### Basic mechanisms of cell communication:

Neuronal communication

Quick answer fast muscular and secretory cells

Endocrine communication:

Slow response durable Cell functions, metabolism

Neuroendocrine communication









# Synaptic Transmission

# Synapses: what are they?

Specialized contact sites that allow the transmission of the nerve signal from one neuron to another

Connecting elements between nerve cells



A synapse is a site of close contact between a neuron and a target cell, such that an electrical signal in the neuron leads to a change in the probability that the target cell will produce an action potential.

-if the probability increases, the synapse is excitatory

-if the probability decreases, the synapse is inhibitory

Synaptic transmission is the closest example of intercellular communication

the target cell is distant <1  $\mu\text{m}$ 

the target cell is usually a few  $\mu\text{m}$  away

♦ Humoral transmission

the target cell can be anywhere in the body

### Neurons and targets



## Neural networks



### General structure

A synapse is made up of specialized structures both in the presynaptic termination and in the postsynaptic cell

### Electrical synapses vs. chemical synapses

### Electrical





The variation of the membrane potential of a cell is transmitted to another adjacent for direct current flow.

They are located in the CNS and peripheral of vertebrates and invertebrates where it is necessary a high speed of connection and synchronization in the activity of neighboring neurons



The variation of the membrane potential of a cell is transmitted to an adjacent cell by releasing a chemical transmitter triggered by the action potential of the presynaptic neuron.

The chemical transmitter acts on specialized proteins of the post-synaptic membrane, changing its permeability to ions

### Electrical synapses vs. chemical synapses





- prevent molecules from passing through the intercellular space.
- (b) Desmosomes: Anchoring junctions bind adjacent cells together like a molecular "Velcro" and help form an internal tensionreducing network of fibers.
- (c) Gap junctions: Communicating junctions allow ions and small molecules to pass for intercellular communication.

## Electrical synapses morphology

spazio

canali idrofili

connessoni

membrana

plasmatica

in int

the wave of depolarization of the action potential passes from one cell to another through a specialized structure, the gap junction

The electric synapses are joined by a plate-shaped gap, between which a space of less than 3 nm runs.

Connexon:

it is made up of 6 protein subunit arranged in hexagon that surround a channel permeable to water.

They are among the largest pores that are known. Their diameter is such as to allow the passage of the largest cell ions and organic micromolecules

# The electric synapses allow direct flow of ionic current from one cell to another



(a) Direct communication through gap junctions

© 2011 Pearson Education, Inc.

Nature Reviews | Neuroscience

Small molecules can easily pass between cells connected through gap-junction (connexons)

♦ Weight molecules up to ~ 2 kDa can pass from one cell to another electrically connected to it

♦ Fluorescent molecules such as yellow lucifer pass through gap-junctions – the connected cells are called "dye-coupled."



## Molecular organization and gap-junction topology



Goran Söhl et al Nature reveiw neurosci 2005

Three types of gap-junction homomeric/homotypic heteromeric heterotypic

Homotypic or heterotypic gap junctions comprise two identical or two different types of hemichannel, respectively.

Homomeric or heteromeric hemichannels are composed of one or more connexin (or possibly pannexin) isoforms, respectively.

Each hemichannel represents an assembly of six connexin protein subunits.

### Connessine



Connexin protein subunits are tetra-spanning membrane proteins that share three conserved extracellular cysteine residues, which are crucial for docking.

The subunits vary mainly in their cytoplasmic loop and carboxy-terminal region.

S-S represents conserved disulphide bonds in the extracellular domains of connexins.

# A cosa servono le gap junction quando coesistono con le sinapsi chimiche?



Improve network activity by synchronizing a large number of neuronal sets and their oscillatory activity (perception, memory and learning)

Due to the size of the pores (16-20 Å diameter) ions passe(above all K +) but also second messengers such as IP3 and cAMP

#### Table 1 | Properties of protein subunits in electrical synapses

Gap junction protein	Amino acid residues	Unitary conductance	Half-inactivation voltage (V <sub>0</sub> )	Ability to form heteromeric channels	Site of main expression
CX36	321	10–15 pS <sup>107</sup>	±75 mV <sup>107</sup>	n.d.	Interneurons <sup>33</sup> , olivary nucleus, cone photoreceptor cells <sup>81</sup>
CX45	396	32±8 pS <sup>108</sup>	±13.4 mV <sup>108</sup>	Yes (CX43/CX45) <sup>109</sup>	Distinct neuronal subtypes <sup>27</sup> , vascular smooth muscle cells <sup>106</sup> , conductive myocardiocytes <sup>106</sup>
CX57	492	n.d.*	n.d.	n.d.	Horizontal cells <sup>29</sup>
PX1	426	550 pS <sup>54</sup> (‡PX1)	n.d.	Yes⁵ (§PX1/§PX2)	<sup>II</sup> Principal neurons <sup>30</sup>
PX2	607	n.d.	n.d.	Yes (§PX1/§PX2)⁵	<sup>II</sup> Principal neurons <sup>30</sup>

\*The mouse CX57 protein with the correct carboxy-terminal region<sup>29</sup> has not yet been analysed. <sup>‡</sup>Human protein. <sup>§</sup>Mouse protein. <sup>§</sup>Expression based on *in situ* hybridization. CX, connexin; n.d., not determined; PX, pannexin.

In the Hippocampus the gamma oscillations depend on gap junctions



Electrical coupling between retinal cells

#### ◇During development

All cells in the initial stage of the embryo are coupled with gap junctions. This coupling, and its subsequent loss, can be related to the fate of the cells.

#### ◊In the epithelia

Many epithelia are made by sheets of paired cells. This coupling is regulated: damaged cells are decoupled from their neighbors.

#### ◊In the heart

the heart muscle cells are coupled in a syncithium, which allows conduction of the cardiac action potential in a coordinated manner.

#### ◇In the central nervous system

the coupling in the CNS seems to be important for the generation of coordinated behaviors of cell populations, and, in invertebrates for the rapid escape reaction.



# **Electrical SYNAPSE**



Action potential moves DIRECTLY between neurons

EXAMPLES:

Smooth Muscle Cardiac Muscle



Gap junction between adjacent cardiac cells





# Activation also subthreshold



### Stimulating current pulse

### The channels pass through 2 membranes





### Metabolic coupling due to gap junction: Schwann cells

### Charcot-Marie-Tooth Syndrome

- $\diamond$  Demyelinating disease of peripheral nerves  $\rightarrow$  sensory-motor neuropathy.
- $\diamond$  There is a form linked to the X chromosome.
- The X-linked form is associated with point mutations in a connexin gene expressed by Schwann cells (Cx32).
- $\Rightarrow$  alteration of metabolite flows from the outer (perinuclear) region of the Schwann cell to the internal myelin layers.



### Electrical coupling due to gap junction:



### Electrical coupling due to gap junction:





### Converting an Electrical Signal to Chemical Signal









# Chemical synapses



Copyright © 2006 Pearson Education, Inc., publishing as Benjamin Cummings.
### Chemical synapses



### The neuromuscular junction



Depending on which the postsynaptic part is invvolved



# CNS synapses – "Gray types"

Depending on the relationship between the pre-and post-synaptic termination thickness, the following is distinguished:

### The Geometry of Excitatory and Inhibitory Synapses



- Excitatory synapses
  - Gray's type I morphology
  - Spines: Excitatory synapses
- Inhibitory synapses
  - Gray's type II morphology
  - Clustered on soma and near axon hillock



Copyright © 2007 Wolters Kluwer Health | Lippincott Williams & Wilkins

### Why have chemical synapses?

Chemical synapses can modify the input signal:

- 1. Amplification
- 2. Reversal
- 3. Time extension
- 4. Plasticity phenomena

# Sequence of events (chemical synapses)

- 1. Arrival of the action potential in the axon terminal
- 2. Activation of activated voltage calcium channels
- 3. Ca2 + input into the terminal that stimulates vesicle fusion with the plasma membrane
- 4. Release of the neurotransmitter in the synaptic space
- 5. Linking the neurotransmitter to postsynaptic membrane receptors
- 6. Postsynaptic changes due to different membrane permeability



### Resting and activated synapses

RESTING STATE ACTIVATED STATE Α Myelin Neurotransmitter Presynaptic membrane sheath molecules are packaged depolarizes, usually as the into membranous result of an action potential. vesicles, which are concentrated and docked at the presynaptic Axon terminal. The depolarization causes Actin Extracellular voltage-dependent Ca24 filament space channels to open and allows Ca<sup>2+</sup> ions to flow into the terminal. Ca2+ channel Vesicles (closed) Ca<sup>2+</sup> Transmitter molecules The resulting increase of intracellular [Ca2+] Presynaptic triggers fusion of nerve terminal vesicles with the (bouton) presynaptic membrane. Synaptic. cleft 1 al

Receptor-G

protein complex

lon /

(closed)

Postsynaptic nerve terminal

of the nerve cell (spine)

Transmitter is released into the extracellular space in quantized amounts and diffuses passively across synaptic cleft.

Some of the transmitter molecules bind to receptors in the postsynaptic membrane, and the activated receptors trigger some postsynaptic event, usually the opening of an ion channel or the activation of a G-protein–coupled signal cascade.

6

-

Transmitter molecules

postsynaptic receptors

diffuse away from

and are eventually

continued diffusion,

or active uptake into

enzymatic degradation,

Transmitter-

degrading

enzymes

Extracellular

space

cleared away by

cells.

Action

potential

# Sequence of events (chemical synapses)



Nature Reviews | Neuroscience

### Resting and activated synapses: minis

The discovery of miniature endplate potentials; "minis"



Sir Bernard Katz



spontaneous

evoked



Ca++ is necessary for transmitter release

Low Ca++ in the perfusion medium



### Ca<sup>2+</sup> channels open in response to depolarization



Sinapsi gigante del calamaro per isolare le correnti al calcio

### Ca<sup>2+</sup> channels open in response to depolarization



A strong depolarization suppresses the calcium currents until the end of the depolarizing impulse



Augustine G, Charlton M, Smith S (1985) Calcium entry into voltage-calmped presynaptic terminals of squid. J Physiol 367:143–162.

# The peak of the calcium currents is in the descending phase of the action potential



© 2001 Sinauer Associates, Inc.

Lavoro basato sulla sinapsi gigante del calmato a varie T

### Voltage gated Ca<sup>2+</sup> channels



They transduce electrical signals into metabolic signals





Pore sodium permeability in the absence of calcium

 Voltage-dependent inactivation of conformational change at the entrance of the canal

◇Inactivation Ca<sup>2+</sup>-dependent



#### Which Ca<sup>2+</sup> channel is responsible for Transmitter release?



#### Agelenopsis Aperta

Conus Granulatus

conotoxins



#### Which Ca<sup>2+</sup> channel is responsible for Transmitter release?



ωconotoxin MVII A :Blocks N-type Ca2+ channels
ω Agatoxin IVa: Blocks P-type Ca2+ channels

### Ca2 + is necessary for the release of NT



Consequences of BAPTA injection into the presynaptic terminal: slowing of release

Adler EM, Augustine GJ, Duffy SN, Charlton MP (1991) Alien intracellular calcium chelators attenuate neurotransmitter release at the squid giant synapse. JNeurosci 11:1496–1507.

# Ca2 + is sufficient for the liberation of NT



Consequences of the arificial increase of [Ca2 +] i in the terminal: increase in the probability of release.

Delaney K, ZuckerRS (1990) Calcium released by photolysis of DM-nitrophenstimulates transmitter release at squid giant synapse. J Physiol 426:473–498.

What happens if we interfere with NT removal?

AChE blockers at the neuromuscular junction –sarin, GB –Malathion®

Blockers of the re-uptake of the amine transmitters

-dopamine -cocaine -serotonin -fluoxetine(Prozac \*)

A + R ↔AR + A ↔A2R ↔A2R\* ↔A2R

Quantal release : (a theory that explains the facts)

The presynaptic terminal releases quanta of ^ neurotransmitter which act on the postsynaptic membrane, depolarizing it







Del Castillo & Katz, 1957





### Release is vesicular

Synaptic Vesicle exocytosis captured by quick freezing; (Hauser & Reese)





BURGECENCE, They Ballion, Pigure 3.8 (Part 2) - 2004 Brauer Assesse



# The exo-endocytotic cycle of synaptic vesicles



### The exo-endocytotic cycle of synaptic vesicles



# The exo-endocytotic cycle of synaptic vesicles



Nature Reviews | Neuroscience

### Synaptic vesicle pools



Figure 1 | **Three vesicle pools. a** | The classic three-pool model. The reserve pool makes up ~80–90% of the total pool, and the recycling pool is significantly smaller (~10–15%). The readily releasable pool (RRP) consists of a few vesicles (~1%) that seem to be docked and primed for release. **b** | Three kinetic components of release (indicating release of three vesicle pools) on depolarization of goldfish bipolar cells. The cell was stimulated in the presence of the styryl dye FM 1-43, and the increase in fluorescence gives a direct measure of exocytosis. Panel **b** modified, with permission, from REF. 12 © (1999) Blackwell Scientific Publishing.

### Vesicular Cycling Kinetics



NEUROSCIENCE, Third Edition, Figure 5.9 (Part 2) @ 2004 Sinauer Associates, Inc.

### Presynaptic proteins involved in neurotransmitter release





Copyright © 2002, Elsevier Science (USA). All rights reserved.





(B) (1) La vescicola si ancora



(2) \$i formano i complessi SNARE per tirare le membrane



(3) Il Ca<sup>2+</sup>che estra silega alla sineptotagmina



(4) La sinoptotagmina che ha legato il Ca<sup>2+</sup> catalizza la fusione della nembrana



### Molecular mechanisms of NT liberation
# What would happen if...



You took a drug that destroyed the Ca<sup>2+</sup> sensitive proteins that fuse synaptic vesicles to the membrane???

## **That's How Botox Works!**



Botox destroys the proteins that fuse synaptic vesicles with the membrane.

By stopping vesicle release, Botox prevents muscle contraction which prevents wrinkles!



#### BOTOX IN THE NEUROMUSCULAR JUNCTION







## Toxins that interfere with the liberation of NT





# Classical and peptide neurotransmitters have different functions

#### CLASSIC NEUROTRASMECTORS:

- rapid and precise synaptic action
- self-sufficient terminal
- specific to SN

#### PEPTIDIC NEUROTRASMETTERS:

- slower and more modulatory paracrine action
- release requires intense electrical activity
- non self-sufficient terminal
- common to the endocrine system



#### Neurotransmitter action



(B) SMALL-MOLECULE TRANSMITTERS





#### Neurotrasmitter receptors





Copyright © 2002, Elsevier Science (USA). All rights reserved.

#### NMJ



## Synthesis and recycling of Acetylcholine



NMJ



#### Scheme



## Excitatory and inhibitory synapses



# Ionotropic vs. metabotropic PSPs



#### EPSP



# Excitatory and Inhibitory Synapses

Presynaptic action potentials result • in either Excitatory (EPSP) or Inhibitory Post Synaptic potentials (IPSP).

Changes in potential reflect ionic currents across the membrane.





The *nature* of the *neurotransmitter* determines the *response* of the post-synaptic membrane

# Inhibition

inhibitory neuron

-inhibitory synapse

-post-synaptic neuron

During hyperpolarisation, the post-synaptic membrane potential becomes more negative than its resting potential and results from either the efflux of positive charge or the influx of negative charge

Inhibition occurs at synapses where transmitter release results in the *hyperpolarisation* of the post-synaptic membrane

### Excitatory and inhibitory synapses coexist and modulate with each other



### Excitatory and inhibitory synapses coexist and modulate with each other



Nature Reviews | Neuroscience

# Presynaptic Inhibition and Facilitation

- Axoaxonic synapses: axon of one neuron synapses with the presynaptic terminal (axon) of another. Many of the synapses of CNS
- Presynaptic inhibition: reduction in amount of neurotransmitter released from presynaptic terminal. Endorphins can inhibit pain sensation
- Presynaptic facilitation: amount of neurotransmitter released from presynaptic terminal increases. Glutamate facilitating nitric oxide production



# **Presynaptic inhibition**

#### •Mechanism:

•GABA opens the CI<sup>-</sup> channels in the excitatory presynatpic terminal  $\rightarrow$ increase CI influx  $\rightarrow$  leading to decrease in the amplitude of the action potentials arriving at the knob  $\rightarrow$ decrease Ca<sup>2+</sup> influx  $\rightarrow$ decrease the amount of neurotransmitter released  $\rightarrow$ decrease in post-synaptic response ( $\downarrow$  EPSP)

