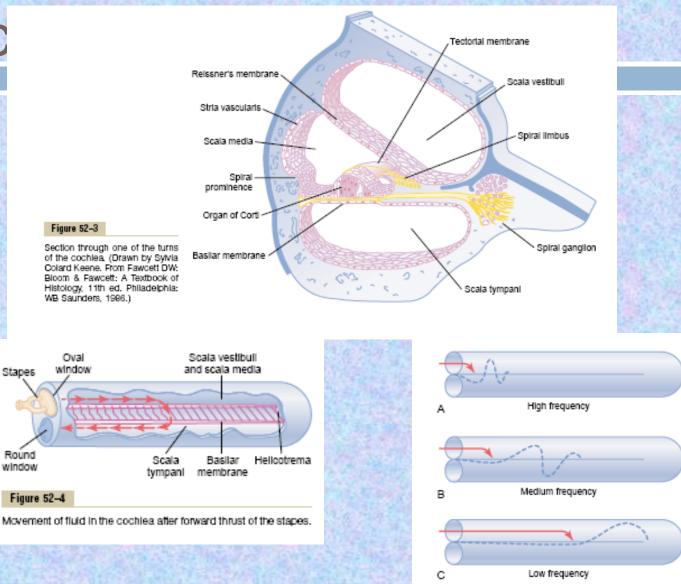


Figure 52-1

Tympanic membrane, ossicular system of the middle ear, and inner ear.

Transmission of Sound Waves in

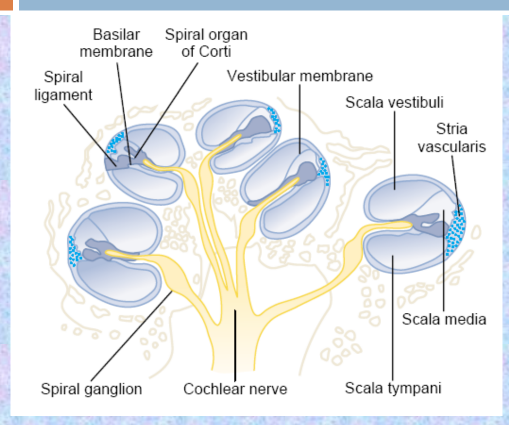
the Coc

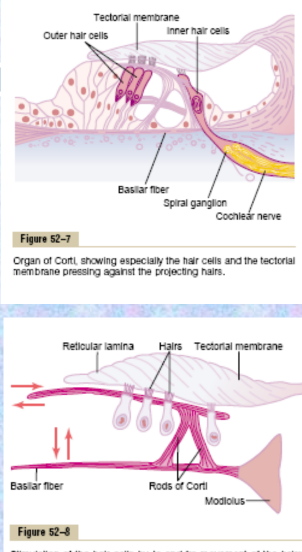


"Traveling waves" along the basilar membrane for high-, medium-, and low-frequency sounds.

Figure 52-5

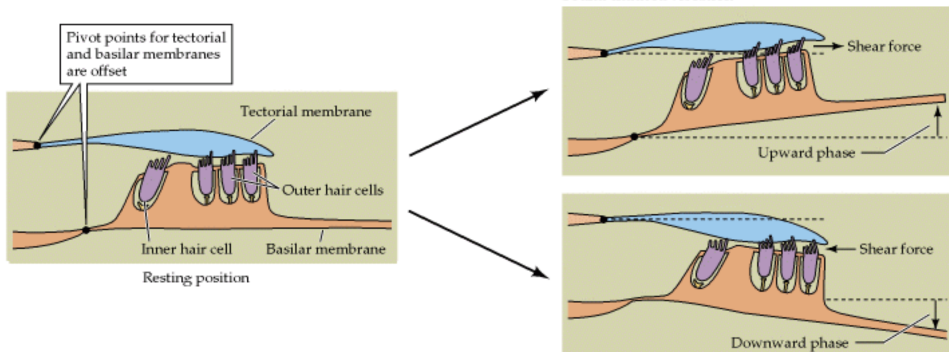
The Organ of Corti





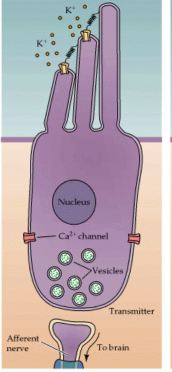
Stimulation of the hair cells by to-and-fro movement of the hairs projecting into the gel coating of the tectorial membrane.

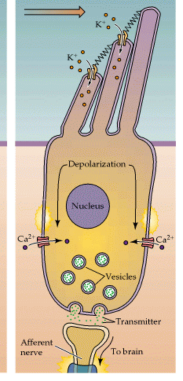
Excitation of the Hair Cells

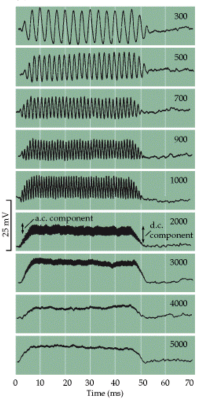


Sound-induced vibration

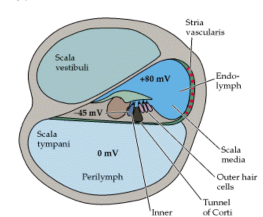
Hair Cell Receptor Potentials



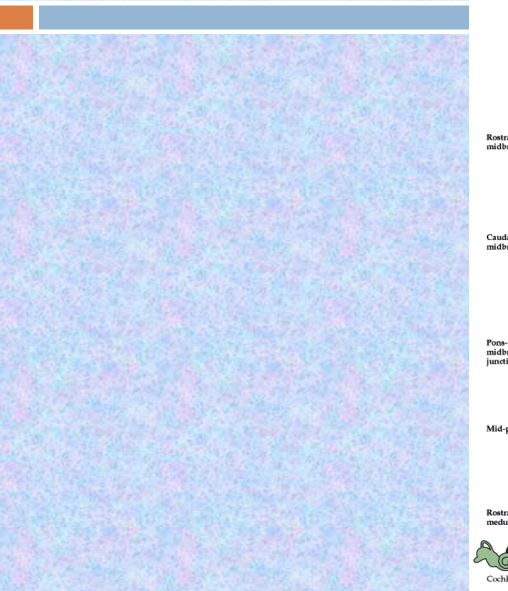


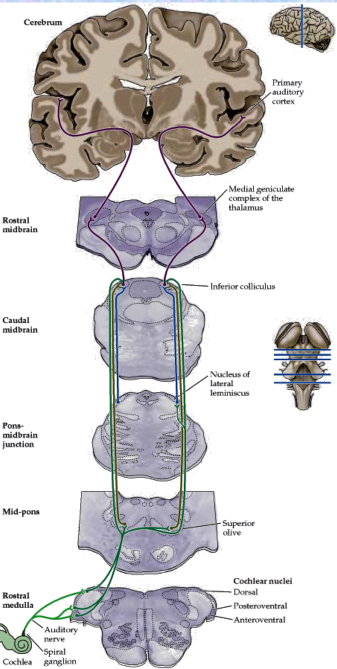


(D)

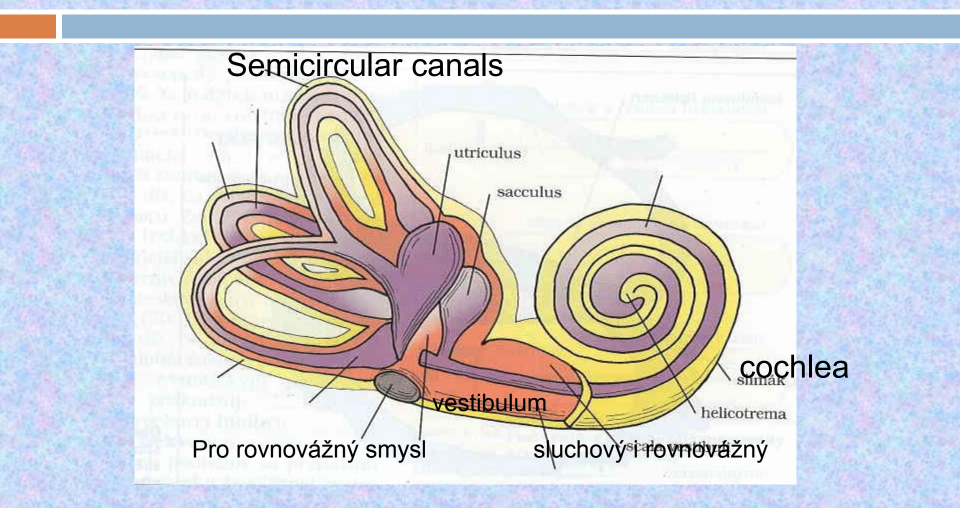


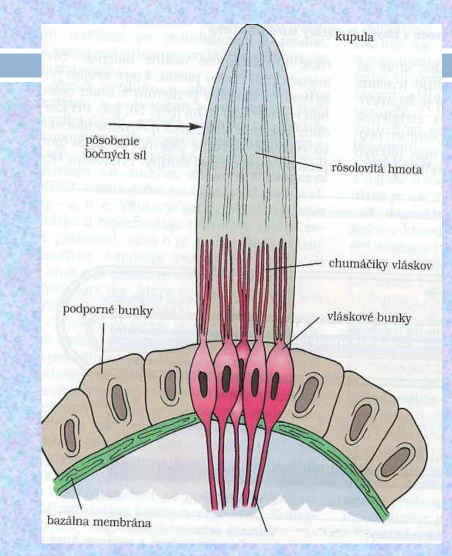
Auditory Pathwa



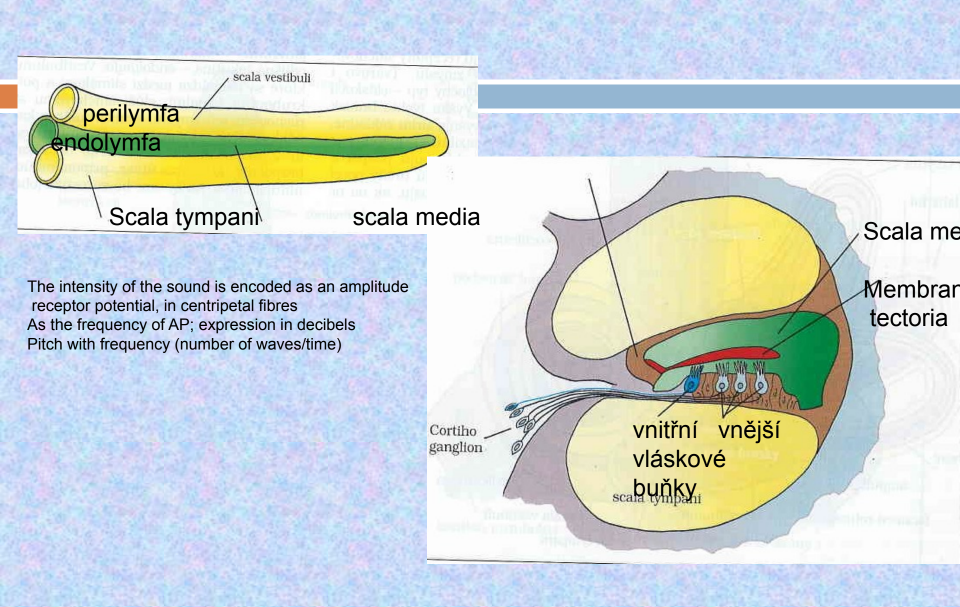


Inner ear

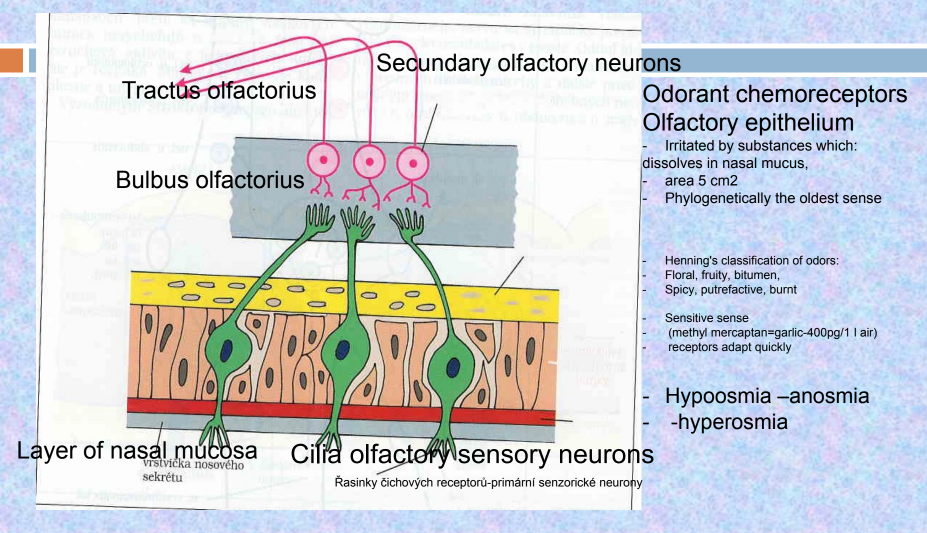




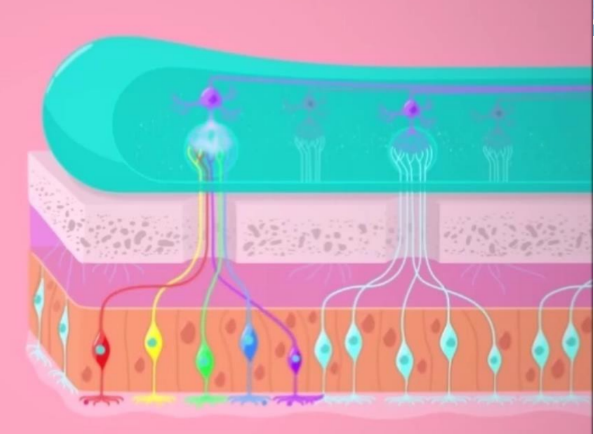
Structure of hair cell Lateral movement stimulate receptor – receptor potencial

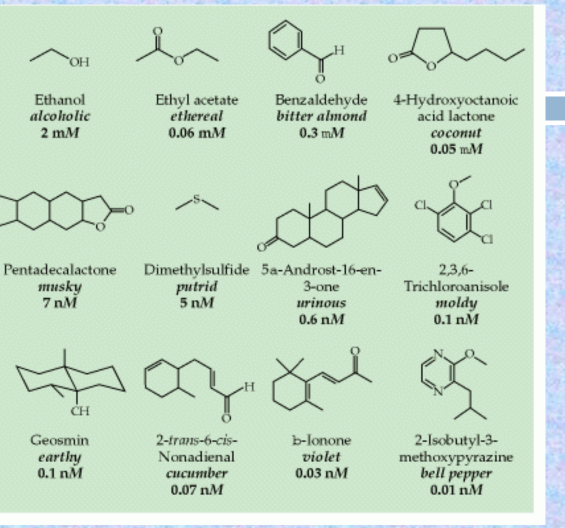


Smell



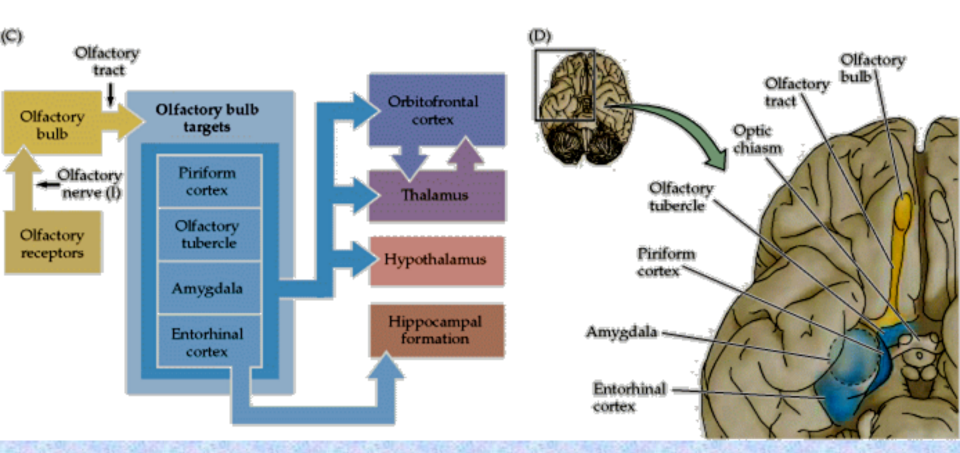
The Olfactory Epithelium and



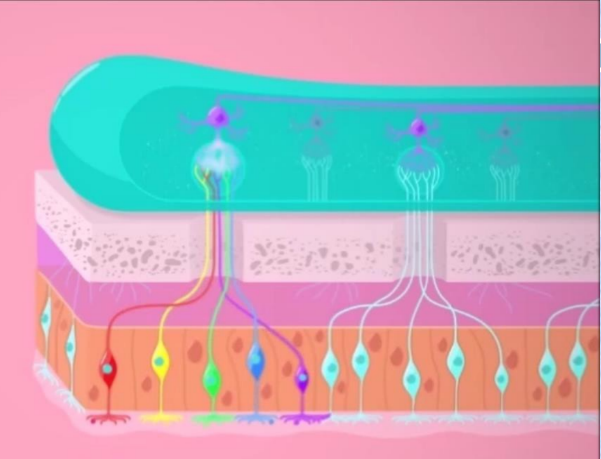


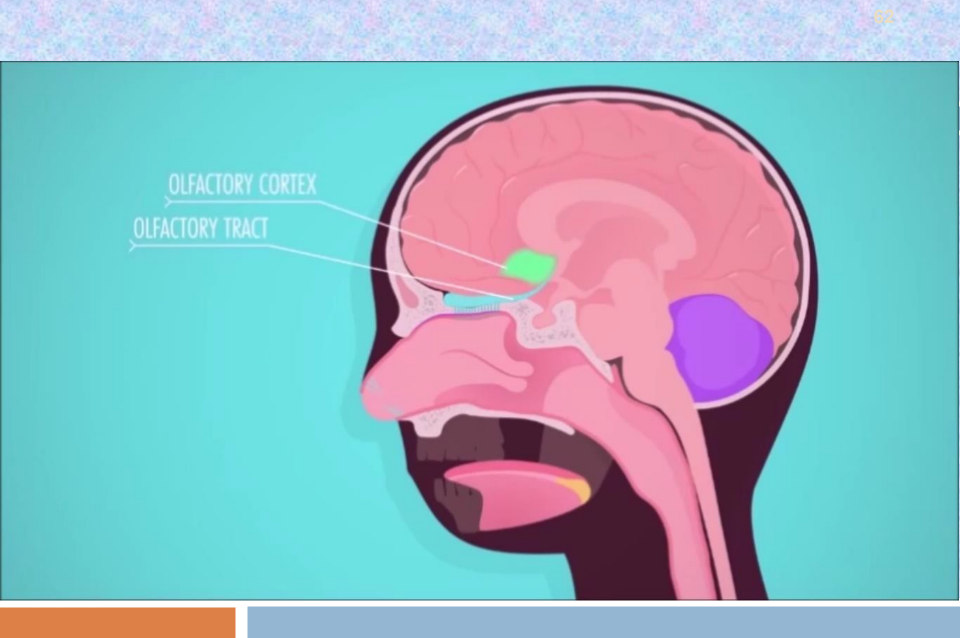


Olfactory System

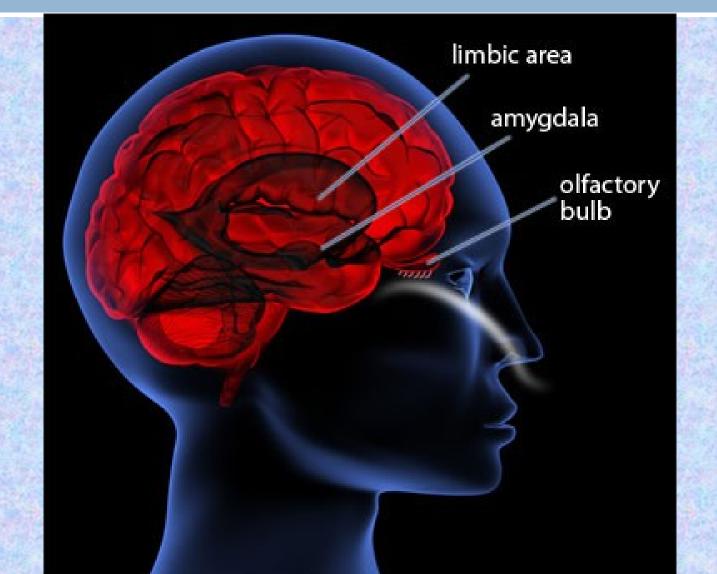


(C) Diagram of the basic pathways for processing olfactory information.(D) Central components of the olfactory system.

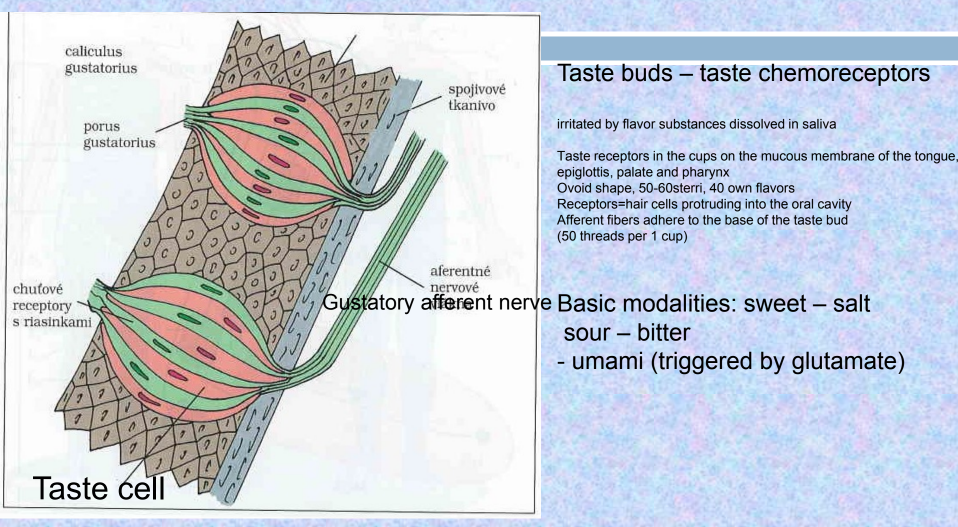




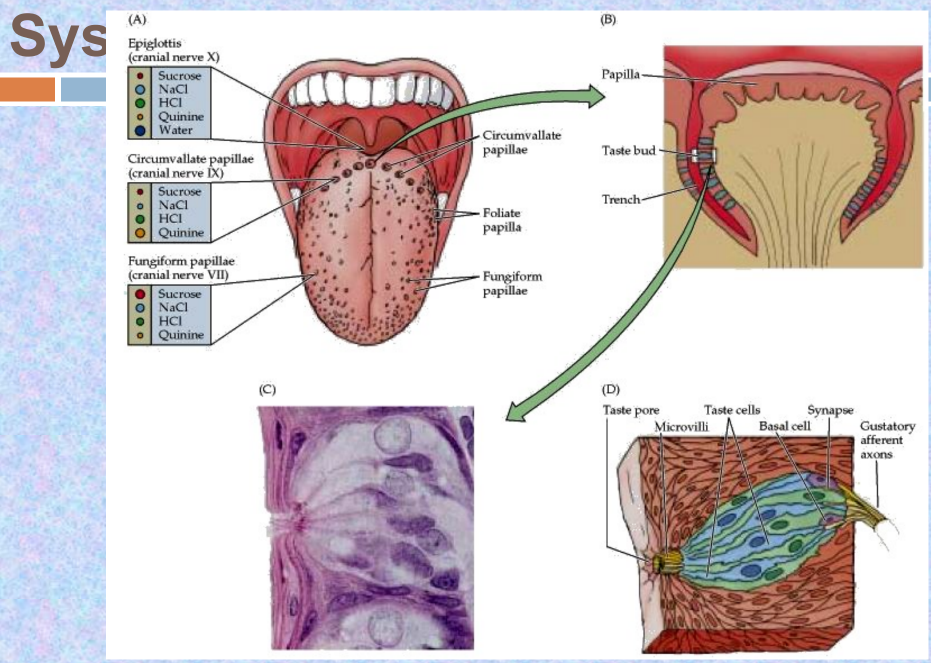
Physiological and Behavioral Responses to Odorants



Taste



The Organization of the Taste



Central Processing of Taste

