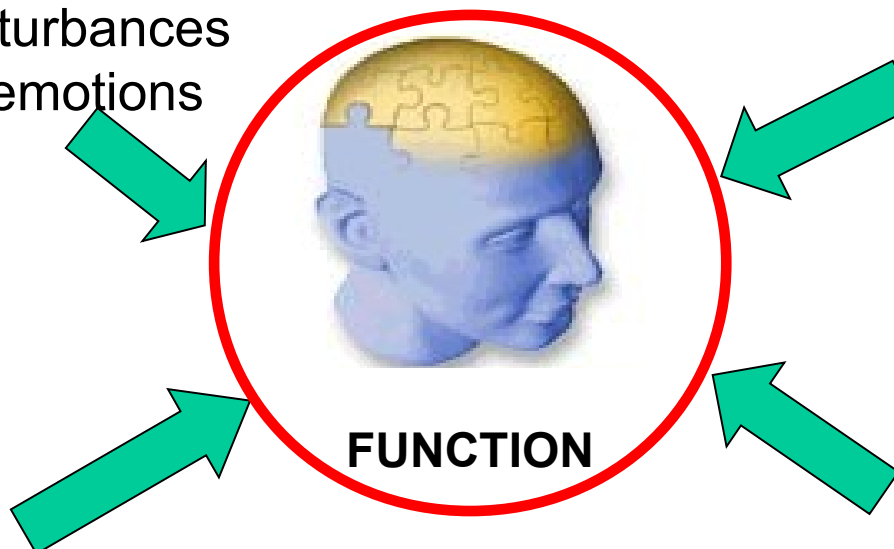


Neuroleptics = antipsychotics = „major tranquilizers“: drugs for treatment, or better for attenuation of symptoms of schizophrenic psychoses

# Schizophrenia - symptoms

## Positive Symptoms

Hallucinations  
Delusions (bizarre, persecutory)  
Disorganized Thought  
Perception disturbances  
Inappropriate emotions



## Cognition

New Learning  
Memory

## Negative Symptoms

Blunted emotions  
Anhedonia  
Lack of feeling

## Mood Symptoms

Loss of motivation  
Social withdrawal  
Insight  
Demoralization  
Suicide

# Historic and alternative treatment of schizophrenia

- insulin coma
- electroconvulsions
- prefrontal lobotomy
  - Egas Moniz, 50 000 surgeries, 1935 Nobel prize
  - patients were just calmer, but also more sluggish and apathetic

## ► Prefrontal Lobotomy Procedure of Moniz and Lima

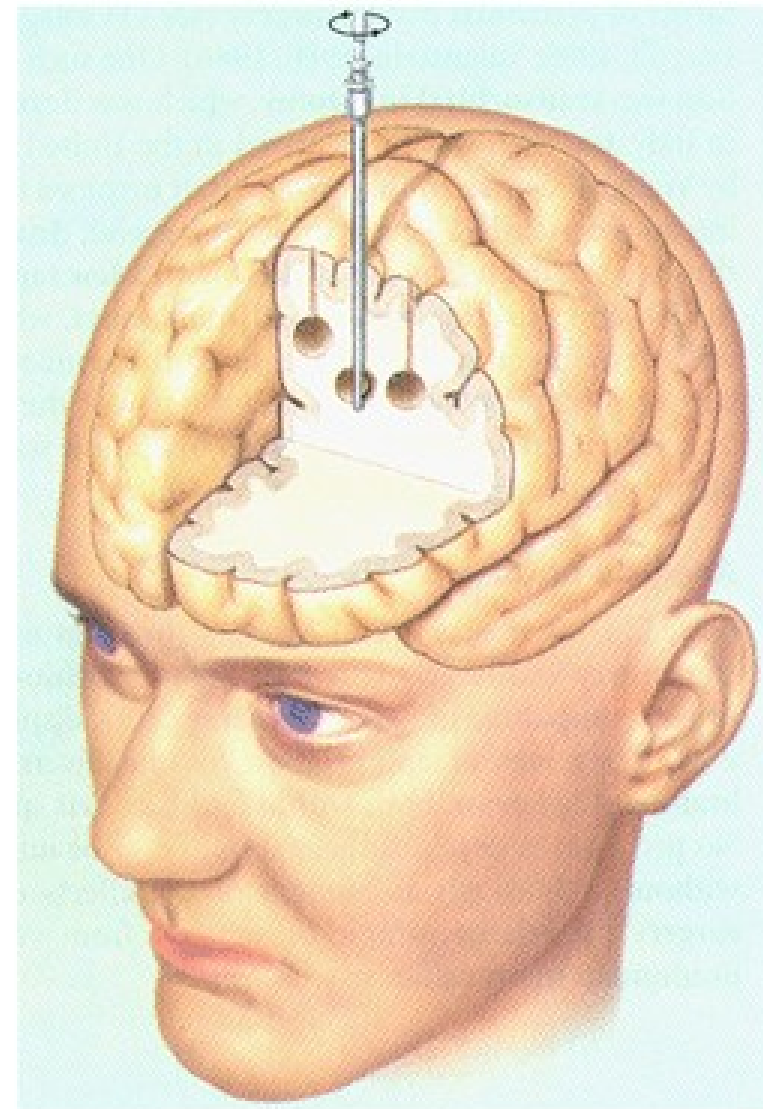
### The Prefrontal Lobotomy Procedure of Moniz and Lima



The leucotome was inserted 6 times into the patient's brain with the cutting wire retracted.



After each insertion the cutting wire was extruded and the leucotome rotated to cut out a core of tissue.



# Schizophrenia Pathophysiology

## Schizophrenia Pathophysiology

**Past** Excess dopaminergic activity

Dopamine D<sub>2</sub>-receptor antagonists

## Present

Renewed interest in the role of serotonin (5-HT)

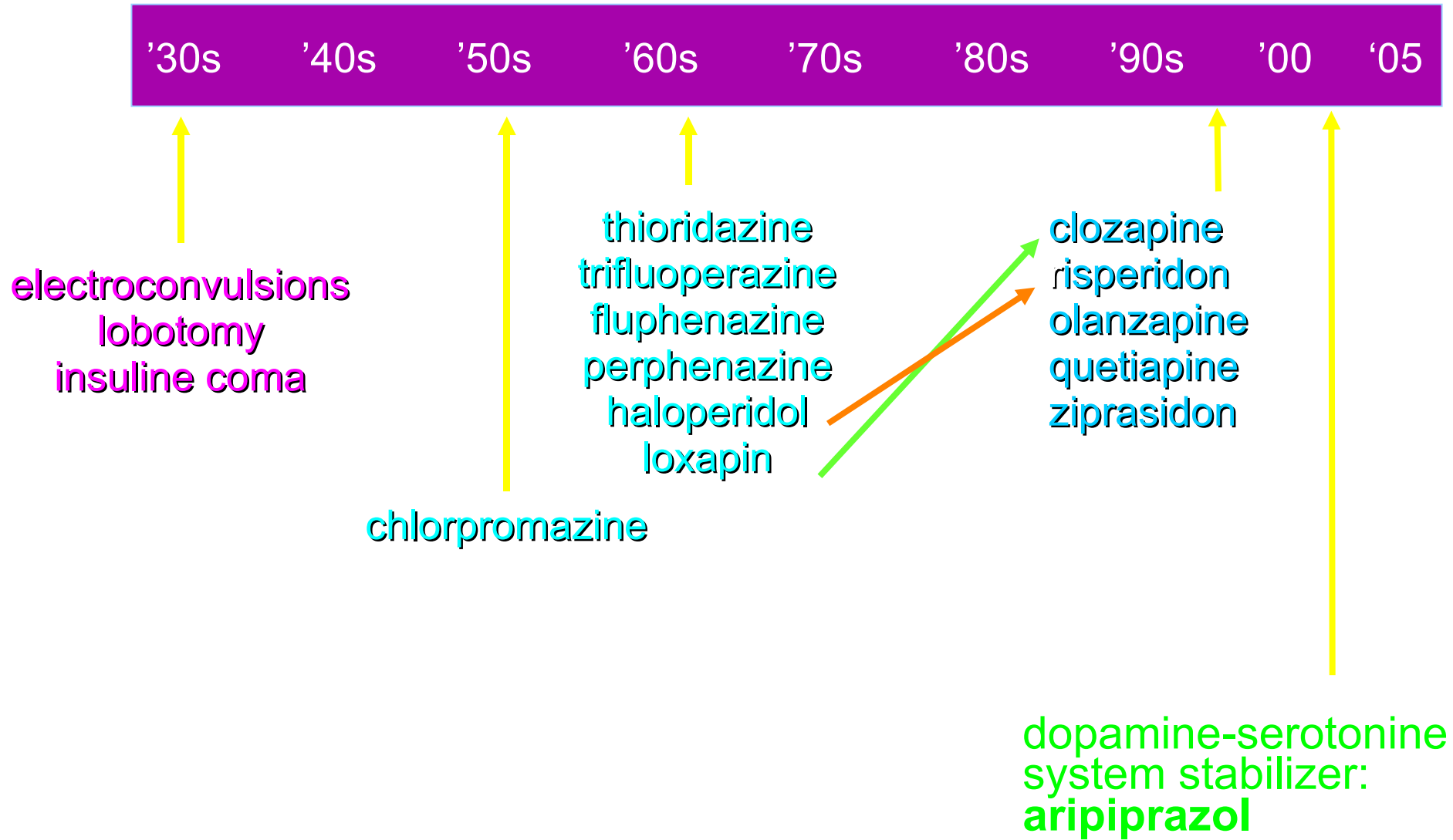
Combined 5-HT<sub>2</sub>/D<sub>2</sub> antagonists

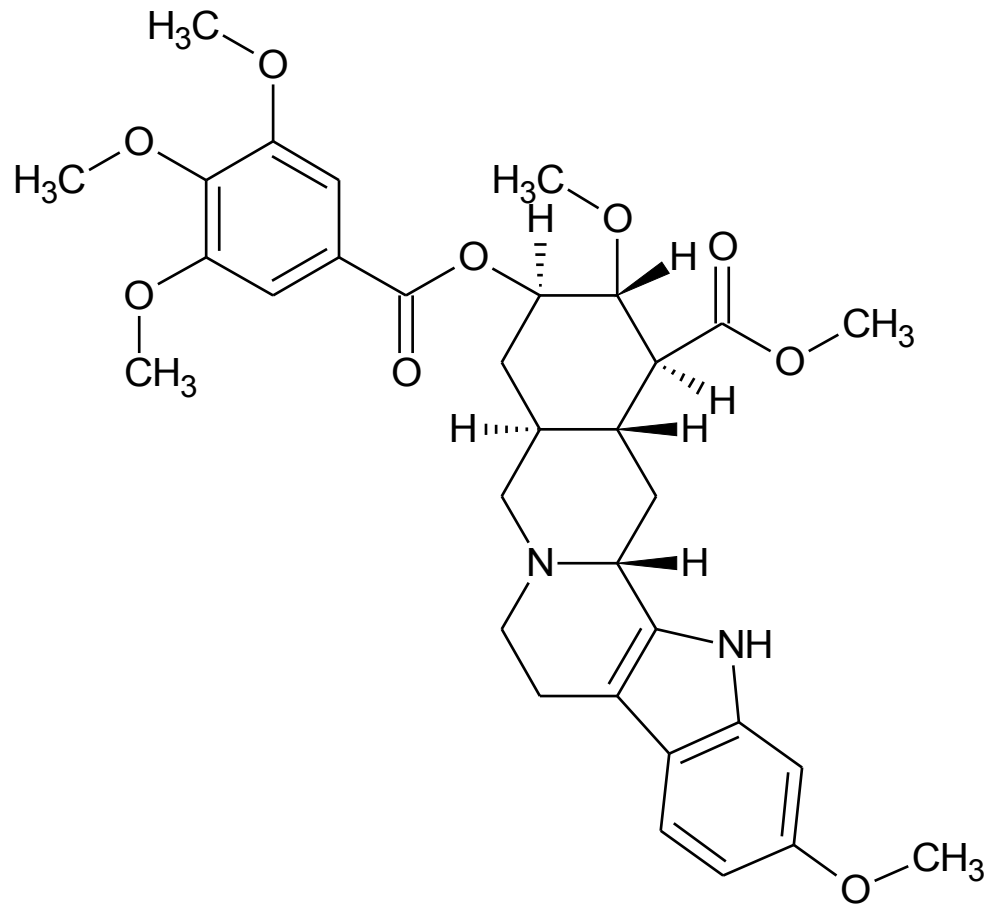
## Future

Imbalance in cortical communication and cortical-midbrain integration, involving multiple neurotransmitters

More selective antagonists  
Mixed agonist/antagonists  
Neuropeptide analogs

# Evolution of therapy of schizophrenic psychoses





**reserpine**

- *Rauvolfia serpentina*
- inhibition of noradrenaline uptake into storing vesicles ⇒ decrease of catecholamines levels in both central and peripheral neuronal ends
- antipsychotic
- antihypertensive
- high toxicity

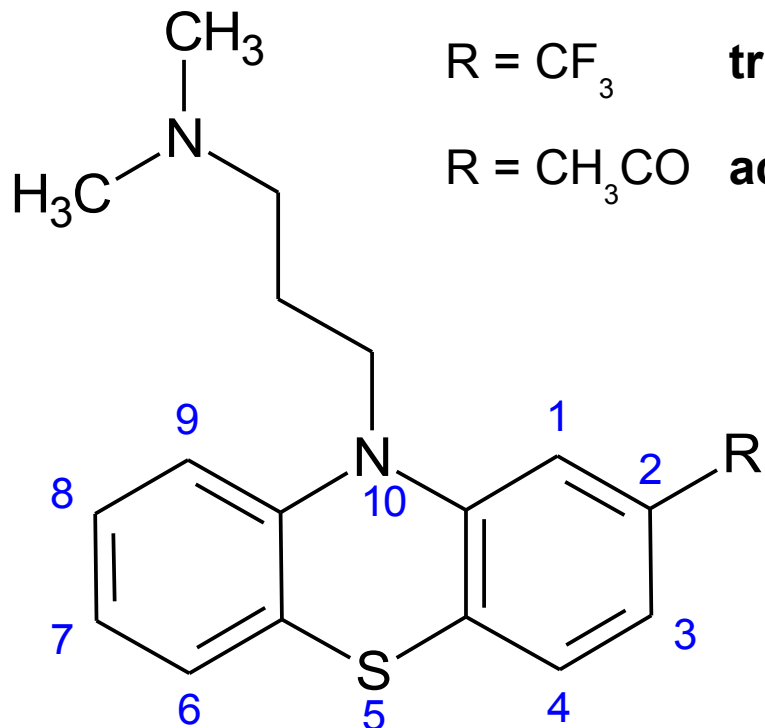
## „Typical“ antipsychotics

Phenothiazines with unbranched aminopropane side chain

R = H      **promazine**

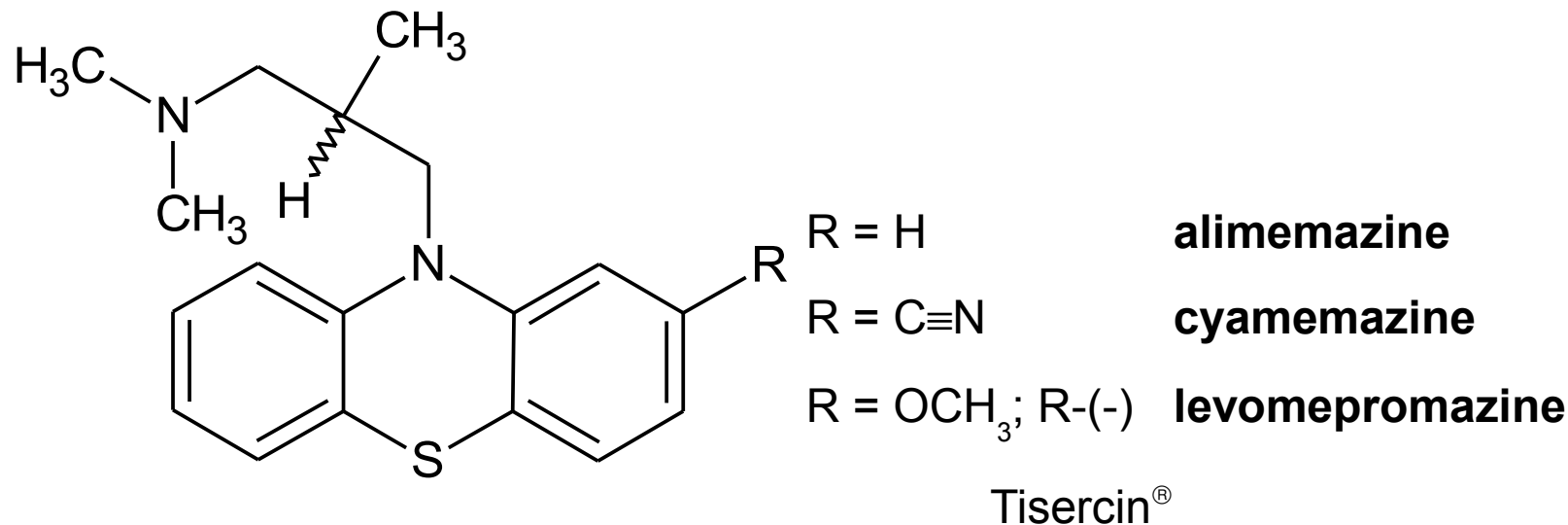
R = Cl      **chlorpromazine** Plegomazin®

- Henri Labroit, French military surgeon: causes „artifitial hibernation“
- in therapy since 1953

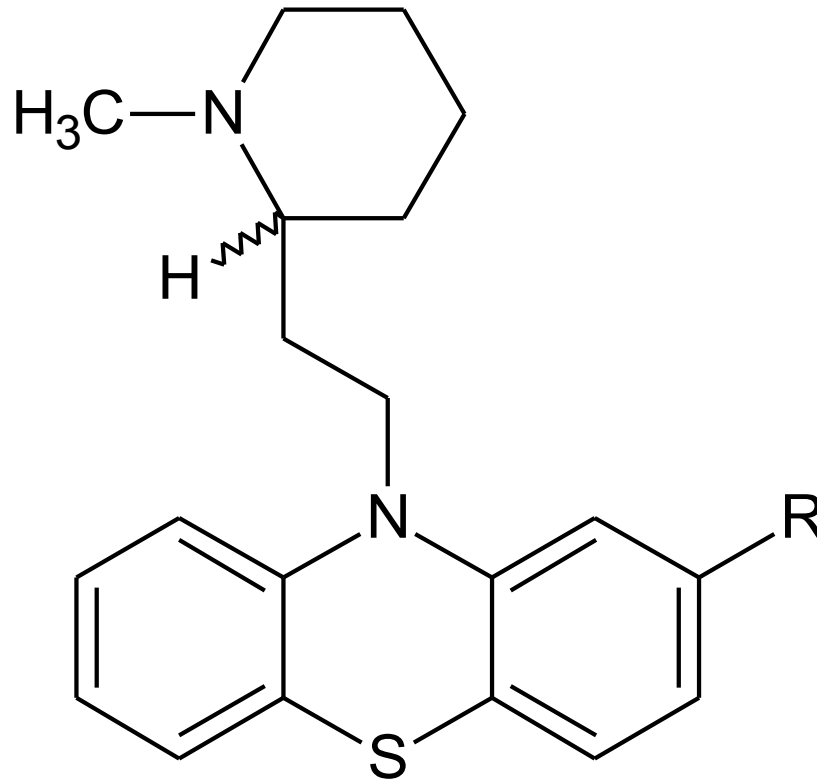


H. Labroit

## Phenothiazines with branched aminoalkane side chain



## Phenothiazines with 2-(piperidine-2-yl)ethyl side chain

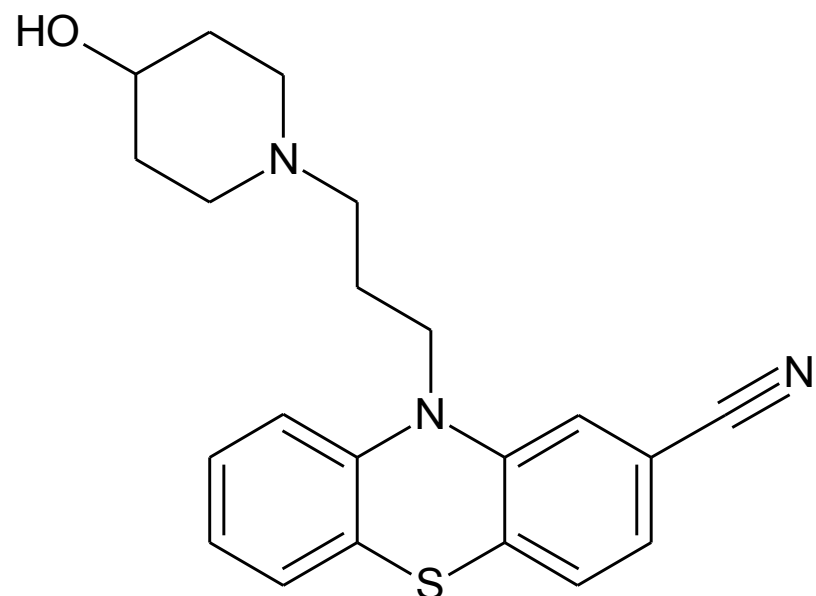


$\text{R} = \text{CH}_3\text{S}$  **thioridazine**

- also antimicrobial activity: *Mycobacterium tuberculosis*, *Listeria monocytogenes*
- in some developing countries used as an antituberculotic

$\text{R} = \text{CH}_3\text{SO}$  **mesoridazine**

## Phenothiazines with 3-(piperidine-1-yl)propyl side chain

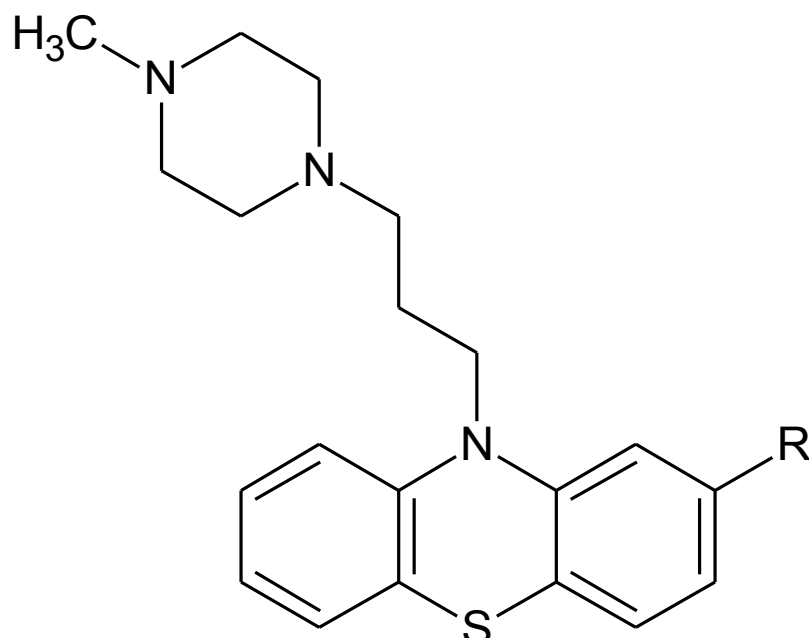


**periciazine**

syn. propericiazine

- AE: hypersensitivity of sensual perception

Perazine series: phenothiazines with 3-(piperazin-3-yl)propyl side chain

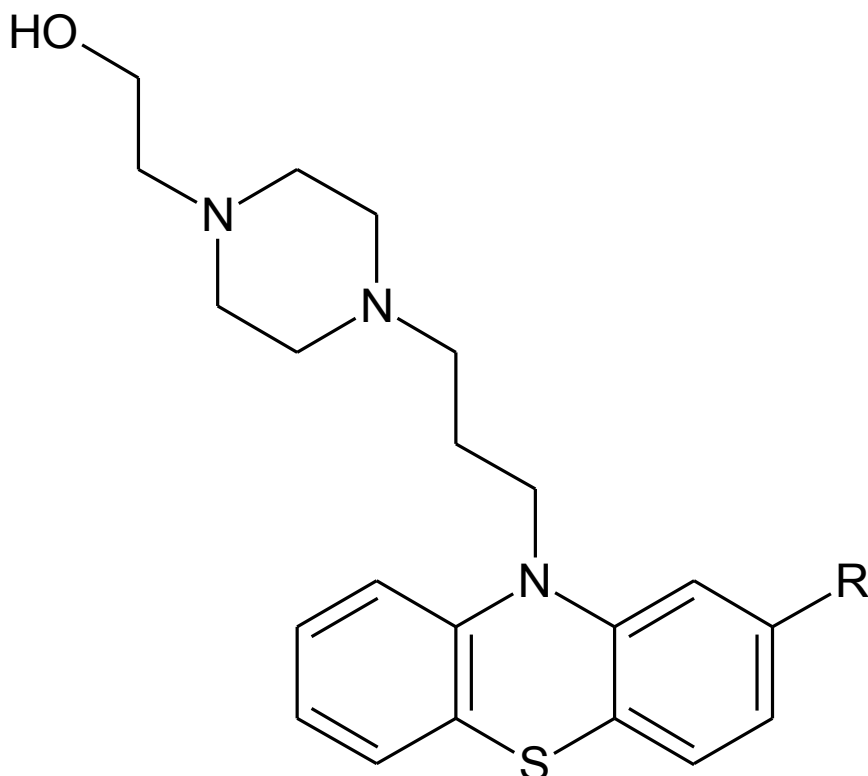


R = H

**perazine**

R = CF<sub>3</sub>

**trifluperazine**



R = Cl

**perphenazine**

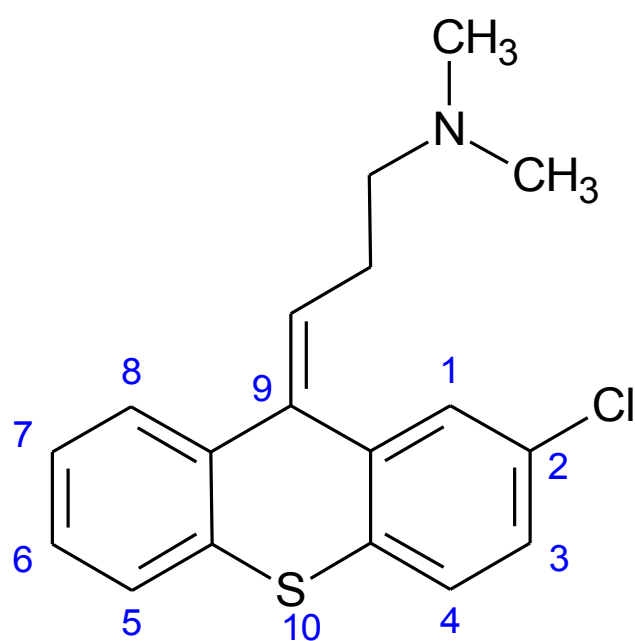
R = CF<sub>3</sub>

**fluphenazine**

Moditen

Depot® inj. sol.

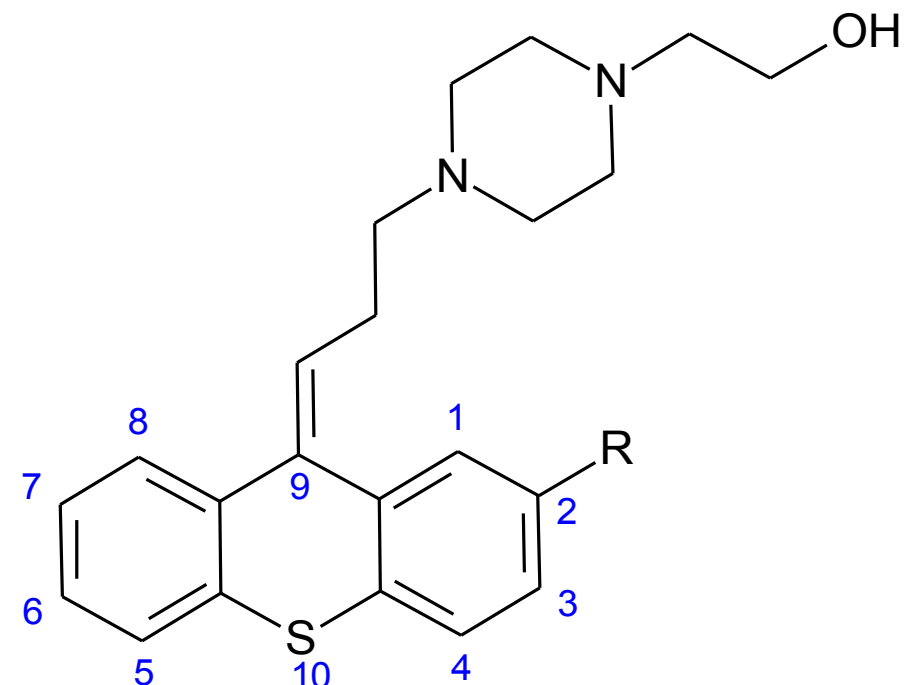
## Thioxanthenes: isosteric analogues of phenothiazines



**chlorprothixene**

•Z-isomer

Chlorprothixen Léčiva®



R = Cl

•mixture E/Z: **clopenthixol**

•Z-isomer: **zuclopentixol**

Cisordinol®

R = CF<sub>3</sub>      **flupenthixol** Fluanxol®

•mixture E/Z

## Structure-activity relationships (SAR) of phenothiazines and thioxanthenes

### 1. linking chain between N(10) and the basic substituent:

- propyl is optimal; compounds with butyl nearly inactive, ethyl  $\Rightarrow$  antihistamine activity
- any substituent in pos. 1 of the side alkyl lowers the activity
- methyl or phenyl in pos. 2 do not decrease the activity while more bulky aliphatic substituents do
- many various substitutions can be proceeded in pos. 3; basic N is often a part of a ring

### 2. substituent in pos. 2 of the tricyclic ring

- the highest effect is linked with electron-accepting lipophilic substituents (-Cl, -CF<sub>3</sub>, -CN), activity increases with lipophilicity and electron-accepting properties, electrondonor substituents (-OCH<sub>3</sub>, -SCH<sub>3</sub>) lower activity

### 3. tricyclic ring

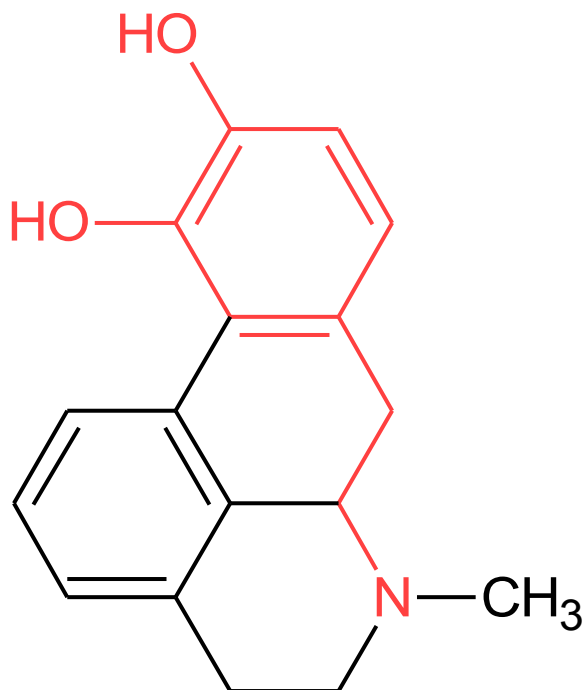
- disubstitution lowers activity, ring opening completely removes it
- substitution of S with C, O, Se etc. lowers activity; substitution of N(10) lowers activity except that with alkylidene substituted C ( $\Rightarrow$  thioxanthenes)
- isosteric substitution C(2) with N keeps activity ( $\Rightarrow$  2-azafenothiazines)
- in thioxanthenes, compounds with Z-configuration on double bond going out from C(9) have higher activity than E-isomers

### 4. modification of amino group of side chain

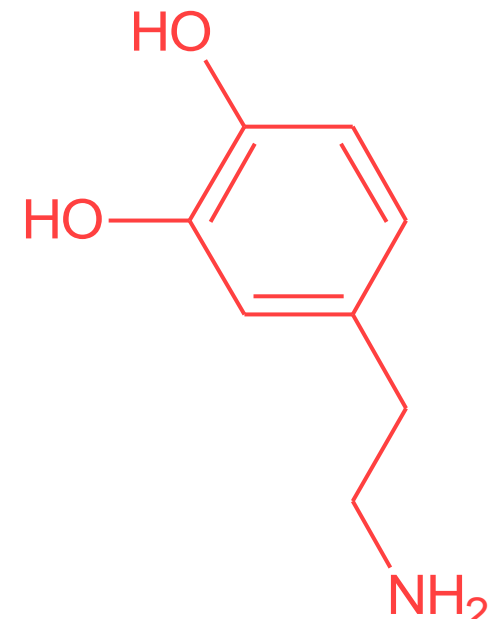
- tertiary amines ( $pK_a$  8-10) have maximum activity
- methyls on nitrogen lead to higher activity than longer alkyls; receptor is long and narrow which is shown by tolerance of phenyl in pos. 2 of the chain
- amino group can be part of a ring; pyrrolidine, piperidine and morpholine belong to useful cyclic substituents; compounds with piperazine are the most active ones

## Mechanism of action of tricyclic antipsychotics

- reversible block of D<sub>2</sub>-subtype of dopamine receptor
- evidence of relationship between antipsychotic antagonism against dopamine agonist apomorphine (displacement of apomorphine from this receptor) and dopamine accumulation in brain



apomorphine

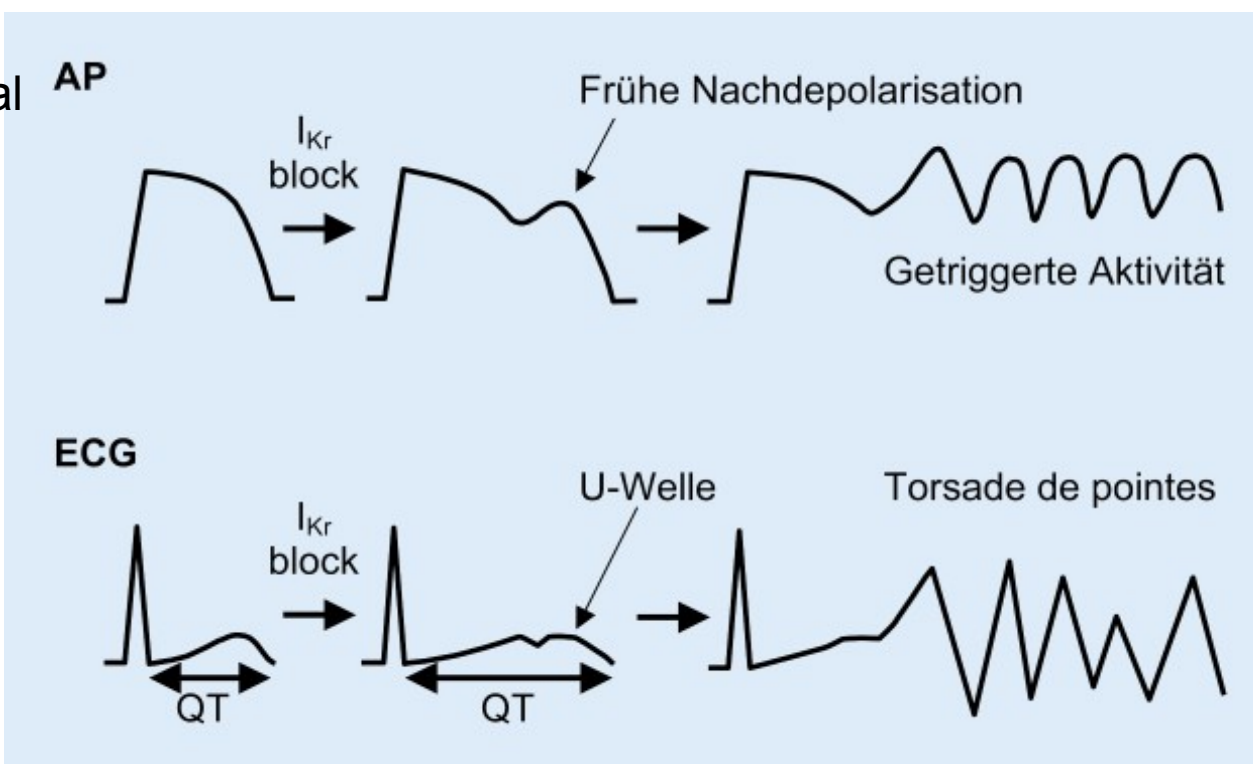


dopamine

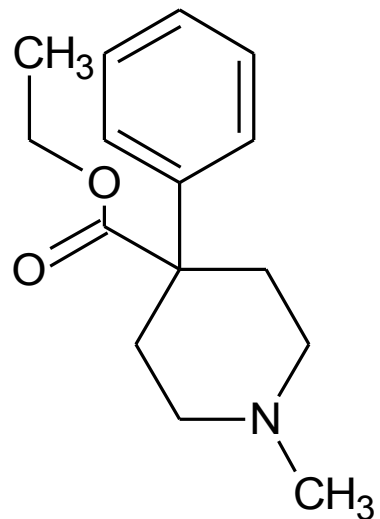
## Unwanted effects of phenothiazines and thioxanthenes

- Parkinsonian = extrapyramidal syndrome – caused by relative excess of acetylcholine in CNS over dopamine
- cardiovascular system: dysrhythmias of type of Torsade de pointes (TdP; „bundle of spikes“) - begins with QT-interval elongation on ECG due to  $K^+$ -channels block – can lead to cardiac arrest and sudden death (mostly thioridazine)
- „amplified“ vision (lights and colours more intensive, objects bigger)

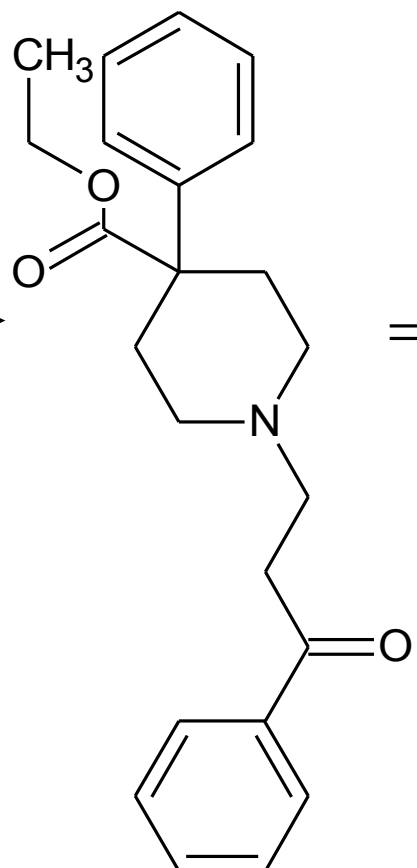
AP = action potential  
 $I_{Kr}$  = stream through  
 $K^+$  channel



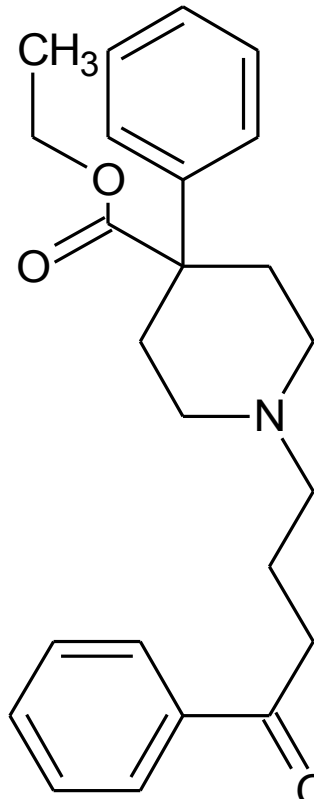
„Typical“ neuroleptics: butyrophenones a diphenylbutylpiperidines  
Origin of butyrophenones



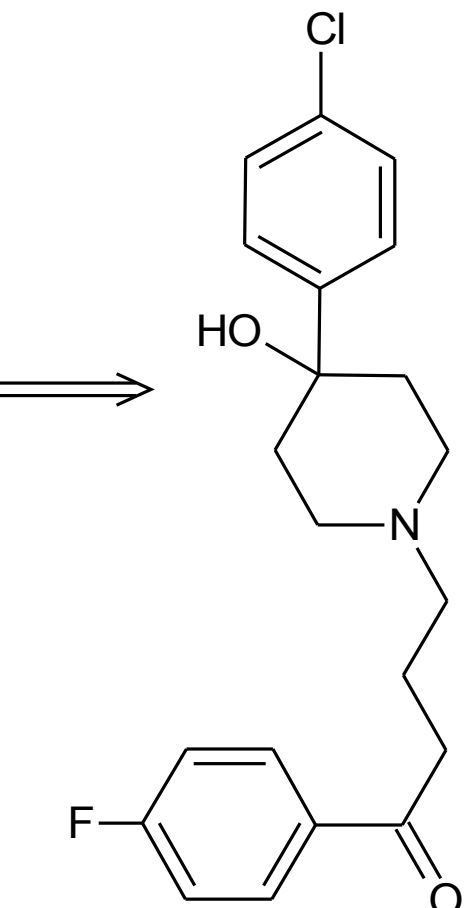
pethidine  
opioid analgesic



propiophenone  
analogue of pethidine  
• 200x highest  
analgesic activity

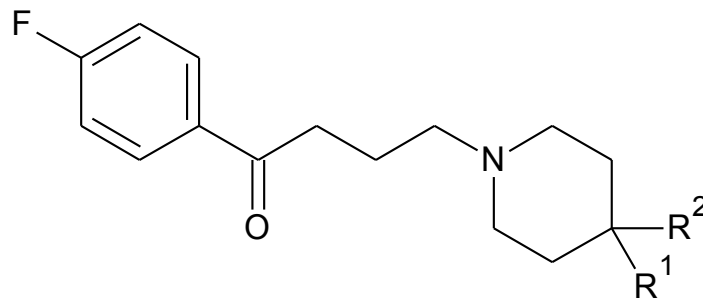


butyrophenone  
analogue of  
pethidine  
• analg. activity  
comparable to  
pethidine, other  
activities similar to  
chlorpromazine



**haloperidol**  
• prototype = lead  
compound of  
butyrophenone  
antipsychotics  
• 10x more active  
than  
chlorpromazine

# Butyrophenones



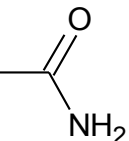
R<sup>1</sup>

OH

OH

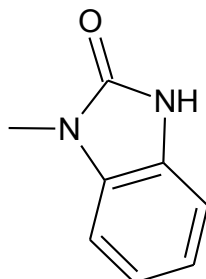
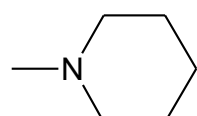
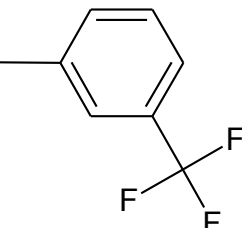
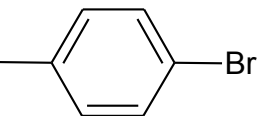
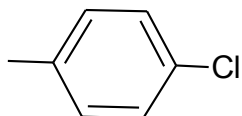
OH

OH



H

R<sup>2</sup>



INN

**melperone**

**haloperidol**

**bromperidol**

**trifluoroperidol**

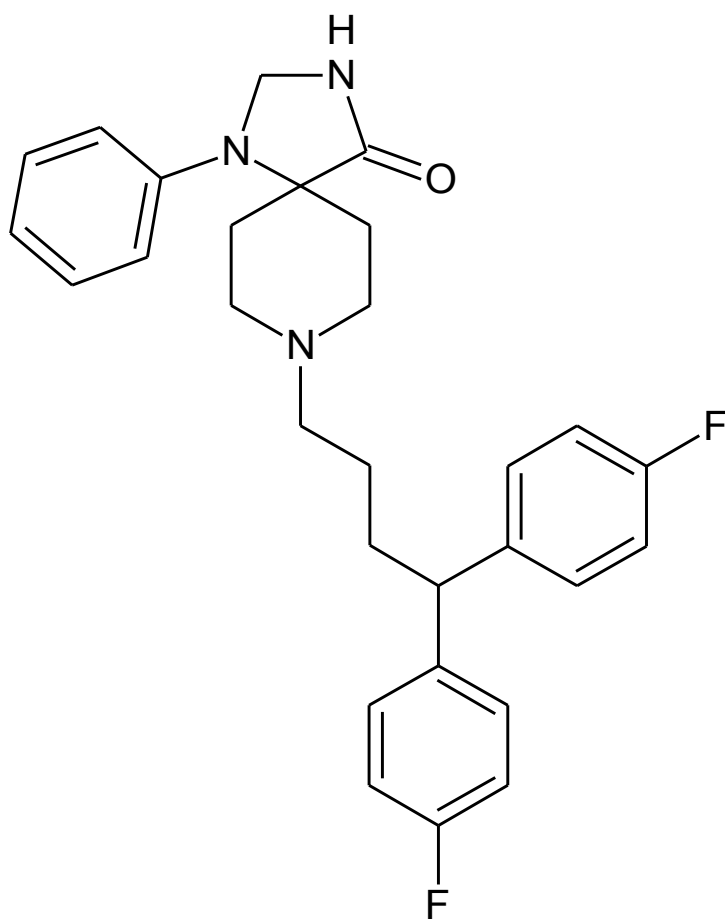
**pipamperone**

LP

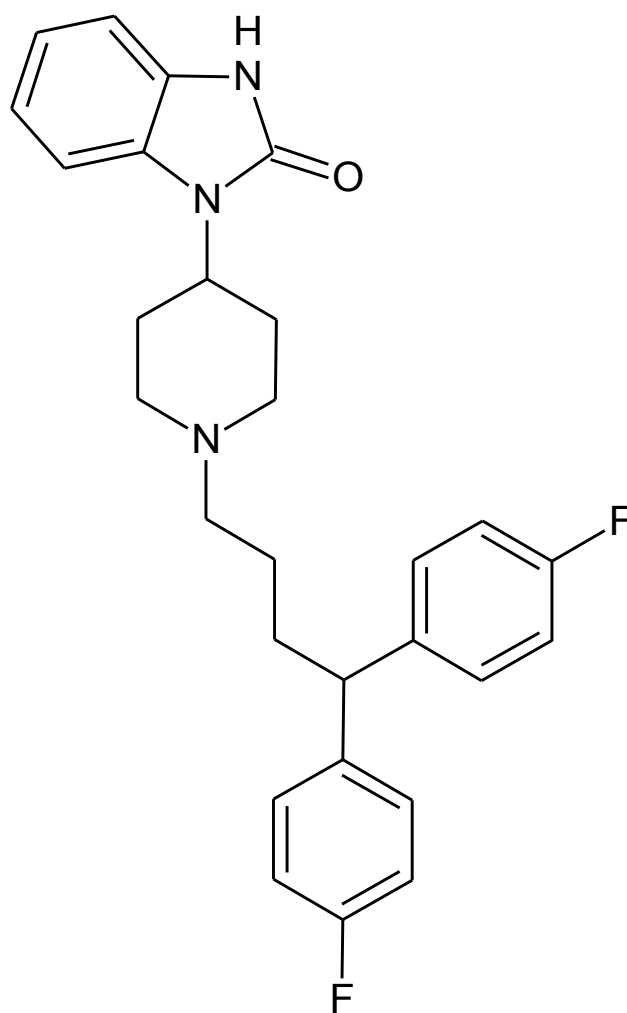
Buronil

Haloperidol Richter

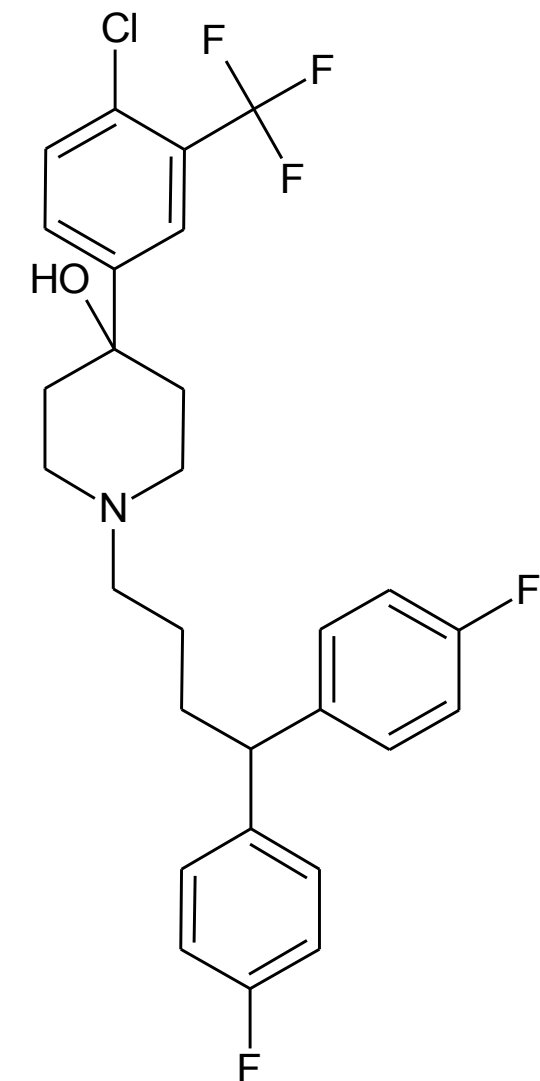
## Diphenylbutylpiperidines



**fluspirilene**



**pimozid**



**penfluridol**

## Butyrophenones and diphenylbutylpiperidines

### Usage:

- treatment of schizophrenia
- neuroleptanalgesia (antipsychotic + opioid analgesic instead general anaesthesia)

### Unwanted effect:

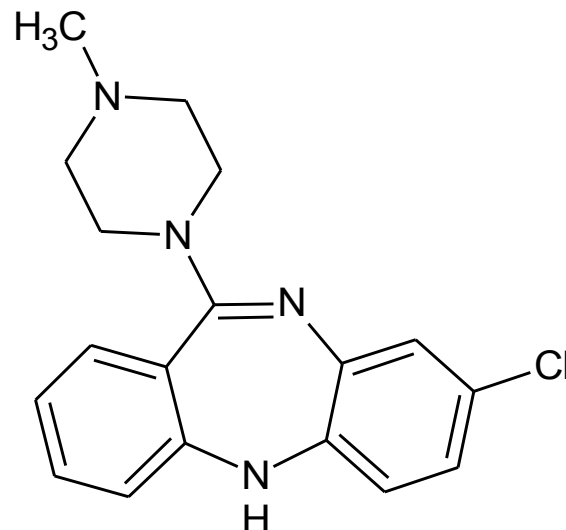
- similar to phenothiazines and thioxanthenes but no extrapyramide syndrome

## „Atypical“ neuroleptics

- influence serotonergic system in addition to the dopaminergic one

### Tricyclic compounds\*

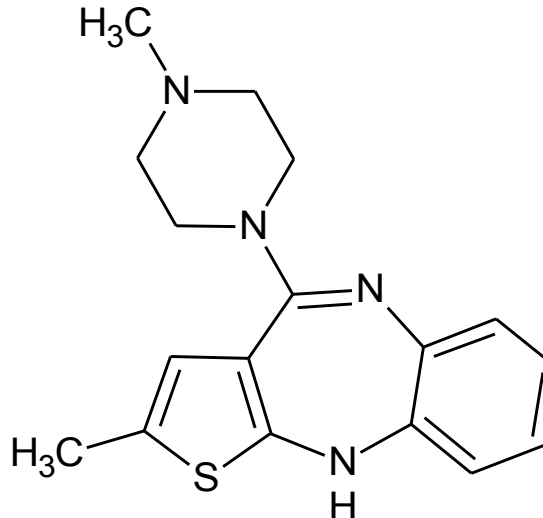
#### Orthocondensed diazepines



8-chloro-11-(4-methylpiperazin-1-yl)-5*H*-dibenzo[*b*,*e*][1,4]diazepine

#### **clozapine**

Clozapin Desitin®, Leponex®

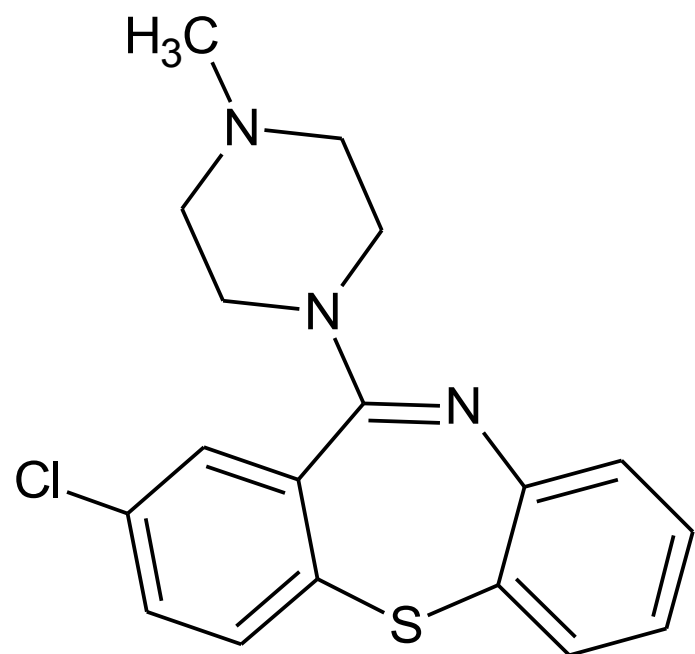


2-methyl-4-(4-methylpiperazin-1-yl)-10*H*-thieno[2,3-*b*][1,5]benzodiazepine

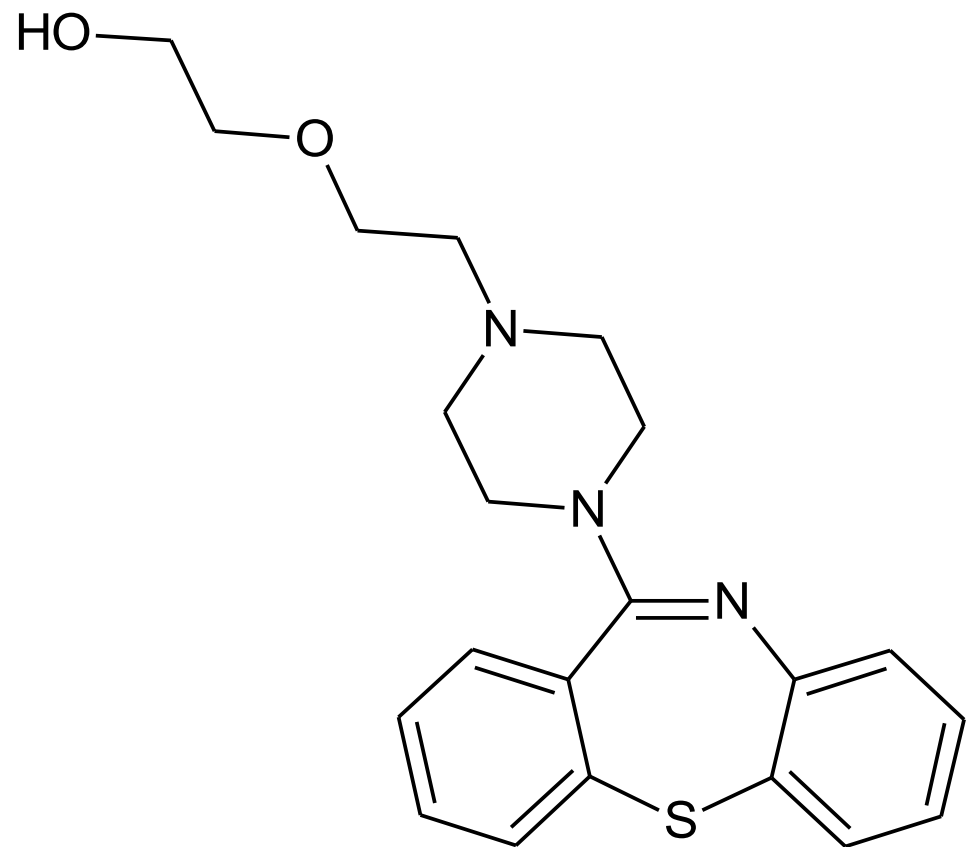
#### **olanzapine**

Zalasta®, Zyprexa® ...

Tricyclic compounds  
Orthokondensed thiazepines



**clothiapine**



**quetiapine**  
Derin<sup>®</sup>, Uxipptra<sup>®</sup> ...

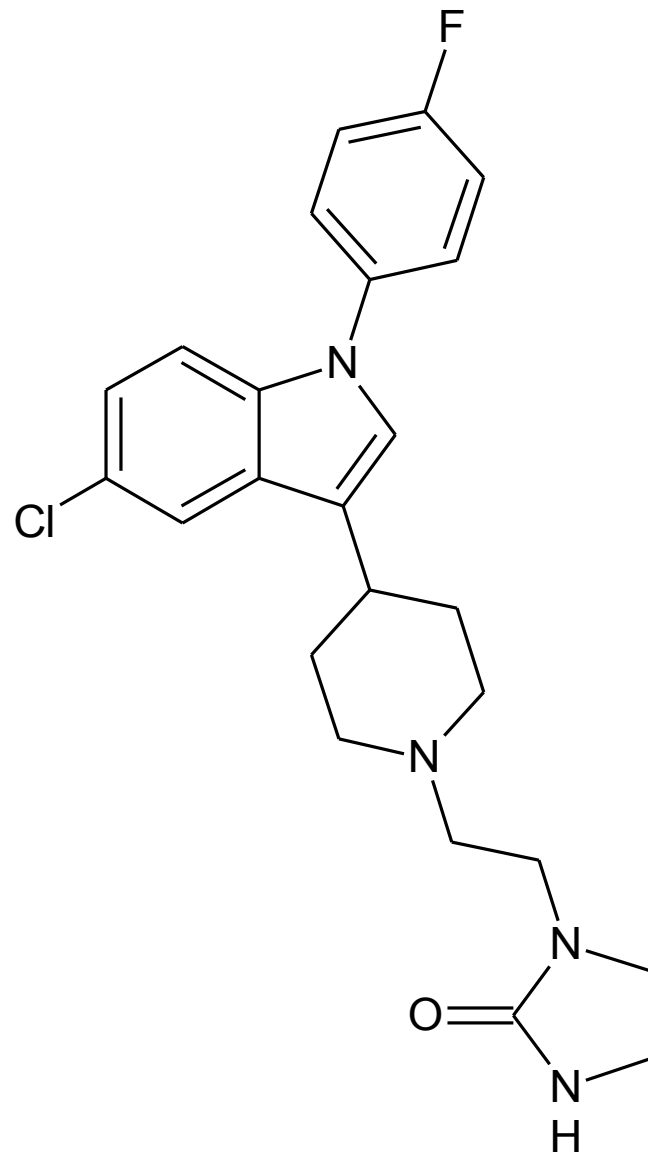
Mechanism of action of tricyclic atyp. neuroleptics:

- serotonine antagonists on 5-HT<sub>2A/2C</sub> receptor subtype
- strong affinity to dopaminergic receptors but weak to D<sub>2</sub> subtype

Unwanted effects:

- agranulocytosis
- cardiovascular system: orthostatic hypotension, TdP dysrhytmias

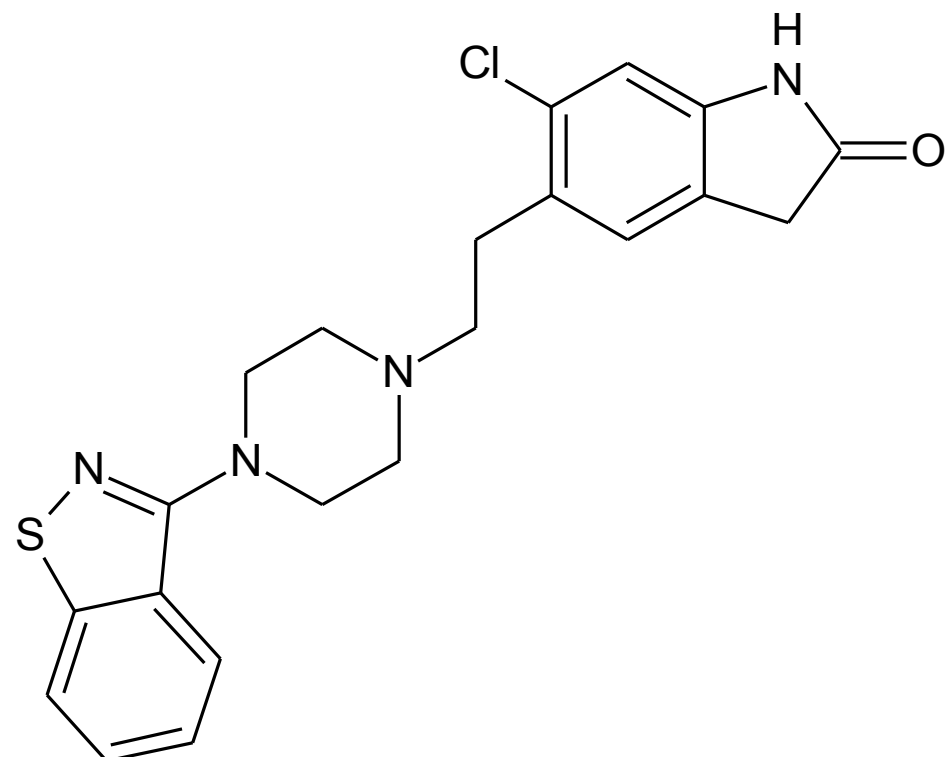
## Indol derivatives



**sertindole**

- $5\text{-HT}_2$  and  $\text{D}_2$ -rp. antagonist

Serdolect®



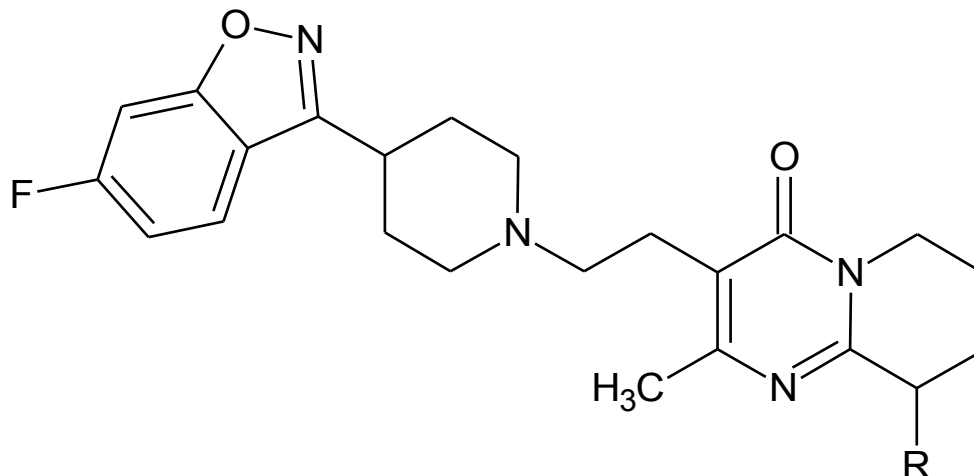
**ziprasidone**

- $\text{D}_2$ -antagonist

- extrapyramidal syndrome occurs but less than in „typical“ antipsychotics

Zeldox®, Zypsila® ...

## Benzoisoxazole derivatives



R = H

**risperidone**

Ridonér®, Rigenin® ...

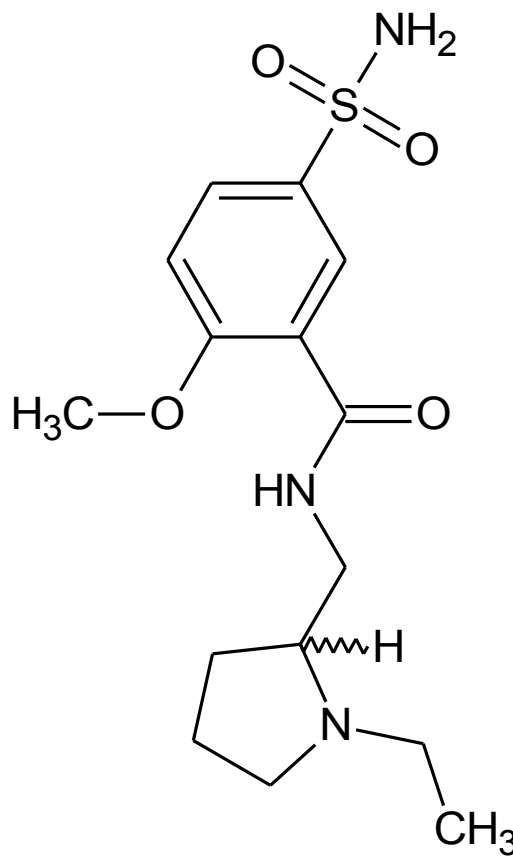
R = OH

**paliperidone**

Invega®

- selectively block D<sub>2</sub> and 5-HT<sub>2</sub> receptors
- inhibit both positive and negative syndroms
- AE & toxicity: somnolence, ECG changes, altered perception

## Benzamide derivatives



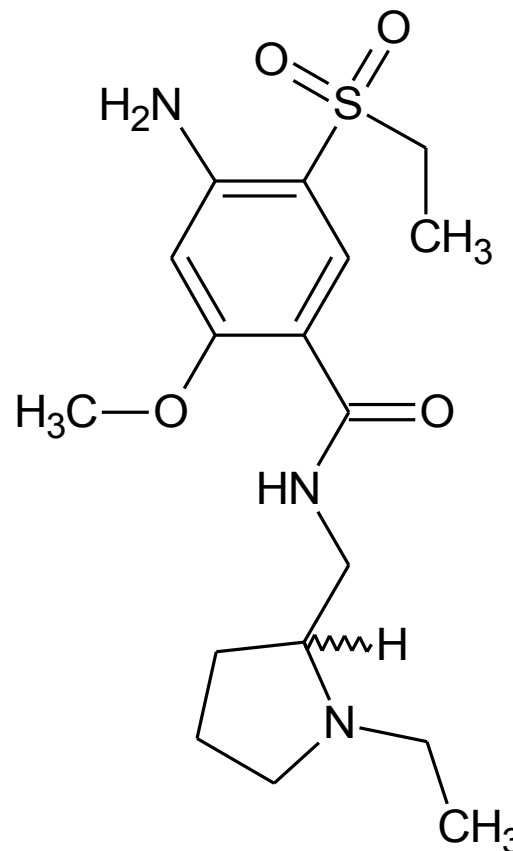
Dogmatil®, Sulpirol® ...

- selective antagonist of D<sub>2</sub>-receptor

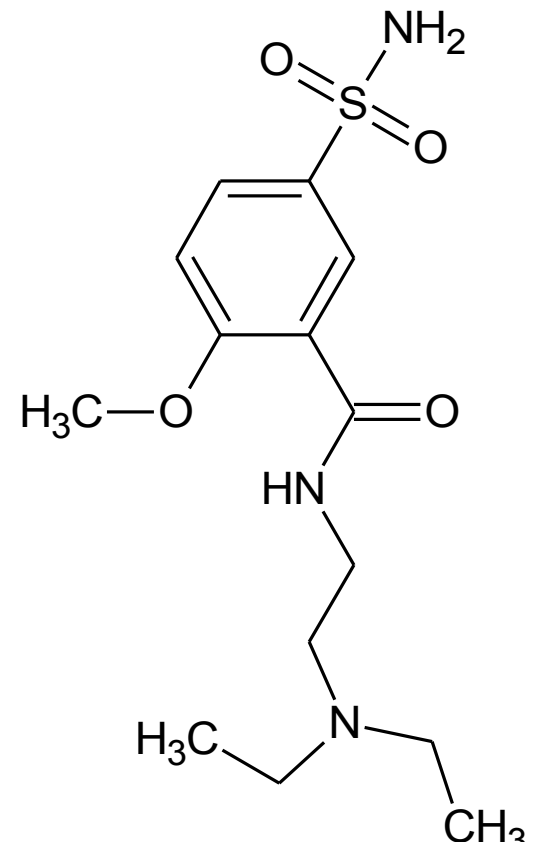
- in lower doses antidepressant – inhibit presynaptic D<sub>2</sub>-receptors, in

- higher doses postsynaptic ones ⇒ antipsychotic

S(-): **levosulpiride**



Amilia®, Deniban® ...



Tiapra®, Tiapridal® ...