



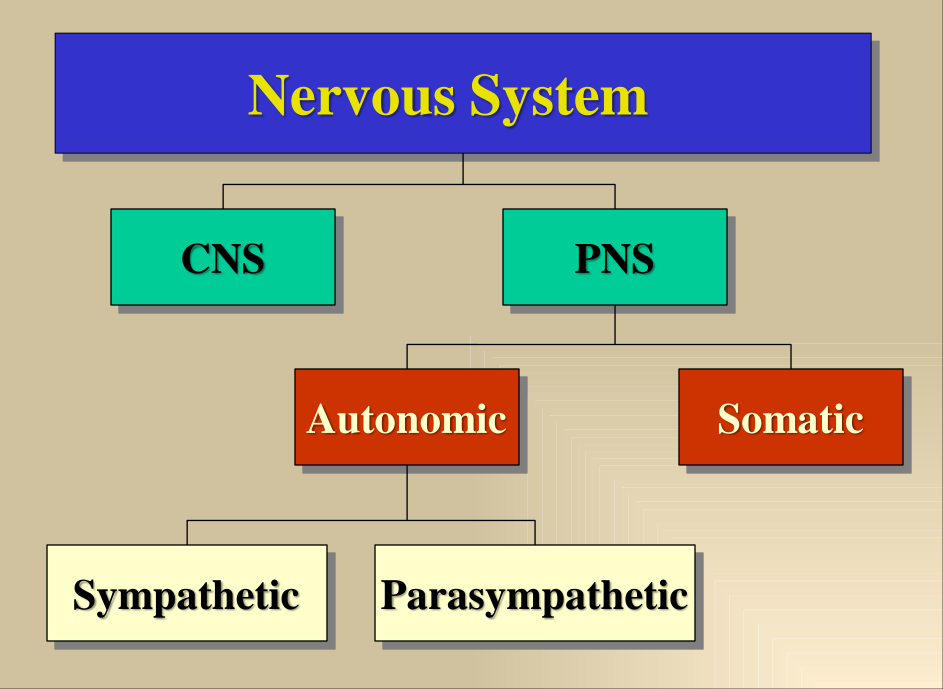
# Drugs That Affect The Nervous System

# **Topics**

- Analgesics and antagonists.
- Anesthetics.
- Anti-anxiety and sedative-hypnotics.
- Anti-seizure / anti-convulsants.
- CNS stimulators.
- Psychotherapeutics.
- ANS/PNS/SNS agents.

### **But first...**

# A colorful review of neurophysiology!



# Analgesics

#### Decrease in sensation of pain.

- Classes:
  - Opioid.
    - Agonist.
    - Antagonist.
    - Agonist-antagonist.
  - Non-opioids.
    - Salicylates.
    - NSAIDs.
    - Adjuncts.

# **Opioids**

- Generic reference to morphine-like drugs/actions.
  - Opiate: derivative of opium
- Prototype: morphine
  - Morpheus: god of dreams.
- Act on endorphin receptors:
  - Mu (most important).
  - Карра.



## **Actions of Opioid Receptors**

Response	Mu	Kappa
Analgesia		
Respiratory		
Depression		
Sedation		Ø
Euphoria		
Physical Dependence		
<b>↓ GI motility</b>		

# **Actions at Opioid Receptors**

Drugs	Mu	Kappa
Pure Agonists	Agonist	Agonist
-morphine, codeine, meperidine (Demerol <sup>®</sup> ), fentanyl (Sublimaze <sup>®</sup> ), remifentanil (Ultiva <sup>®</sup> ), propoxyphene (Darvon <sup>®</sup> ), hydrocodone (Vicodin <sup>®</sup> ), oxycodone (Percocet <sup>®</sup> )		
Agonist-Antagonist	Antagonist	Agonist
-nalbuphine (Nubaine®), butorphanol (Stadol®)		
Pure Antagonist	Antagonist	Antagonist
-naloxone (Narcan®)		

### **General Actions of Opioids**

- Analgesia.
- Respiratory depression.
- Constipation.
- Urinary retention.
- Cough suppression.
- Emesis.
- Increased ICP
  - Indirect through
    - CO<sub>2</sub> retention.

- Euphoria/Dysphoria.
- Sedation.
- Miosis
  - Pupil constriction.
  - **Preload & afterload** 
    - Watch for hypotension!

### **Non-opioid Analgesics**

- Salicylates:
  - Aspirin (Bayer®) \* (prototype for class).
- Non-Steroidal Anti-Inflammatory Drugs:
  - Ibuprofen (Motrin®, Advil®)
    - Propionic Acid derivative.
  - Naproxen (Naprosyn®).
  - Naproxen sodium (Aleve®).
  - All compete with aspirin for protein binding sites.
  - Ketorolac (Toradol®).

# **NSAID Properties**

Drug	Fever	Inflammation	Pain
Aspirin			
Ibuprofen			
Acetaminophen			

### **Aspirin Mechanism of Action**

Inhibit synthesis of cyclooxygenase (COX)
 – Enzyme responsible for synthesis of:

#### **Prostaglandins**

- -Pain response.
- -Suppression of gastric acid secretion.
- -Promote secretion of gastric mucus and bicarbonate.
- -Mediation of inflammatory response.
- -Production of fever.
- -Promote renal vasodilation (\blood flow).
- -Promote uterine contraction.

**Thromboxane A<sub>2</sub>** –Involved in platelet aggregation.

# **Aspirin Effects**

#### Good

Pain relief.
∀ ↓ Fever.
∀ ↓ Inflammation.

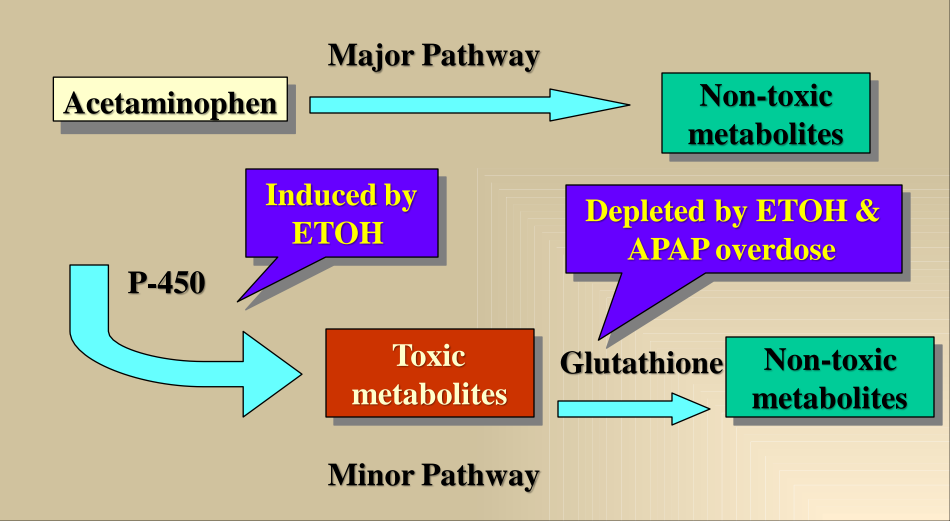
#### Bad

 GI ulceration: **f** Gastric acidity.  $\Downarrow$  GI protection.  $\forall \uparrow \mathbf{B}$  Bleeding.  $\forall \Downarrow \mathbf{Renal}$  elimination. ∀ **Uterine** contractions during labor.

# **Acetaminophen (Tylenol®)**

- NSAID similar to aspirin.
- Only inhibits synthesis of CNS prostaglandins.
  - Does <u>not</u> have peripheral side effects of ASA:
    - Gastric ulceration.
    - $\forall \Downarrow$  Platelet aggregation.
    - $\forall \Downarrow$  Renal flow.
    - $\forall \Downarrow$  Uterine contractions.

### Acetaminophen Metabolism



### Anesthetics

#### Loss of all sensation:

- Usually with loss of consciousness.
  - $\Downarrow$  propagation of neural impulses.
- General anesthetics:
  - Gases
    - Nitrous oxide (Nitronox®), halothane, ether.
  - -IV
    - Thiopental (Pentothal<sup>®</sup>), methohexital (Brevitol<sup>®</sup>), diazepam (valium<sup>®</sup>), remifentanil (Ultiva<sup>®</sup>).

### Anesthetics

#### • Local:

- Affect on area around injection.
- Usually accompanied by epinephrine
  - Lidocaine (Xylocaine ®), topical cocaine.

# Anti-anxiety & Sedativehypnotic Drugs

- Sedation: ↓ anxiety & inhibitions.
- Hypnosis: instigation of sleep.
- Insomnia:
  - ↑ Latent period.↑ Wakenings.
- Classes:
- Barbiturates.
- Benzodiazepines.
- Alcohol.

Chemically different, Functionally similar

### **Mechanism of action**

- Both promote the effectiveness of GABA receptors in the CNS:
  - Benzodiazepines promote only.
  - Barbiturates promote and (at high doses) stimulate GABA receptors.
- GABA = chief CNS inhibitory neurotransmitter:

Promotes hyperpolarization via Î Cl<sup>-</sup> influx.

# Benzodiazepines vs. Barbiturates

Criteria	BZ	Barb.
Relative Safety		Low
Maximal CNS depression		High
Respiratory Depression	Low	High
Suicide Potential	Low	High
Abuse Potential	Low	High
Antagonist Available?	Yes	No

### Benzodiazepines

#### **Benzodiazepines:**

- Diazepam (valium®).
- Midazolam (versed®).
- Alprazolam (xanax®).
- Lorazepam (atiavan<sup>®</sup>).
- Triazolam (Halcion<sup>®</sup>).

#### "Non-benzo benzo":

- Zolpidem (ambien®).
- Buspirone (BusPar®).

### **Barbiturates**

Subgroup	Prototype	Typical Indication
Ultra-short acting	thiopental (Pentothol®)	Anesthesia
Short acting	secobarbital (Seconal®)	Insomnia
Long acting	phenobarbital (Luminal®)	Seizures

### Barbiturates

- Amobarbital (amytal®).
- Pentobarbital (nembutal®).
- Thiopental (pentothal®).
- Phenobarbital (luminal <sup>®</sup>).
- Secobarbital (seconal <sup>®</sup>).

### **Anti-seizure Medications**

- Seizures caused by hyperactive brain areas.
- Multiple chemical classes of drugs:
  - All have same approach.
  - Decrease propagation of action potentials
     ∀↓ Na+, Ca++ influx (delay depolarization/prolong repolarization).
     ∀↑ Cl-influx (hyperpolarize membrane).

### **Anti-Seizure Medications**

#### <u>Benzodiazepines</u>

- Diazepam (valium®).
- •Lorazepam (Ativan®).

<u>Barbiturates</u>

 Phenobarbital (Luminal<sup>®</sup>).

#### Ion Channel Inhibitors

- Carbamazepine (Tegretol®).
- •Phenytoin (Dilantin®).

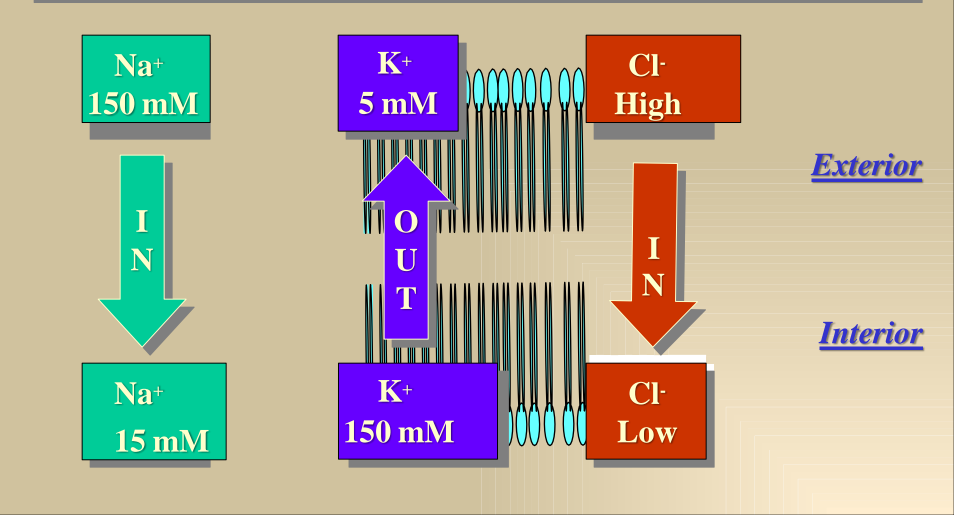
Misc. Agents

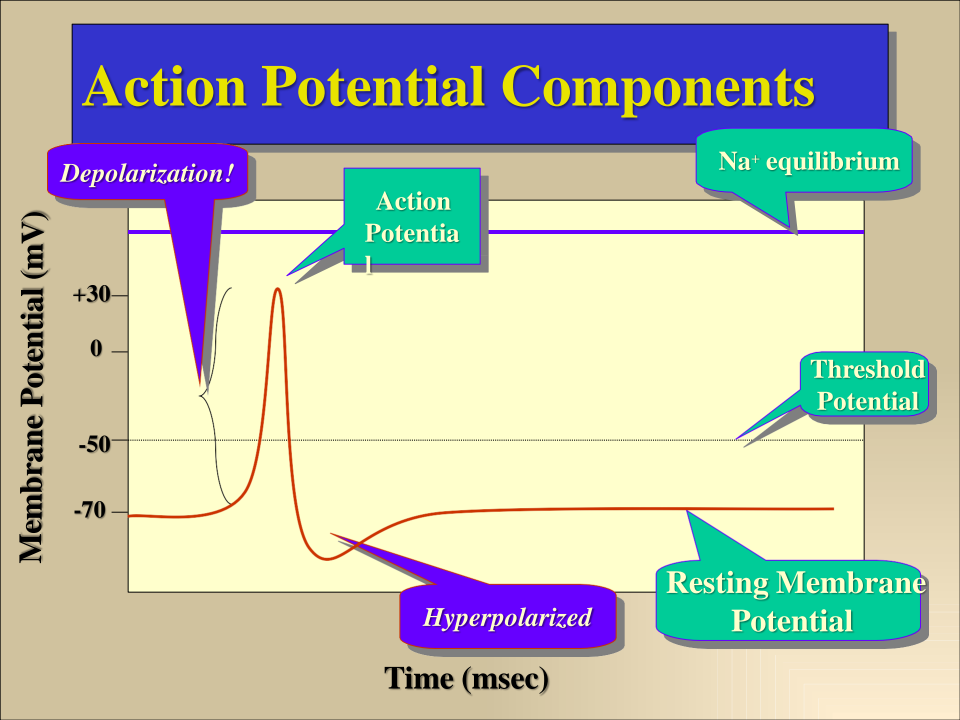
• Valproic acid (Depakote®).

### **Ion Diffusion**

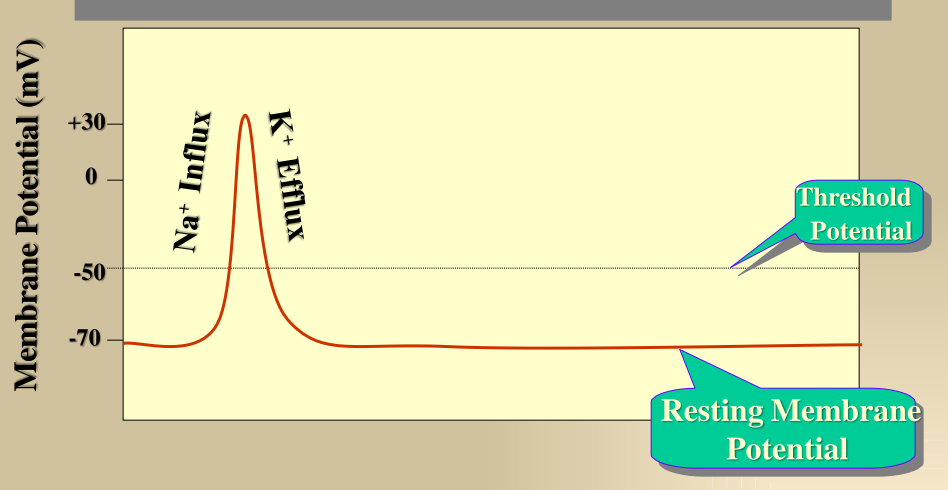
- Key to neurophysiology.
- Dependent upon:
  - Concentration gradient.
  - Electrical gradient.
- Modified by:
  - 'Gated ion channels'.

# Where Does Diffusion Take the Ion?



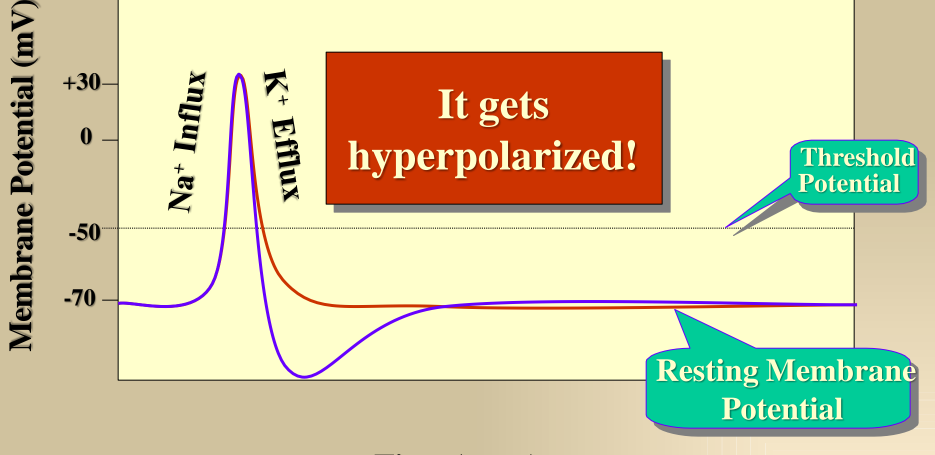


### **Membrane Permeability**



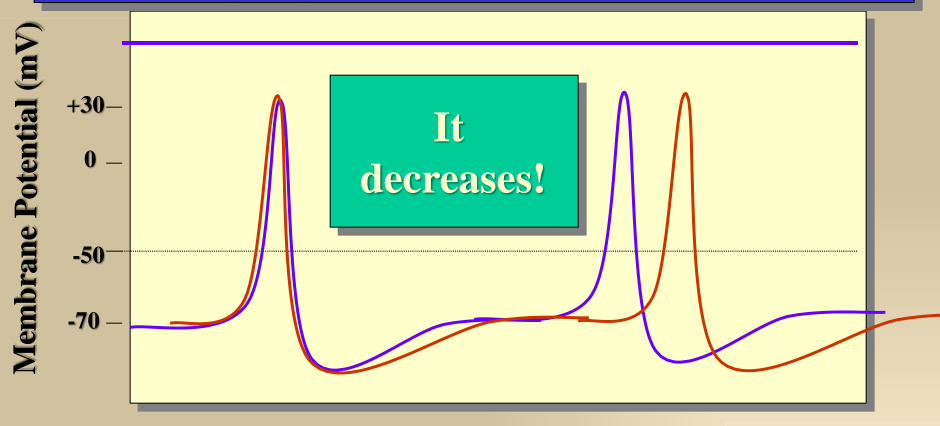
Time (msec)

### What Happens to the Membrane If Cl-Rushes Into the Cell During Repolarization?



Time (msec)

### What Happens to the Frequency of Action Potentials If the Membrane Gets Hyperpolarized?



Time (msec)

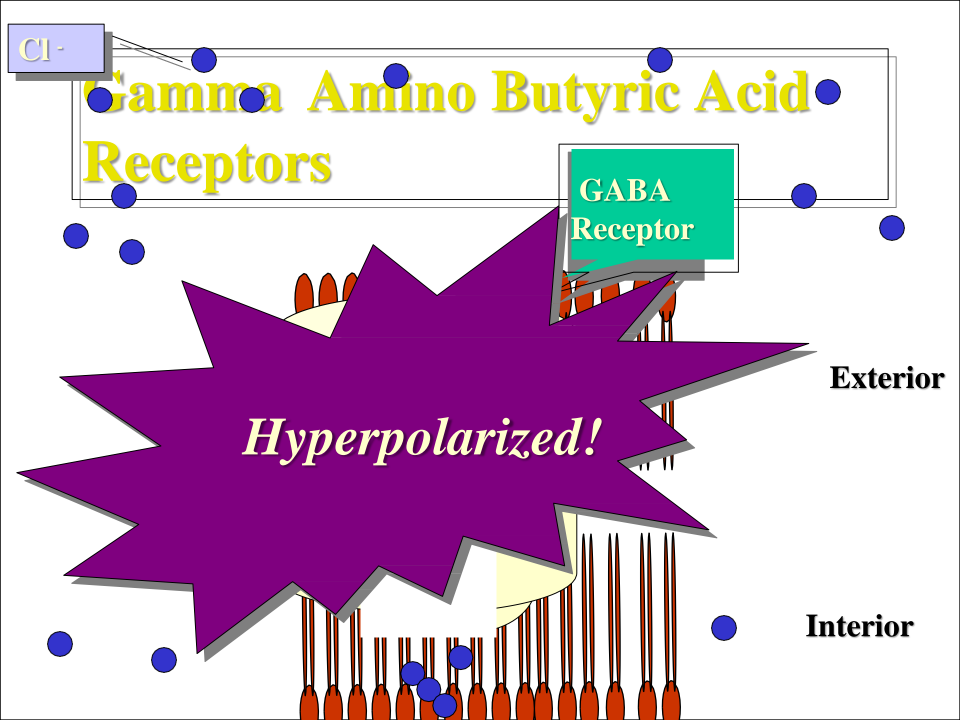
#### **Clinical Correlation**

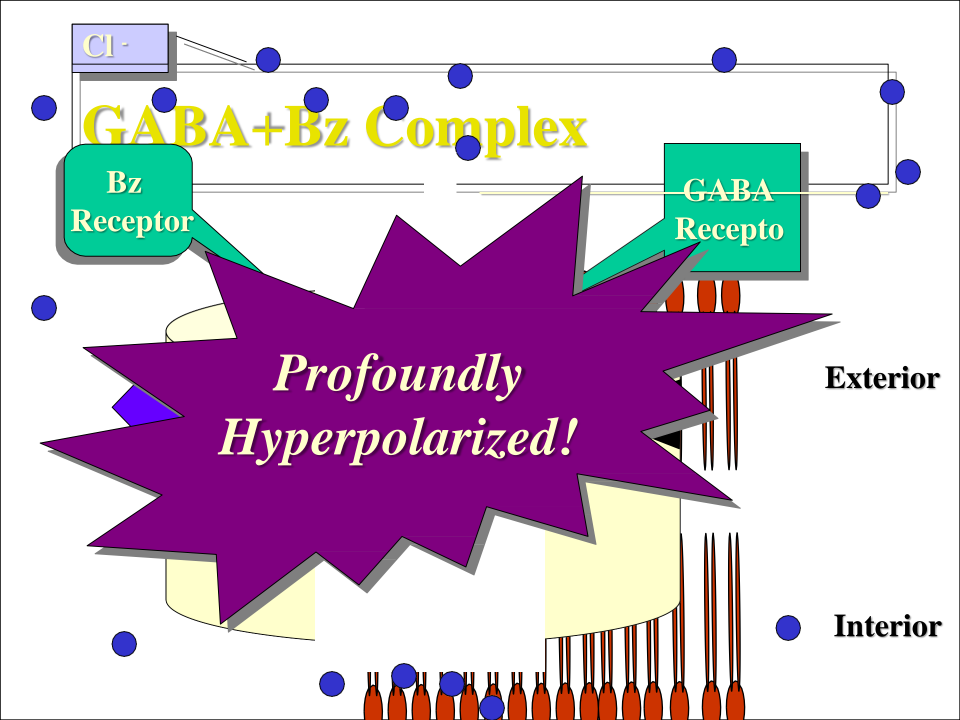
- Remember that it is the rate of action potential propagation that determines neurologic function.
  - Determined by frequency of action potentials.

What would be the effect on the membrane of ↑ Cl· influx during a seizure?

#### What is a seizure?

Hyperpolarization & ...





# Are You Ready for a Big Surprise?

Many CNS drugs act on GABA receptors to effect the frequency and duration of action potentials!

 $\bigcirc$ 

# **SNS Stimulants**

### Two general mechanisms:

- Increase excitatory neurotransmitter release.
- Decrease inhibitory neurotransmitter release.

#### • Three classes:

- Amphetamines.
- Methylphendidate.
- Methylxanthines.

# Amphetamines

Amphetamine. Methamphetamine. Dextroamphetamine. (Dexedrine®)

### **Indications**

Diet suppression.
∀↓ Fatigue.
∀↑ Concentration.

### **MOA:**

promote release of norepinephrine, dopamine.

### Side Effects

- •Tachycardia.
- Hypertension.Convulsion.
- •Insomnia.
- •Psychosis.

# Methylphenidate (Ritalin®)

- Different structure than other stimulants:
  - Similar mechanism.
  - Similar side effects.
- Indication: ADHD
  - Increase ability to focus & concentrate.

# Methylxanthines

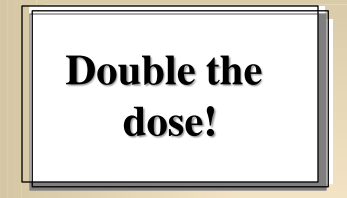
- Caffeine.
- Theophylline (Theo-Dur®).
- •Aminophylline.

**Mechanism of action** 

• Reversible blockade of adenosine receptors.

A patient is taking theophylline and becomes tachycardic (SVT). You want to give her adenosine. Is there an interaction you should be aware of? How should you alter your therapy?

Methylxanthines blocks adenosine receptors. A typical dose of adenosine may not be sufficient to achieve the desired result.



### News You Can Use...

Source	Amount of Caffeine
Coffee	
•Brewed	40 – 180 mg/cup
•Instant	30 – 120 mg/cup
<b>Decaffeinated Coffee</b>	2 - 5 mg/cup
Tea	20 – 110 mg/cup
Coke	40 – 60 mg/12 oz

# **Psychotherapeutic Medications**

- Dysfunction related to neurotransmitter imbalance.
  - Norepinephrine.
  - Dopamine.
  - Seratonin.

Monoamines

 Goal is to regulate excitory/inhibitory neurotransmitters.

# **Anti-Psychotic Drugs** (Neuroleptics)

- Schizophrenia:
  - Loss of contact with reality & disorganized thoughts.
  - Probable cause: increased dopamine release.
  - Tx. Aimed at decreasing dopamine activity.



#### Phenothiazines

- chlorpromazine (Thorazine ®)
- **Butyrophenones** 
  - haloperidol (Haldol®)

# **Other Uses for Antipsychotics**

- Bipolar depression.
- Tourette's Syndrome.
- Prevention of emesis.
- Dementia (OBS).
- Temporary psychoses from other illness.

# **Antipsychotic MOA**

- Mechanism is similar.
- Strength ([]) vs. Potency ('oomph'):
  - Phenothiazines low potency.
  - Butyrophenones high potency.
- Receptor Antagonism:
  - Dopamine<sub>2</sub> in brain
    - Muscarinic cholinergic <sup>–</sup>
    - Histamine
    - -Norepi at alpha<sub>1</sub>

Therapeutic effects

**Uninteded** effects

# **Antipsychotic Side Effects**

- Generally short term.
- Extrapyramidal symptoms (EPS).
- Anticholinergic effects (atropine-like):
  - Dry mouth, blurred vision, photophobia, tachycardia, constipation).
- Orthostatic hypotension.
- Sedation.
- Decreased seizure threshold.
- Sexual dysfunction.

# **Extrapyramidal Symptoms**

Reaction	Onset	Features
Acute dystonia	Hours to 5 days	Spasm of tongue, neck, face & back
Parkinsonism	5 – 30 days	Tremor, shuffling gait, drooling, stooped posture, instability
Akathesia	5 – 60 days	Compulsive, repetitive motions; agitation
Tarditive dyskinesia	Months to years	Lip-smacking, worm-like tongue movement, 'fly- catching'

## **Treatment of EPS**

- Likely caused by blocking central dopamine<sub>2</sub> receptors responsible for Movement.
- Anticholinergic therapy rapidly effective – diphenhydramine (Benadryl<sup>®</sup>).

# **Antipsychotic Agents**

- Chlorpromazine (thorazine®).
- Thioridazine (mellaril®).
- Trifluoperazine (stelazine®).
- Haloperidol (Haldol®).

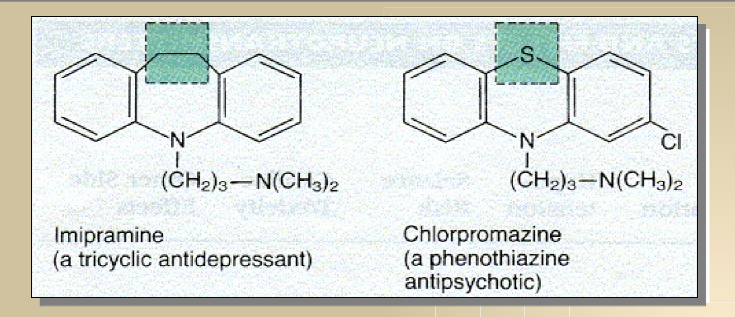
## Antidepressants

- Likely cause: Inadequate monoamine levels.
- Treatment options:
  - Increasing NT synthesis in presynaptic end bulb.
  - Increasing NT release from end bulb.
  - Blocking NT 'reuptake' by presynaptic end bulb.

# **Tricyclic Antidepressants** (TCAs)

- Block reuptake of both NE & serotonin

  Enhance effects.
- Similar side effects to phenothiazines.



## **TCA Side Effects**

- Orthostatic hypotension.
- Sedation.
- Anticholinergic effects.
- Cardiac toxicity
  - Ventricular dysrythmias.

# **Selective Serotonin Reuptake Inhibitors (SSRIs)**

- Block only serotonin (not NE) reuptake – Elevate serotonin levels.
- Fewer side effects than TCS
  - No hypotension.
  - No anticholinergic effects.
  - No cardiotoxicity.
- Most common side effect

- Nausea, insomnia, sexual dysfunction.

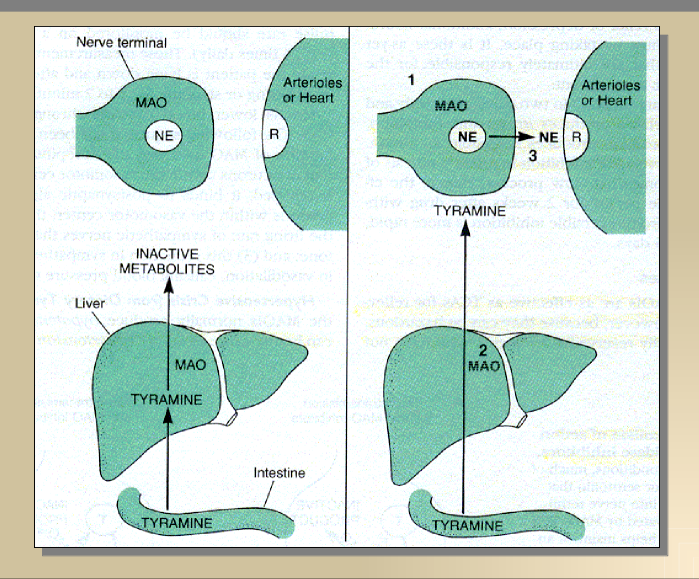
# Monoamine Oxidase Inhibitors (MAOIs)

- Monoamine oxidase
  - Present in liver, intestines & MA releasing neurons.
  - Inactivates monoamines.
  - Inactivates dietary tyramine in liver
    - Foods rich in tyramine: cheese & red wine.

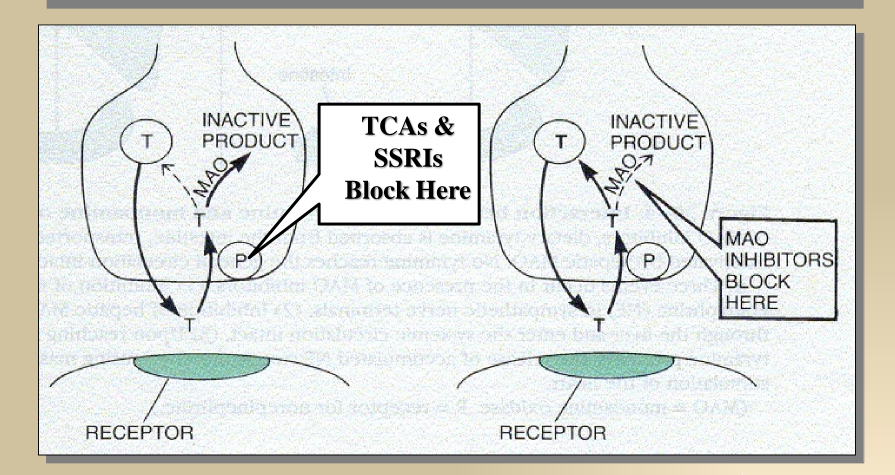
# **MAOI Side Effects**

- CNS Stimulation
  - Anxiety, agitation.
- Orthostatic hypotension.
- Hypertensive Crisis
  - From increased tyramine consumption
    - Excessive arteriole constriction, stimulation of heart.

# **MAOI & Dietary Tyramine**



## **Antidepressant Mechanism**



## **Antidepressants Agents**

### **TCAs**

- Imiprimine (tofranil®)
- Amitriptyline (elavil<sup>®</sup>)
- Nortriptyline (pamelor <sup>®</sup>)
   <u>SSRIs</u>
- Fluoxetine (prozac<sup>®</sup>)
- Paroxetine (paxil<sup>®</sup>)
- Sertraline (zoloft<sup>®</sup>)

#### MAOIs

• Phenelzine (Nardil®)

#### <u>Atypical</u> <u>Antidepressants</u>

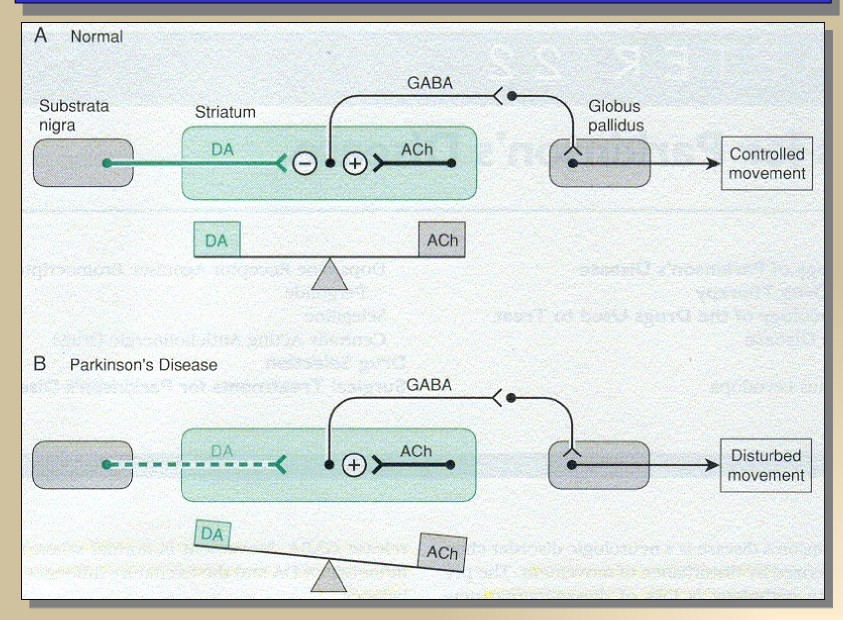
Bupropion (Wellbutrin®)

## **Parkinson's Disease**

- Fine motor control dependent upon balance between excitatory and inhibitory NT.
- Acetylcholine = excitatory
- –Dopamine =inhibitory
- **GABA**= inhibitory

Control - GABA release

# **Parkinson's Disease**



# **Parkinson's Symptoms**

- Similar to EPS.
- Dyskinesias

Tremors, unsteady gait, instability.

- Bradykinesia.
- Akinesia in severe cases.

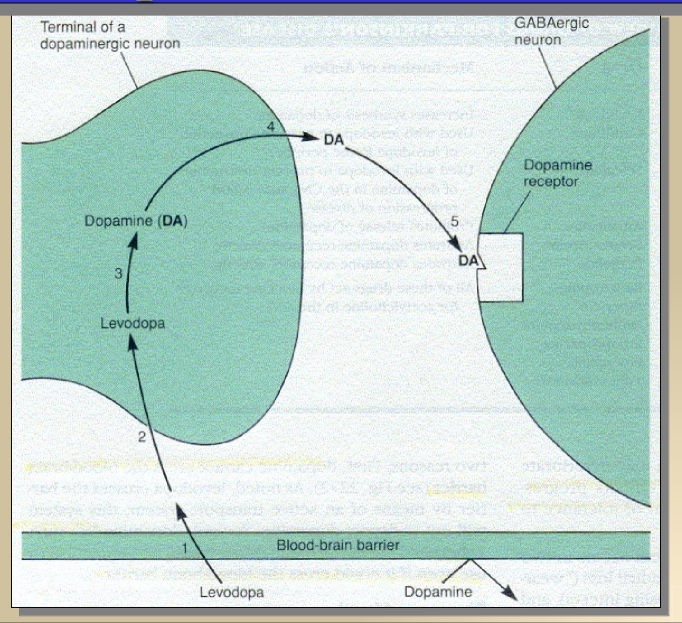
### **Parkinson's Treatment**

**Dopaminergic approach: 1** Release of dopamine. **[]** [Dopamine]. **↓** Dopamine breakdown. **Cholinergic approach: ↓** Amount of ACh released. - Directly block ACh receptors. All treatment is symptomatic and temporary.

## Levodopa

- Sinemet <sup>®</sup> = levodopa + carbidopa.
- Increase central dopamine levels.
- Side effects:
  - Nausea and vomiting.
  - Dyskinesia (~80% of population).
  - Cardiovascular (dysrythmias).

# Levodopa Mechanism



# **Other Agents**

- Bromocriptine (Parlodel<sup>®</sup>)
  - Directly stimulated dopamine receptors.
- Selegiline (Carbex<sup>®</sup>, Eldepryl<sup>®</sup>)
   MAOI selective for dopamine (MAO-B).
- Benztropine (Cogentin®)

Centrally acting anticholinergic.

