

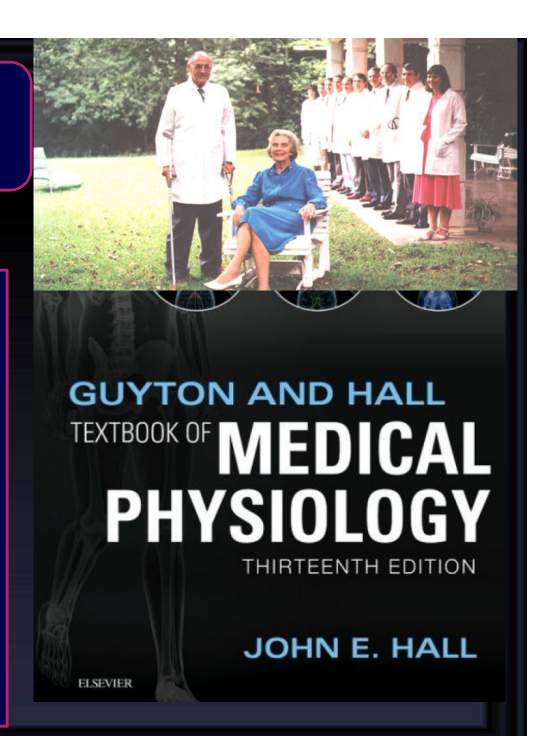
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REMEMBER

- These handouts will facilitate what you have to study and are not an alternative to your text book.
- The main source of this Lectures is from Guyton & Hall 13th Edition
- Ch46-Pages 546-561
- Ch47-Pages 568-574





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OBJECTIVES

At the end of this lecture the student should be able to:

- Define synapses and enumerate functions of synapses.
- Classify types of synapses: anatomical & functional.
- Draw and label structure of synapses
- Describe Synaptic transmission & neurotransmitters
- Explain the fate of neurotransmitters.
- Explain electrical events at synapses (EPSPs & IPSPs).
- Elaborate Properties and Patterns of synaptic transmission in neuronal pools
- Explain factors affecting synaptic transmission

Amazing Facts about human Nervous System On an average, humans experience around 70,000 thoughts each day

ree-year-old child's brain has an estimated one quadrillion (10¹⁵) synapses in total. The number, however, decreases with age, and an average adult has between 100 to 500 trillion synapses

Loss of blood supply to brain for 8-10 seconds can lead to unconsciousness

Everyone dreams, even blind people

There are no nociceptors in the brain, so the brain itself cannot feel pain

Most *amazing* thing is that the brain in our head isn't our only brain. There's a "second brain" in our intestines that contain 100,000 neurons.

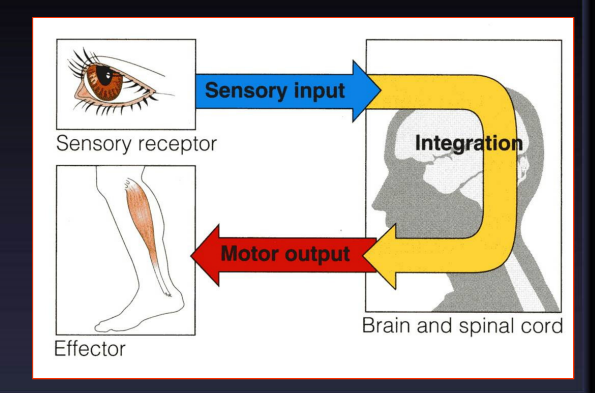
Two percent of the body's weight, the human brain consumes 20% of the energy used by the entire body 15% of all cardiac output, 20%t of total body oxygen consumption, and 25% of total body glucose consumption

The human brain has a capacity of 2.5 petabytes, that is 2.5 million gigabytes

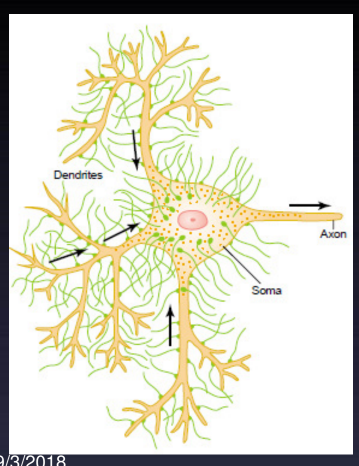
The brain is so plastic that it can rewire itself

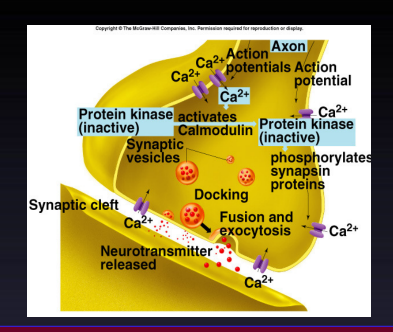
HOW BRAIN FUNCTIONS?

- Collection of sensory input
- Central Integration
- Motor output



The Synapse **Synaptic Transmission/neurotransmitters**





Information is transmitted in the central nervous system mainly in the form of nerve action potentials, called **NERVE** IMPULSES, through a succession of neurons, one after another.

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The Synapse

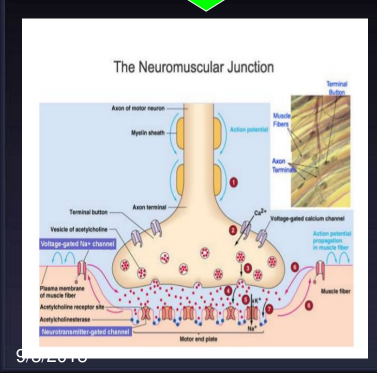
- A synapse is the connection between a neuron and a second cell.
- In the CNS, this other cell is also a neuron.
- In the PNS, the other cell may be either a neuron or an effector cell eg; gland or muscle

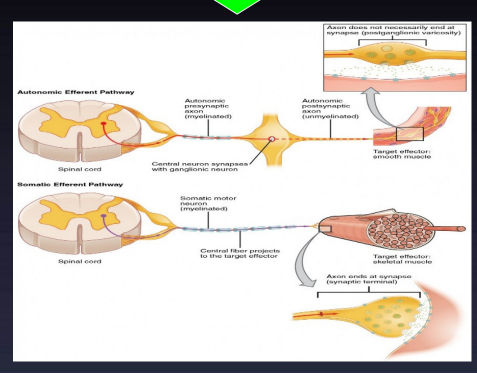
A junction where the axon or some other portion of one cell (presynaptic cell) terminates on the dendrites, soma, or axon of another neuron (post synaptic cell).

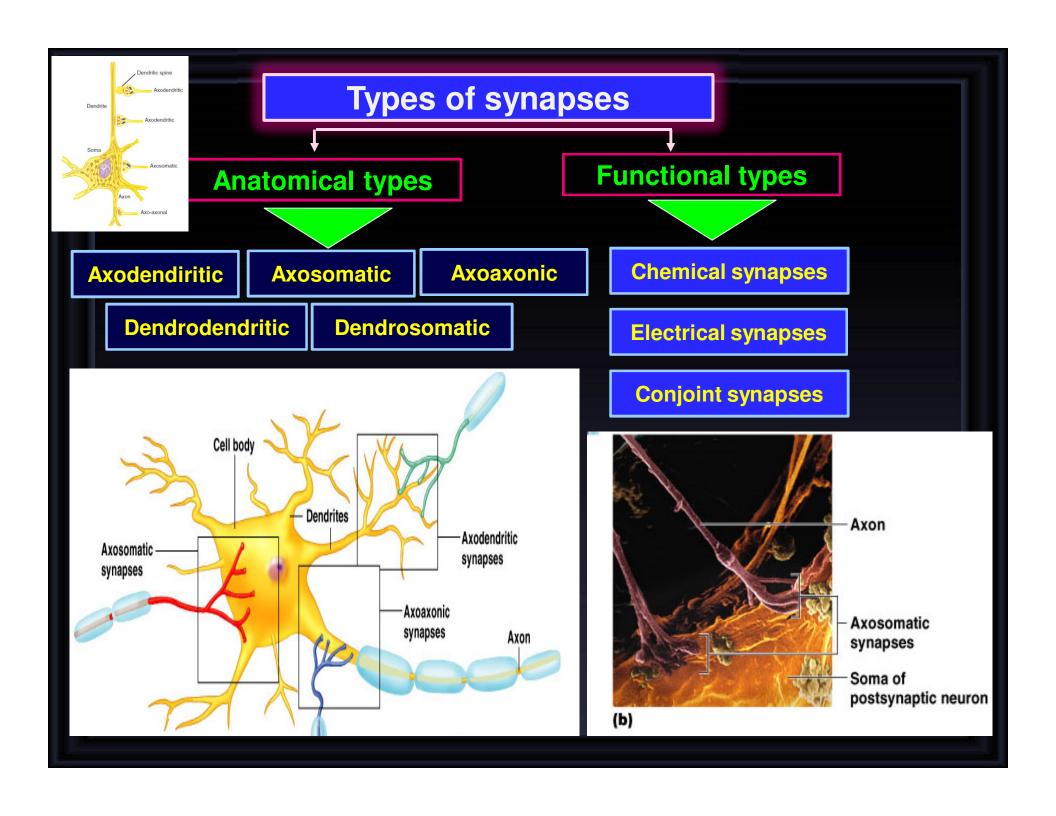
Junctions outside the CNS

Neuromuscular junction

Contact between: autonomic neurons and smooth, cardiac muscles and any other effector cells.



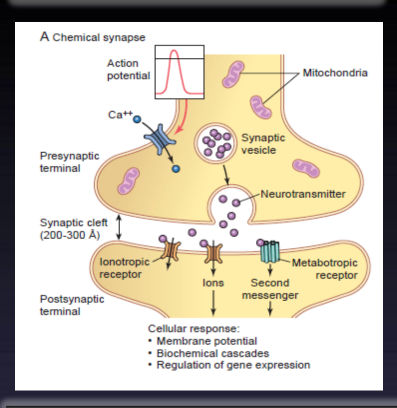




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Functional Types

Electrical Synapse



Chemical Synapse

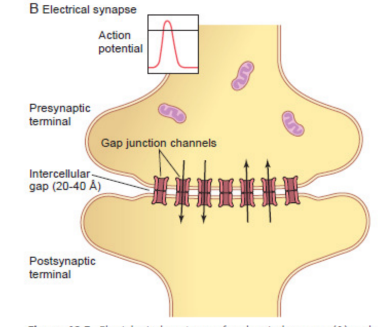
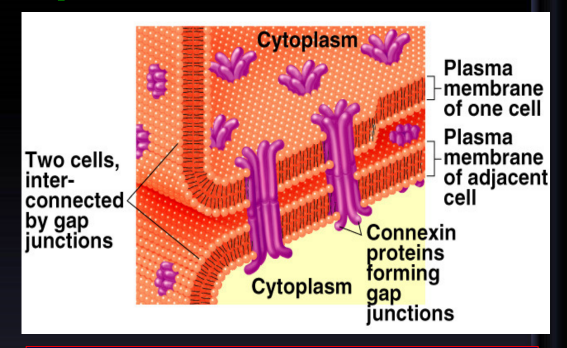


Figure 46-5. Physiological anatomy of a chemical synapse (A) and an electrical synapse (B).

Conjoint synapse: Both electrical and chemical. **Examples** → neurons in lateral vestibular nucleus.

Electrical Synapse

- Impulses can be regenerated without interruption in adjacent cells.
- Gap junctions:
 - Adjacent cells electrically coupled through a channel.
 - Each gap junction is composed of 12 connexin proteins.



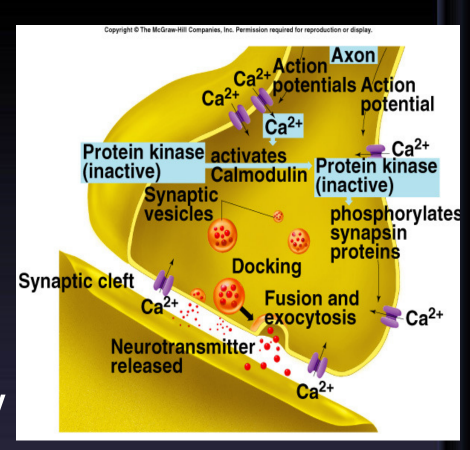
• Examples:Smooth and cardiac muscles, brain, and glial cells.

The bidirectional transmission of electrical synapses permits them to help coordinate the activities of large groups of interconnected neurons. Promotes synchronous firing of a group of interconnected neurons. For example, in

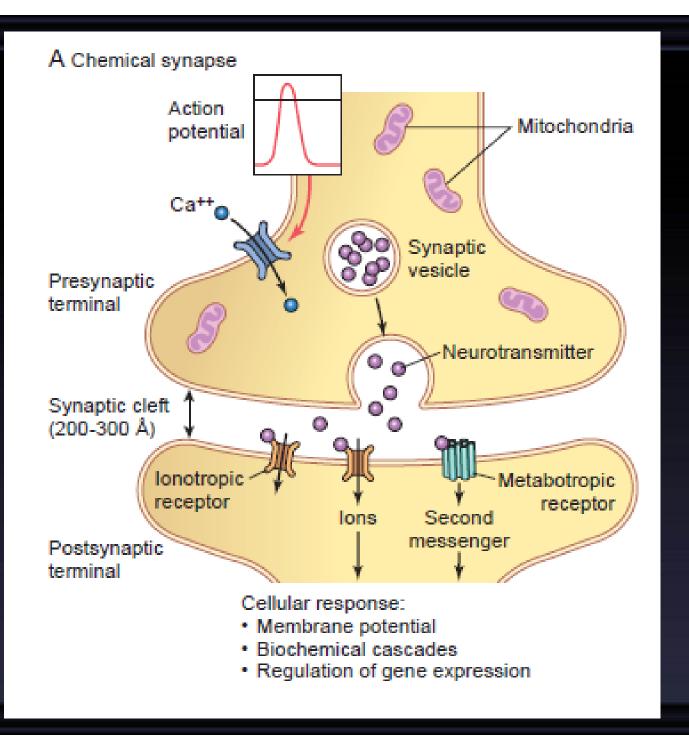
Mental attention, Emotions and Memory
Arousal from sleep

Chemical Synapse

- Terminal bouton is separated from postsynaptic cell by synaptic cleft.
- NTs are released from synaptic vesicles.
- Vesicles fuse with axon membrane and NT released by exocytosis.
- Amount of NTs released depends upon frequency of AP.



"One-Way" Conduction at Chemical Synapses



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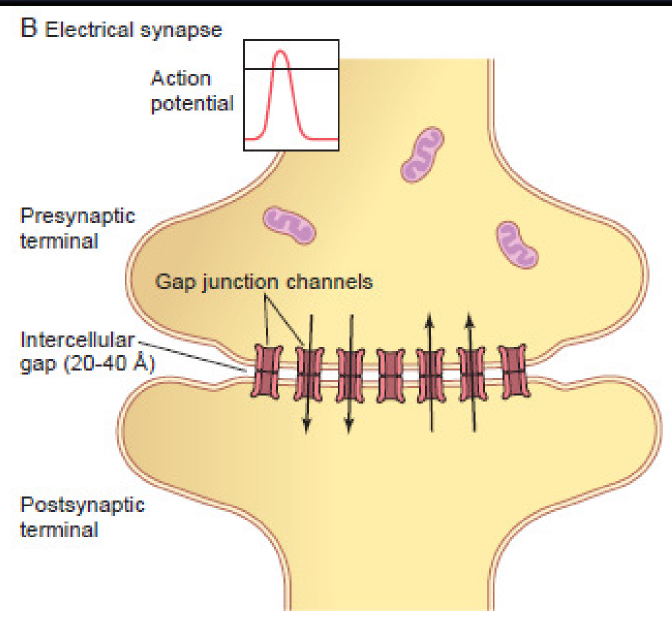


Figure 46-5. Physiological anatomy of a chemical synapse (A) and an electrical synapse (B).

Synaptic Transmission (Events)

- NT release is rapid because many vesicles form fusion-complexes at "docking site."
- AP travels down axon to bouton.
- VG Ca²⁺ channels open.
 - Ca²⁺ enters bouton down concentration gradient.
 - Inward diffusion triggers rapid fusion of synaptic vesicles and release of NTs.
- Ca²⁺ activates calmodulin, which activates protein kinase.
- Protein kinase aid in the fusion of synaptic yesicles.

Synaptic Transmission (continued)

- NTs are released and diffuse across synaptic cleft.
- NT (ligand) binds to specific receptor proteins in postsynaptic cell membrane.
- NT effects are produced

FATE OF NEUROTRANSMITTER

<u>Diffusion</u> out of synaptic cleft into surrounding fluid <u>Enzymatic destruction</u> e.g. Ach esterase for Ach <u>Active transport</u> back into pre-synaptic terminal itself e.g. norepinephrine

Transmitter Substance acts on the Postsynaptic Neuron via "Receptor Proteins"

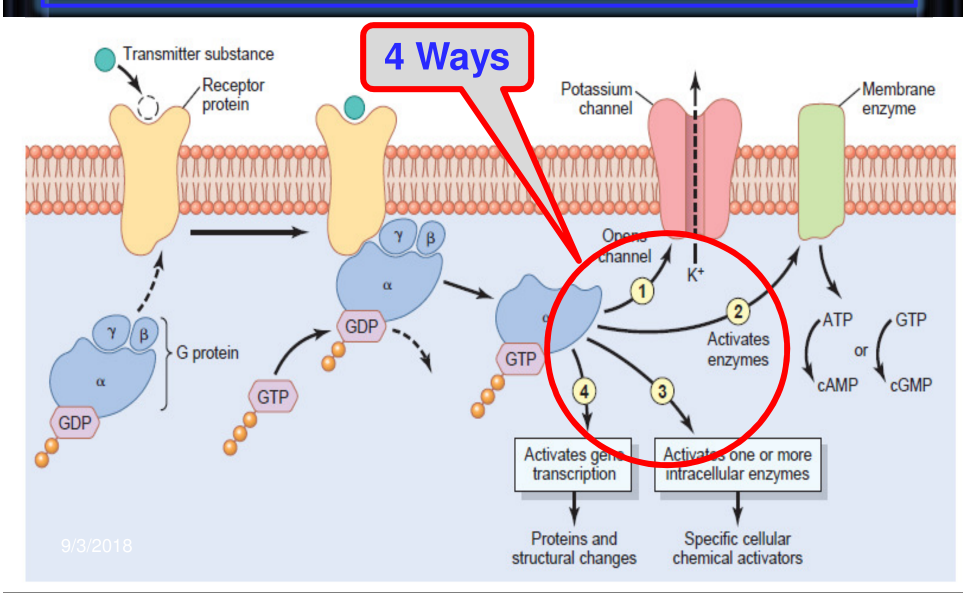
Have binding & intracellular component

Receptor activation acts in one of two ways:

(1) By gating ion channels directly and allowing passage of specified types of ions through the membrane (ionotropic receptors)

(2) By activating a "second messenger" that is not an ion channel but a molecule that protrudes into the cell cytoplasm and activates one or more substances inside the postsynaptic neuron (metabotropic receptors)

"Second Messenger" System in the Postsynaptic Neuron acts in....



EPSP & IPSP at Chemical Synapses

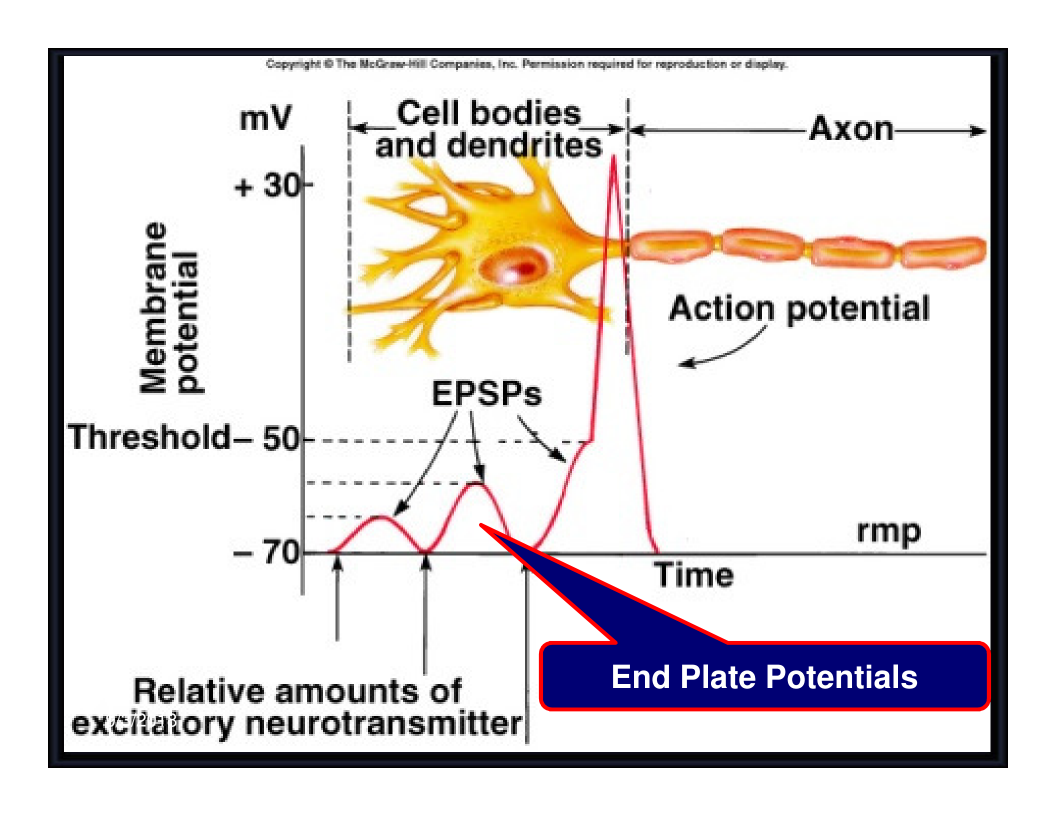
EPSP (excitatory postsynaptic potential):

- 1. Opening of Na channels to threshold level (Most Common).
- 2. ♥ conduction through Cl or K channels, or both.
- 3. Various changes in the internal metabolism of the postsynaptic neuron to excite or, in some instances, to ↑ excitatory membrane receptors or ↓ inhibitory membrane receptors.

IPSP (inhibitory postsynaptic otential):

- 1. Opening of Cl ion channels through the postsynaptic neuronal membrane.
- 2.

 in conductance of K ions out of the Neuron
- 3. Activation of receptor enzymes that inhibit cellular metabolic functions that ↑ inhibitory membrane receptors or ♥ excitatory membrane receptors.



Synaptic properties

- 1. One-way conduction: Synapses generally permit conduction of impulses in one-way i.e. from presynaptic to post-synaptic neuron "Bell- Magendie law".
- 2. Synaptic delay: 0.5 ms for transmission across one synapse
- 3. Synaptic inhibition: 4 Types: Direct, Indirect, Reciprocal, Inhibitory interneuron
- 4. Summation: Spatial & Temporal
- 5. Convergence and divergence:
- 6. Fatigue (synaptic depression):

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2. Synaptic delay (Central Delay)

Is the minimum time required for transmission across the synapse. It is 0.5 ms for transmission across one synapse.

This time is taken by.....

- Discharge of transmitter substance by pre-synaptic terminal
- Diffusion of transmitter to post-synaptic membrane
- Action of transmitter on its receptor
- Action of transmitter to \(\bar\) membrane permeability
- Increased diffusion of Na+ to ↑ post-synaptic potential

Clinical Importance is that we can know number of synapses involved in neuronal pathways by time lag

- A. Direct inhibition: Occurs when an inhibitory neuron (releasing inhibitory substance) acts on a post-synaptic neuron leading to → hyperpolarization due to opening of Cl⁻ [IPSPs] and/or K+ channels. Example: Glycine at the level of the spinal cord to block pain impulses.
- **B. Indirect Inhibition:** (Pre-synaptic inhibition): This happens when an inhibitory synaptic knob lie directly on the termination of a pre-synaptic excitatory fiber.
- The inhibitory synaptic knob inhibits the release of excitatory transmitter from the pre-synaptic fiber. e. g. GABA (Pain modification)
- C. Reciprocal inhibition: Inhibition of antagonist muscle whine agonist is excited.
- **D. Inhibitory interneuron (Renshaw cells):** Negative feedback inhibitory interneuron of a spinal motor neuron .

PATTERNS OF SYNAPTIC TRANSMISSION IN NEURONAL POOLS

Divergence into multiple tracts

Figure 47-11. "Divergence" in neuronal pathways. *A*, Divergence within a pathway to cause "amplification" of the signal. *B*. Divergence

1-DIVERGENCE

2-CONVERGENCE

3-REVERBERATORY CIRCUIT

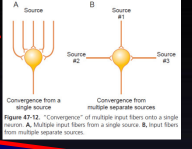
4-AFTER-DISCHARGE

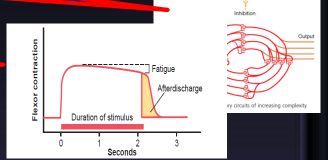
5-IRRADIATION

6-RECIPROCAL INHIBITION

7-LOCAL SIGN







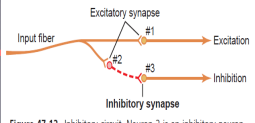


Figure 47-13. Inhibitory circuit. Neuron 2 is an inhibitory neuron.

PATTERNS OF SYNAPTIC TRANSMISSION IN NEURONAL POOLS

4-AFTER-DISCHARGE

A prolonged maintained output discharge of AHCs called after discharge, lasting a few milliseconds or many minutes after the incoming signal is over. Due to:

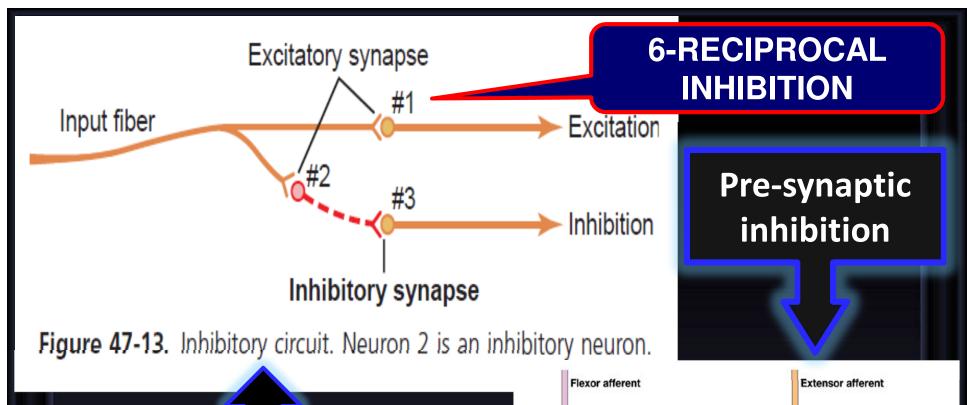
- 1- EPSP can continue to excite the neuron to transmit (a series of continuous repetitive discharges).
- 2- Reverbrating circuits: Presence of reverberating circuit restimulate AHCs

5- IRRADIATION

Spread of impulses up & down to different segments and motor neurons in the spinal cord Ex; A strong stim in sensory afferent irradiate to many segments of S.C due to <u>divergence</u>

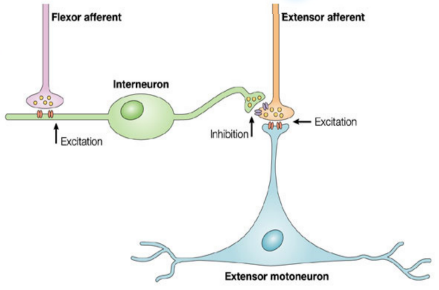
7- LOCAL SIGN

The response to the stimulus by the body will be determined by the location of the stimulus on the body – i.e., a noxious stimulus to the lateral land the lateral lat



Neuronal Circuit With Both Excitatory and Inhibitory Output Signals

This type of circuit is characteristic for controlling all antagonistic pairs of muscles, and it is called the reciprocal inhibition circuit.

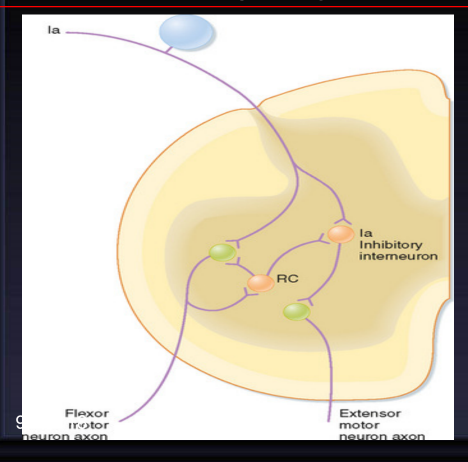


Nature Reviews | Neuroscience

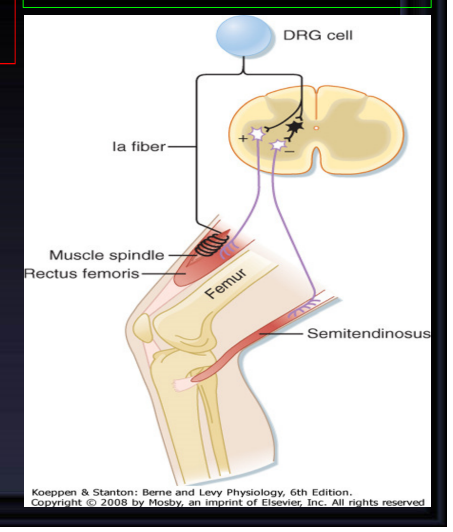
D. Inhibitory interneuron (Renshaw cells)

Negative feedback inhibitory interneuron of a spinal motor neuron;

Send inhibitory cells that transmit inhibitory signals to the surrounding motor neurons → lateral inhibition → Sharpens Signals



C. Reciprocal inhibition: Inhibition of antagonist muscle whine agonist is excited.



Reverberatory (Oscillatory) Circuit

Cause of Signal Prolongation.

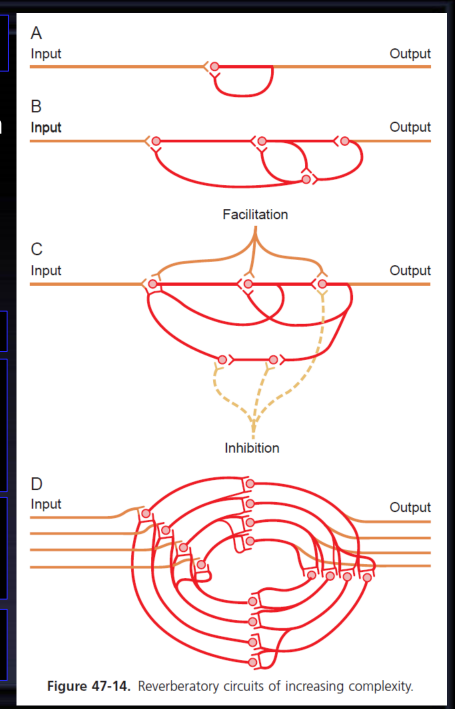
- ☐ caused by positive feedback within the neuronal that re-excite the input of the same circuit.
- ☐ Once stimulated, the circuit may discharge repetitively for a long time called long term potentiation

The simplest Fig A, involves single neuron

Fig B shows additional neurons in the feedback circuit, which causes a longer delay between initial discharge and the feedback signal.

Fig C shows a more complex system in which both facilitatory and inhibitory fibers impinge on the reverberating circuit.

Fig D shows reverberating pathways with 9/3/2018 parallel fibers.



The cause of this sudden cessation of reverberation is fatigue of synaptic junctions in the circuit. Fatigue beyond a certain critical level lowers the stimulation of the next neuron in the circuit below threshold level so that the circuit feedback is suddenly broken.

Synaptic fatigue

short-term (synaptic depression), is an activity-dependent form of short term synaptic plasticity that results in the temporary inability of neurons to fire and therefore transmit an input signal.

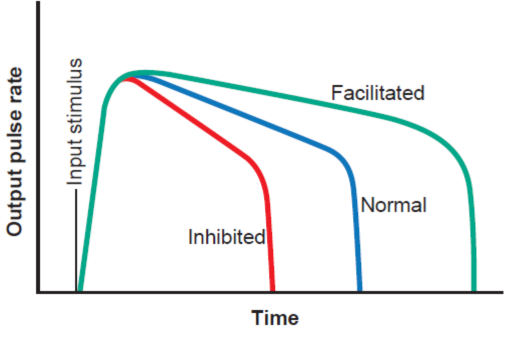


Figure 47-15. Typical pattern of the output signal from a reverberatory circuit after a single input stimulus, showing the effects of facilitation and inhibition.

Almost these exact patterns of output signals are recorded from the motor nerves exciting a muscle involved in a flexor reflex after pain stimulation of the foot (as shown later in **Figure 47-18**).

TRANSMISSION OF SIGNALS OF DIFFERENT INTENSITY IN NERVE TRACTS BY SUMMATION

Spatial Summation

Increasing signal strength is transmitted by using progressively greater numbers of fibers.

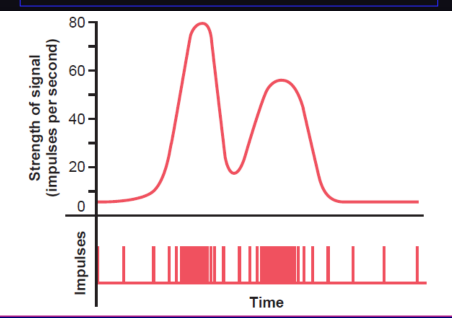
Weak stimulus Strong stimulus

Figure 47-7. Pattern of stimulation of pain fibers in a nerve leading from an area of skin pricked by a pin. This pattern of stimulation is an example of *spatial summation*.

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Temporal Summation

Transmitting signals of increasing strength is by increasing the frequency of nerve impulses in each fiber



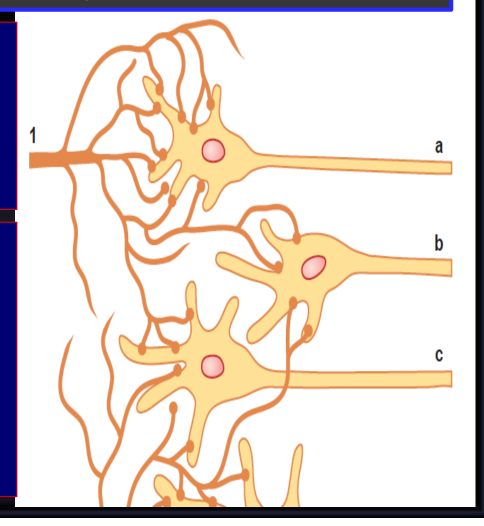
If EPSPs in a pre-synaptic knob are successively repeated without significant delay so the effect of the previous stimulus is summated to the next.

The neuronal area stimulated by each incoming nerve fiber is called its stimulatory field. Large numbers of the terminals from each input fiber lie on the nearest neuron in its "field," & fewer terminals lie on the neurons farther away.

Discharge zone of the incoming fiber, also called the excited zone (a with suprathreshold stimulus)

To each side, the neurons are facilitated but not excited, and these areas are called the facilitated zone, also called the subthreshold zone or subliminal zone. (b & c not enough to cause excitation)

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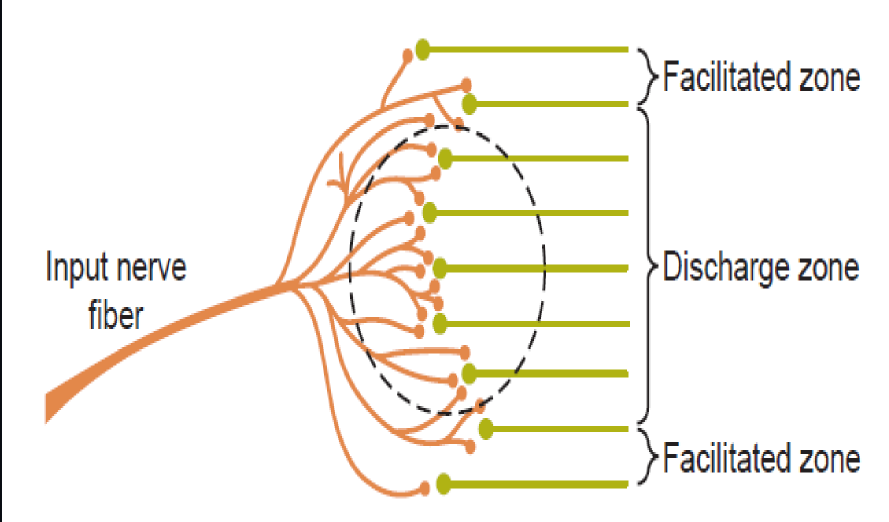


Figure 47-10. "Discharge" and "facilitated" zones of a neuronal pool.

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In divergence weak signals entering a neuronal pool are amplified. Two major types

Divergence into multiple tracts

Divergence into multiple tracts Amplifying type Α В

Figure 47-11. "Divergence" in neuronal pathways. A, Divergence within a pathway to cause "amplification" of the signal. B, Divergence into multiple tracts to transmit the signal to separate areas.

Eg; dorsal columns: of the spinal cord takes two courses in the lower part of the brain: (1) into the cerebellum and (2) on through the lower regions of the brain to the thalamus and cerebral cortex.

변혈?연8rticospinal pathway

Divergence in same tract

Convergence means signals from multiple inputs uniting to excite a single neuron



Action potentials converging on the neuron from multiple terminals provide enough spatial summation to bring the neuron to the threshold required for discharge.

Convergence can also result from input signals (excitatory or inhibitory) from multiple sources 9/3/2018

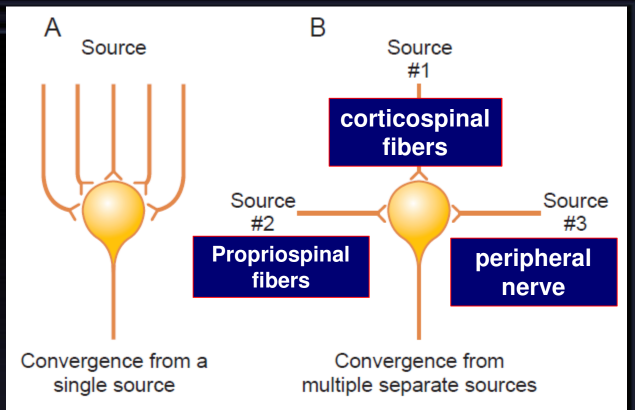


Figure 47-12. "Convergence" of multiple input fibers onto a single neuron. **A,** Multiple input fibers from a single source. **B,** Input fibers from multiple separate sources.

the interneurons of the spinal cord receive converging signals from

- (1) peripheral nerve fibers entering the cord,
- (2) Propriospinal fibers passing from one segment of the cord to another,
- (3) corticospinal fibers from the cerebral cortex, and
- (4) several other long pathways descending from the brain into the spinal cord.

From interneurons converge on the anterior motor neurons to control muscle function. By summation

Factors affecting synaptic transmission

Alkalosis:

neuronal excitability;

Causes cerebral epileptic seizures (Increased excitability cerebral neurons) e. g. overbreating in person with epilepsy

Acidosis:

Drugs:

Caffeine found in coffee, tea, strychnine, theophylline and theobromine increases neuronal excitability, by reducing the threshold for excitation of neurons.

Hypoxia:

Depression of neurons

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