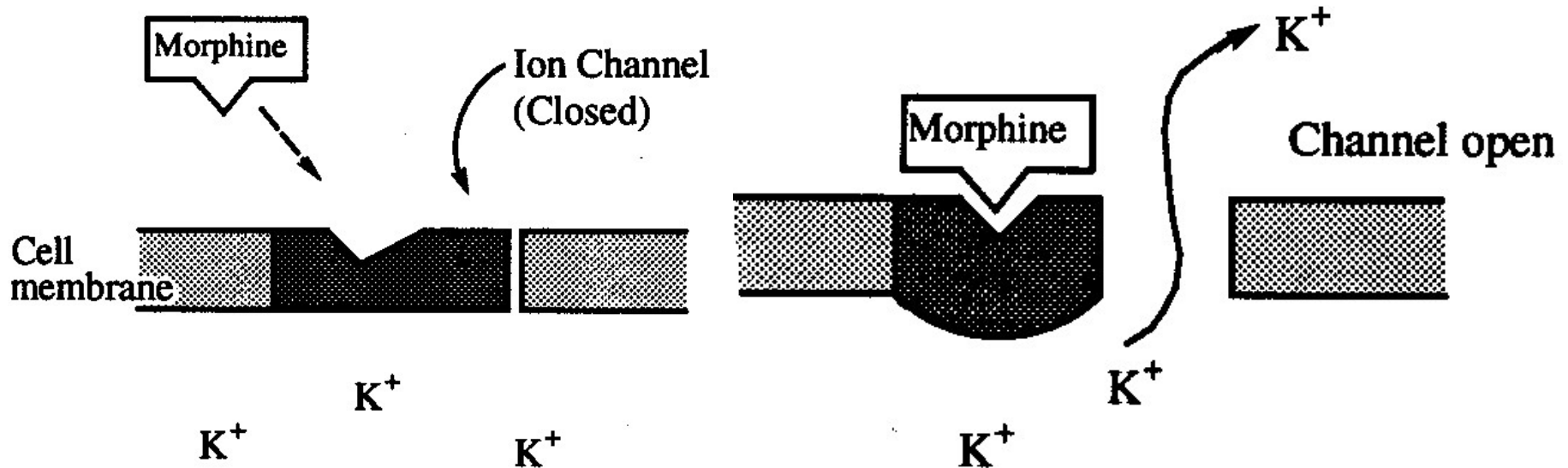


Analgesics – anodyns = opioid = „strong“ = „narcotic“
analgesics

Opioid receptors

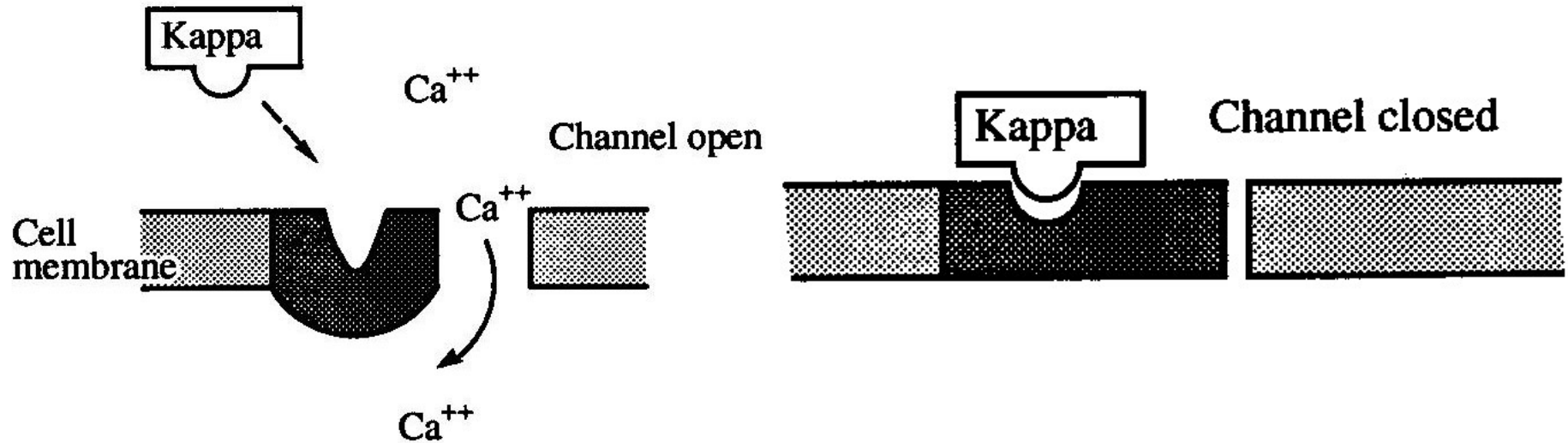
- 4 main types: μ , κ , δ and ORL-1
- (σ receptors currently not recognized as opioid)
- every type has several subtypes
- μ receptor activation leads to analgesic activity, breathing attenuation etc.
- κ receptor activation: also analgesic activity, takes part in diuresis and neuronal activity regulation
- δ receptor activation: also analgesic activity, attenuation of breathing and peristalsis of GIT

μ receptor



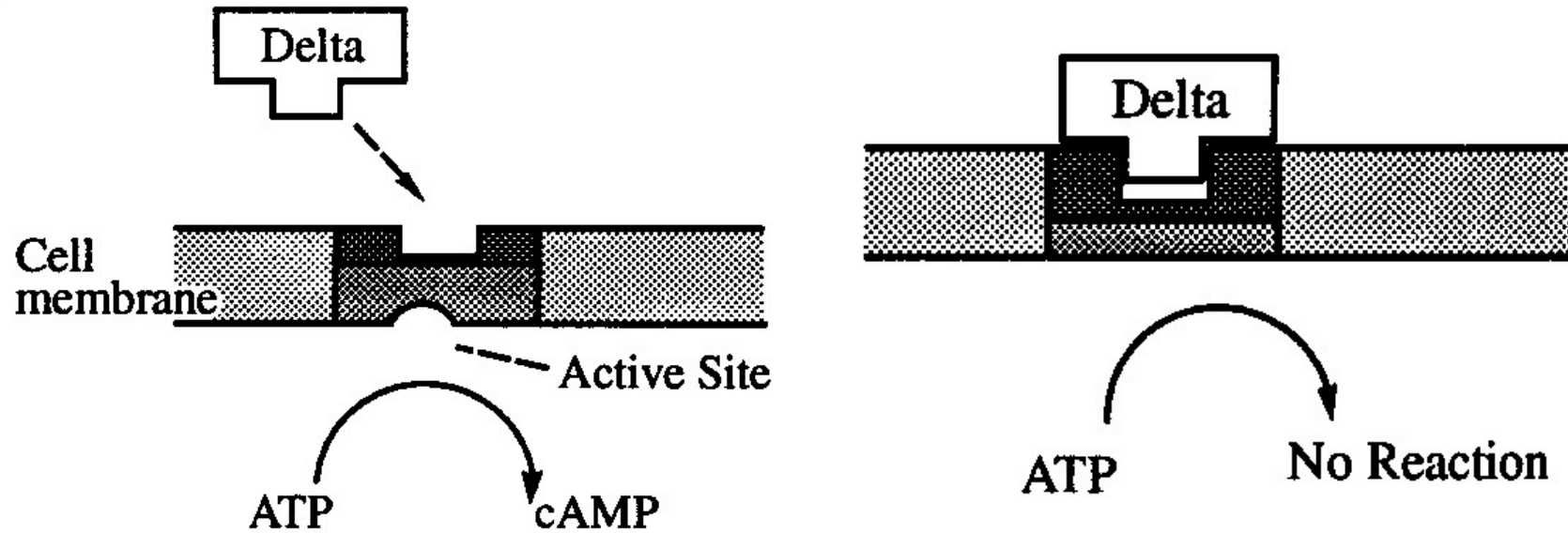
- opens ion channel in cell membrane
- K^+ can stream into the cell \Rightarrow decrease of neurone excitability
- also decreases input of Ca^{2+} into terminal nerve which decreases neurotransmitter release

K receptor



- directly linked with Ca²⁺ channel
- binding of an agonist to the receptor causes channel closing
- inhibition of all nociceptive signals
- activation leads to myosis, diuresis, analgesia and dysphoria

δ Receptor



- not linked with any ion channel
- activation of the receptor probably leads to a change of adenylate cyclase geometry \Rightarrow active site closure
- activation leads to pain relief, attenuation of breathing and peristalsis of GIT

ORL-1 receptor

- also „orphan“, discovered quite recently
- natural agonist nociceptine = orfanine (peptide)
- linked with many activities: memory, cardiovascular functions, kidneys
- probably influences dopamine concentrations in CNS and is involved in neurotransmitters release in anxious conditions

Natural opioid receptors agonists – endogenous analgesics

- morphine receptors exist although it is not endogenous \Rightarrow body own opioids must exist!
- all endo-opioids are fragments of β -lipotropin, adenohypophyse hormone consisting of 91 amino acid rests which has no opioid effects

Enkephalins – binding preferably to δ -receptors

Met-enkephalin

H-Tyr-Gly-Gly-Phe-Met-OH

Leu-enkephalin

H-Tyr-Gly-Gly-Phe-Leu-OH

- **pentapeptide, all activities of morphine, occurs in all animals including man**

Endorphins (= „endo-morphines“) -
in CNS

α : 16 AA

β : 31 AA – after *i.v.* application has morphine effects

χ : 17 AA

- β -lipotropin is not direct precursor of opioid peptides; more precursor peptides exist:

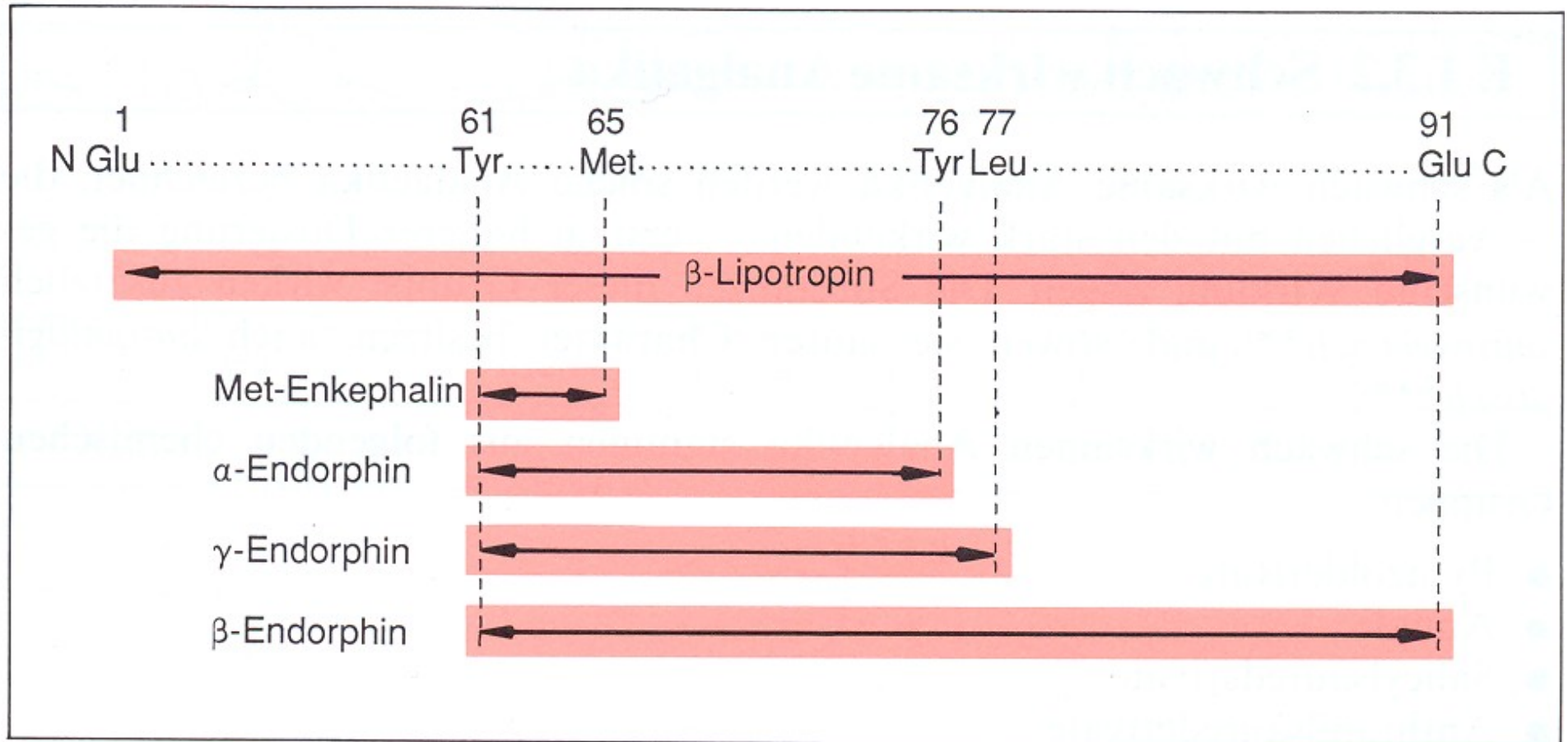
preproenkephalin A \Rightarrow enkephalins

preproopiomelanocortin \Rightarrow endorphins

preproenkephalin B \Rightarrow dynorphins

Dynorphins – peptides from 8 -32 AA, analgesic effect, neurotransmitters in CNS, functions not completely clear

Endorphines and encephalins as parts of β -lipotropine sequence



Dynorphine A sequence (1 - 18)

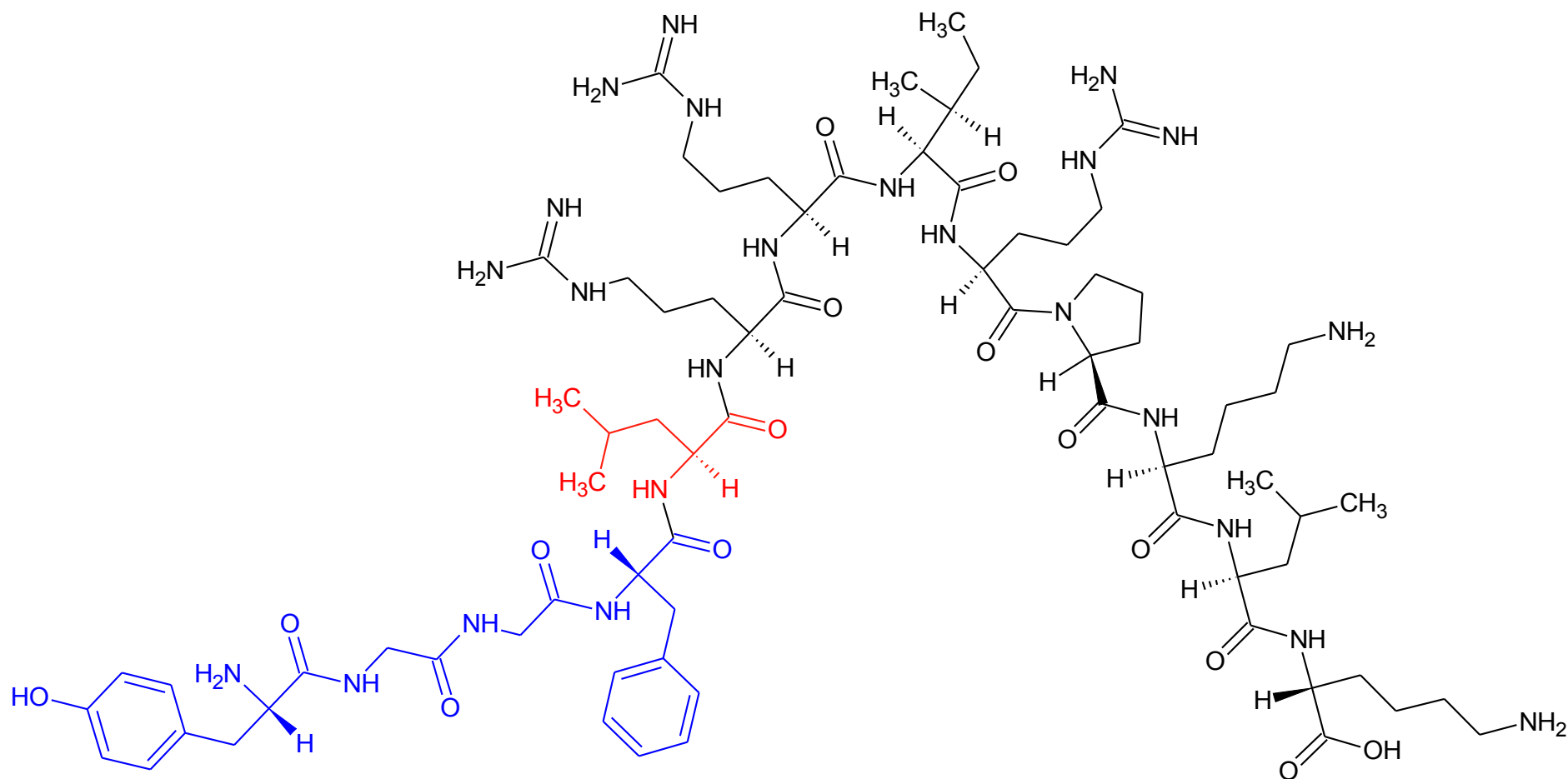
H-Tyr-Gly-Gly-Phe-Leu-Arg-Arg-Ile-

5

Arg-Pro- Lys-Leu-Lys-Trp-Asp-Asn-Gln-OH

10

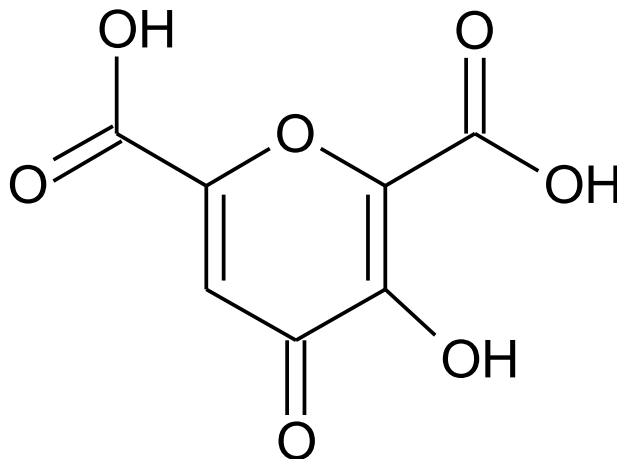
15



Primary structure of dynorphine A (1 – 13) - swine

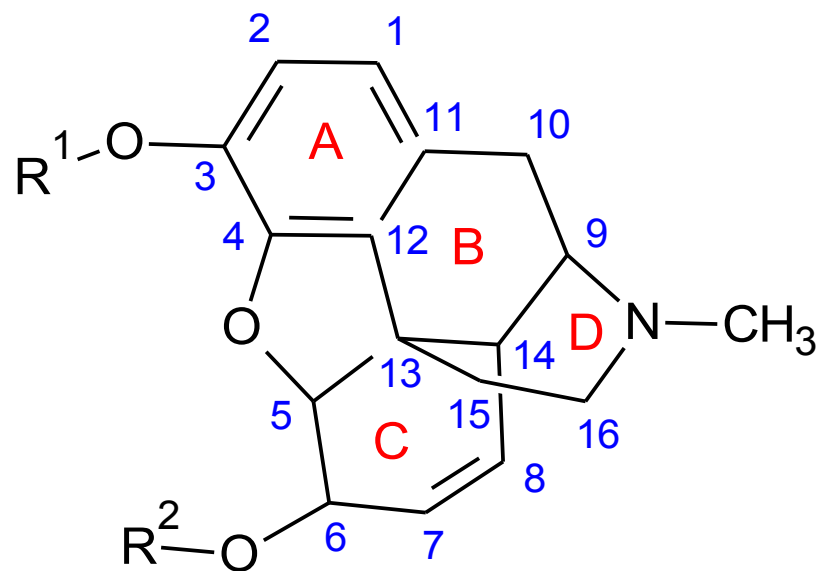
Opium

- dry milky juice (latex) from immature poppy heads (*Papaver somniferum*)
- known from Assyrian manuscripts from 7th century b. C.
- contains 20 – 25 % alkaloids: morphine 3 – 23 %, narcotine 2 – 12 %, codeine 0.3 – 3 %, papaverine 0.8 – 1.2 %; in sum up about 40 various alkaloids
- morphine the most important
- alkaloids in form of salts with carboxylic acids; meconic acid typical
- beaten-out empty dry poppy heads are alternative resource of opium alkaloids (CZ, SK)



meconic acid

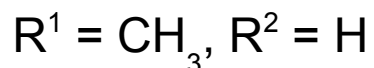
Morphine and its simple derivatives



morphine

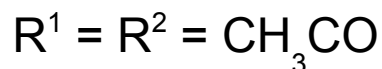
MSI[®], MST[®], Sevredol[®] ...

- isolated by Friedrich Wilhelm Sertürner, pharmacist in Paderborn, from opium in 1806
- structure elucidation: 1925 Robinson and Gunland proposed structural formula, 1952 Gates and Tschudi confirmed structure including stereochemistry by means of total synthesis
- basic anodyne isolated from opium or beaten-out empty dry poppy heads



codein

- basic antitussive
- semi-synthetic; prepared from morphine by selective methylation of phenolic group
- potencuje účinek slabých analgetik potentiates effect of weak analgesics
- abot 10 % methabolized to morphine

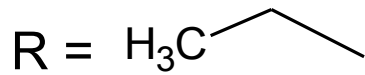
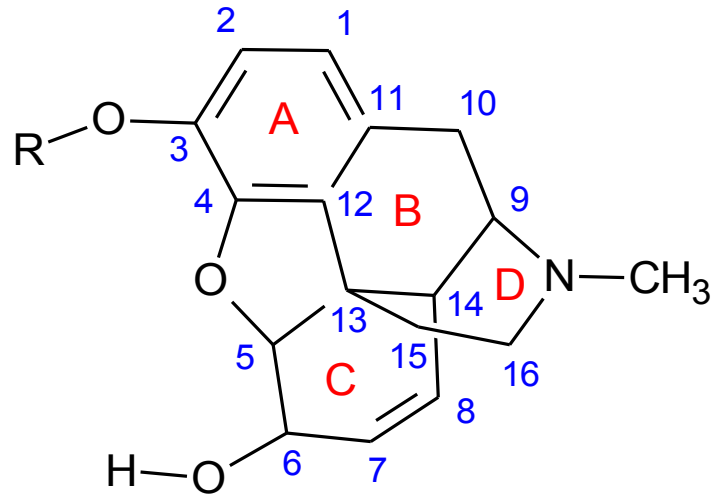


diamorphine

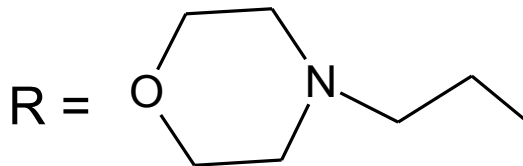
syn. heroine

- 2x more effective than morphine, better penetrates into CNS
- misused as an illegal drug of abuse

Morphine and its simple derivatives: further ethers used as antitussives



ethylmorphine
Diolan ®

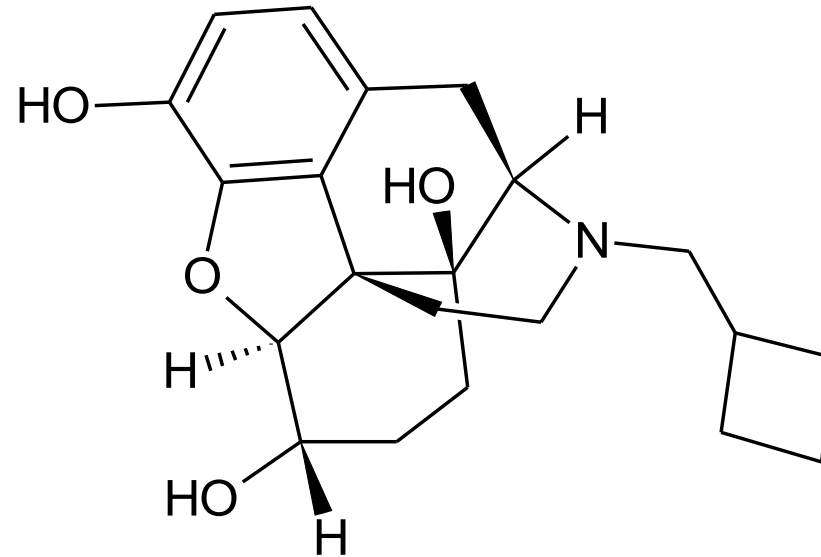


pholcodine
Neocodin ®

Effects of opioid analgesics

- analgesic
- antidiarrhoic (σ -, δ -receptors in gut)
- antitussive – from attenuation of cough reflex to expiratory centre inhibition
- euphoriant
- physical addiction – very slowly formed during relieving of strong pain

Semi-synthetic morphine and codeine derivatives

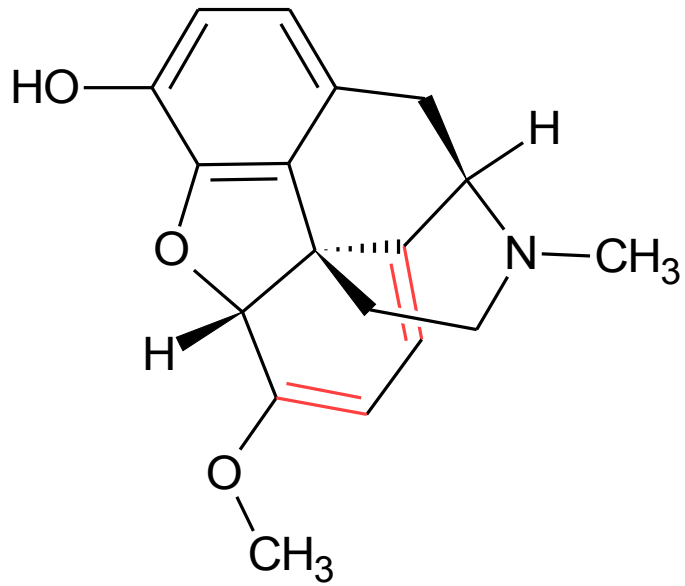


nalbuphine

Nalbuphin[®] OrPha inj.

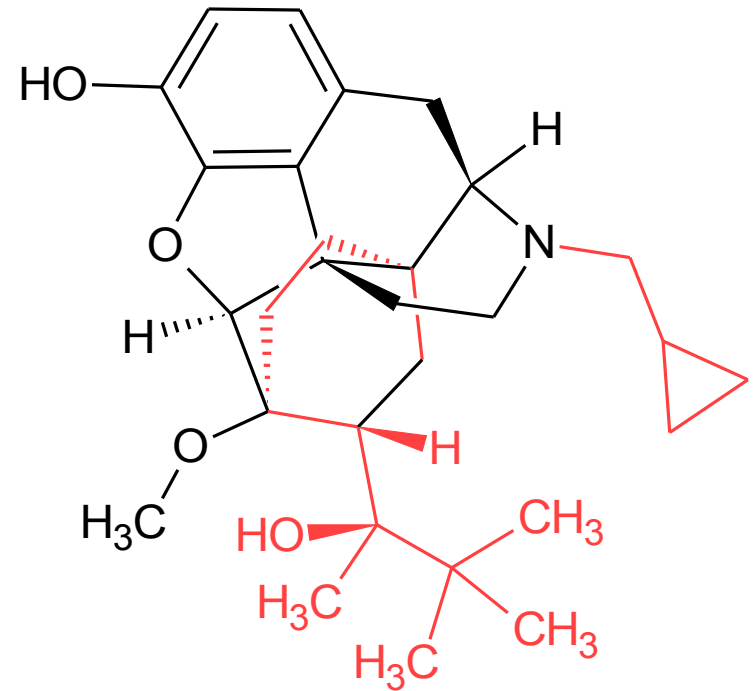
- short period treatment of medium to strong pains, before- and post-operating analgesia

Oripavine derivatives



oripavine

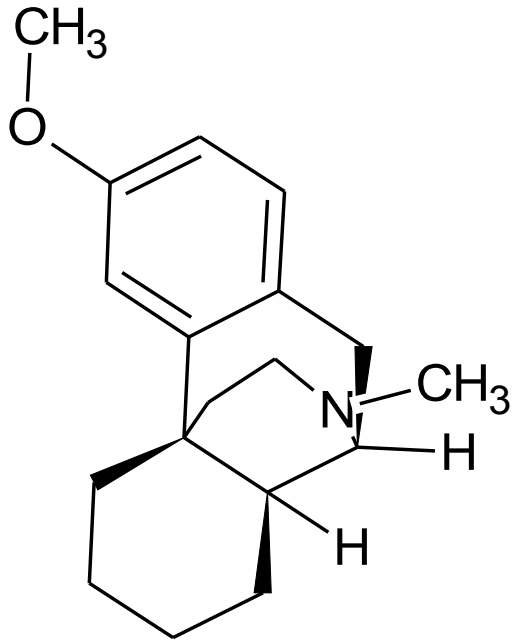
- alkaloide from *Papaver orientale*



buprenorphine

- relief of non-malignant pain of medium intensity
- opioid withdrawal therapeutic programs
Norspan[®] emp. tdr., Transtec[®] emp. tdr.

Morphinane derivatives



dextromethorphan

Dextrometorphani hydrobromidum

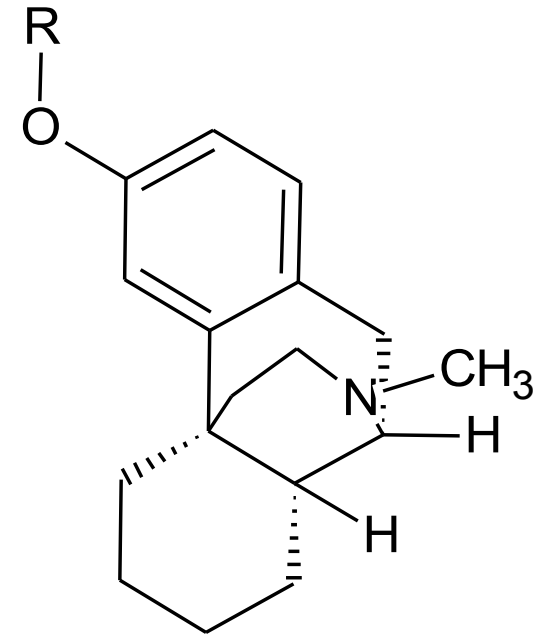
monodydricum PhEur

•antitussive

•euphoriant in higher doses

Humex ® , Robitussin ® , Stopex ® ,

Tussidril ® - OTC

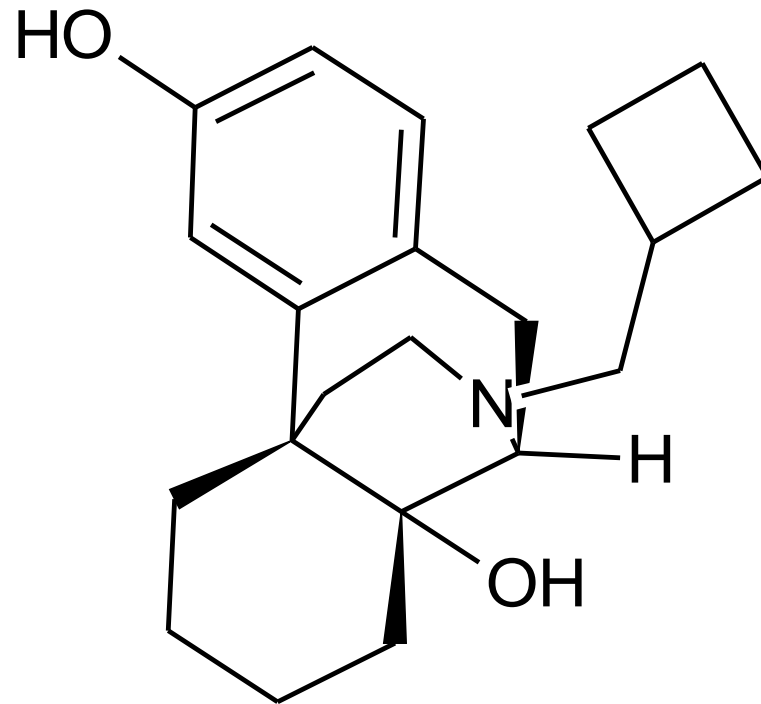


R = H **levorphanol**

•better analgesic than morphine

R = CH₃ levomethorphan

Morphinane derivatives (continued)

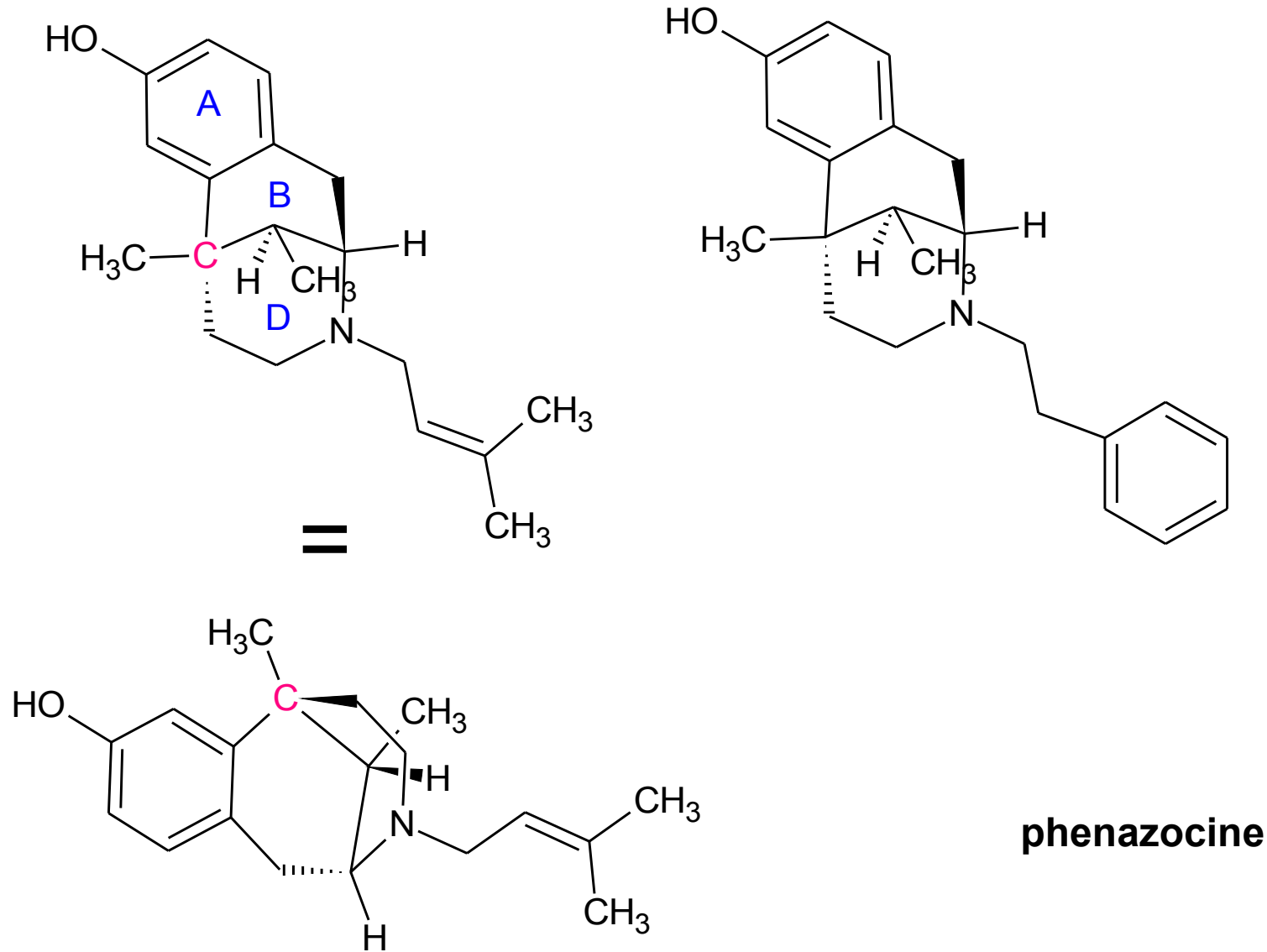


butorphanol

- treatment of moderate to severe pain
- a potent κ -receptor agonist and an antagonist at μ -receptor
- intensive hepatic first-pass metabolism \Rightarrow parenteral administration (nasal sprays)

Benzomorphone derivatives

- removal of the C-ring \Rightarrow greater affinity for κ -receptor; weak for μ -receptor
- central C-atom remained quaternary \Rightarrow truncated open analogues of the C-ring



pentazocine

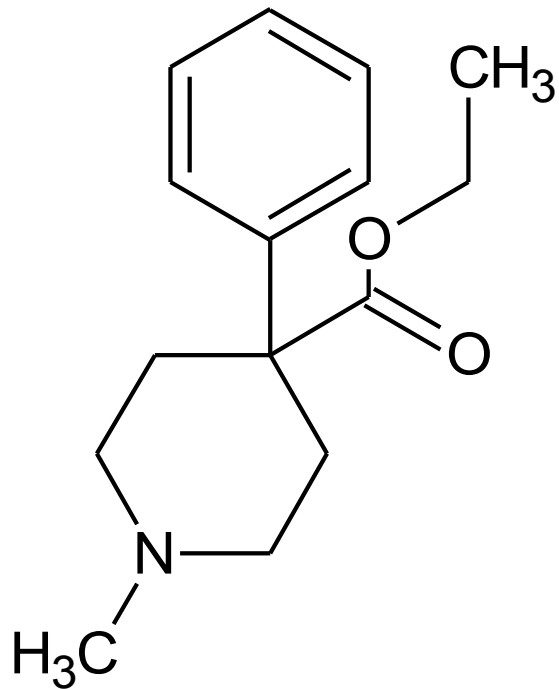
- treatment of moderate pain

Fortral[®] tbl., inj. sol.

phenazocine

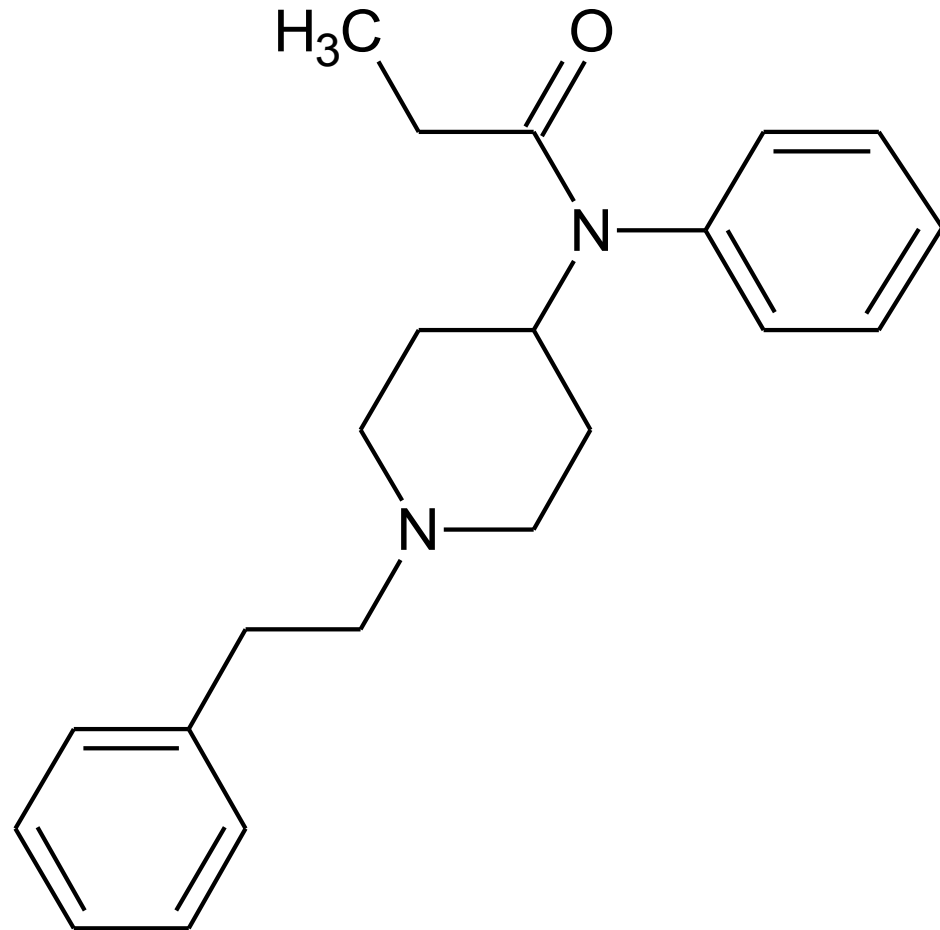
Phenylpiperidine derivatives and compounds derived from them

- originated by removal of the B,C and E rings which are not necessary for the activity
- faster onset and shorter lasting of action
- remaining AE: addiction, expiratory centre attenuation



pethidine

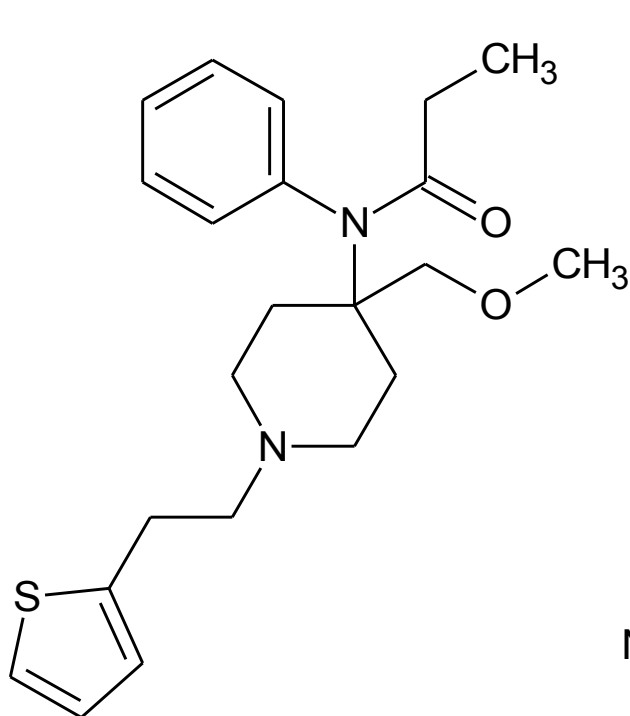
syn. meperidine [USAN]
Dolsin ® inj. sol.



fentanyl

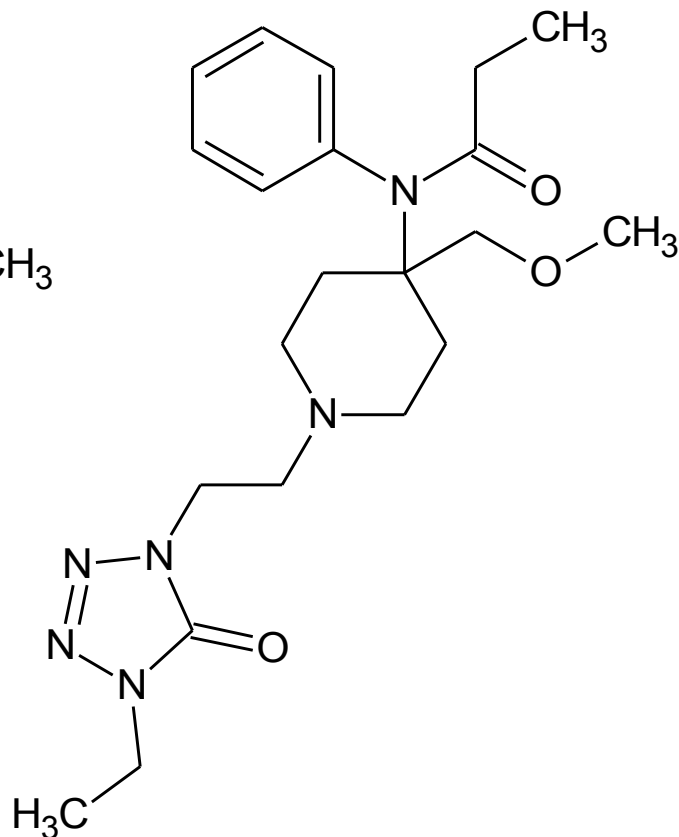
Fentanylum PhEur (free base) – transdermally
Fentanylli citras PhEur – i.m., i.v.
Durogesic ® derm. emp. tdr.

Phenylpiperidine derivatives and compounds derived from them
Fentanyl analogues - 4-anilidopiperidines



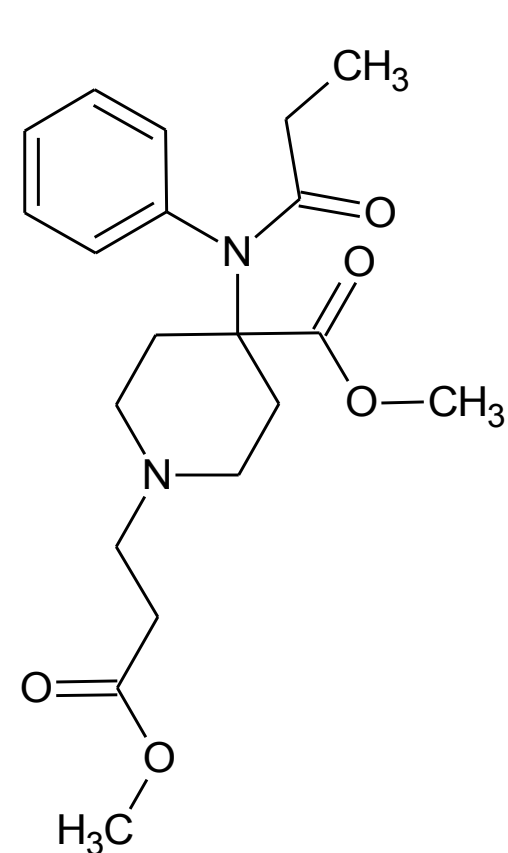
sufentanil

•also in anaesthesia
Sufenta[®] inj.



alfentanil

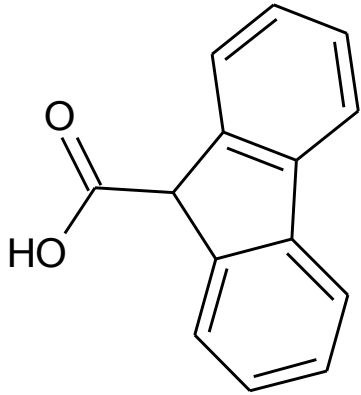
Rapifen[®] inj.



remifentanyl

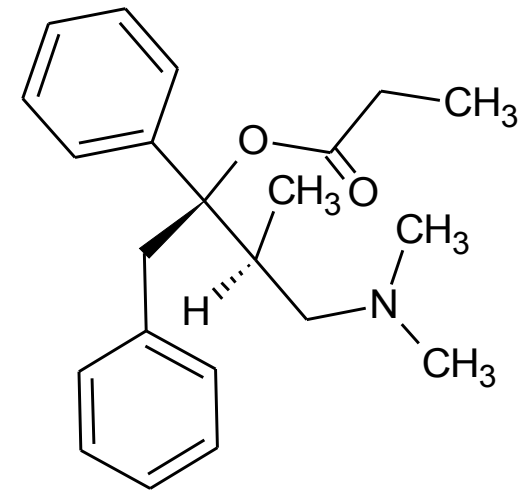
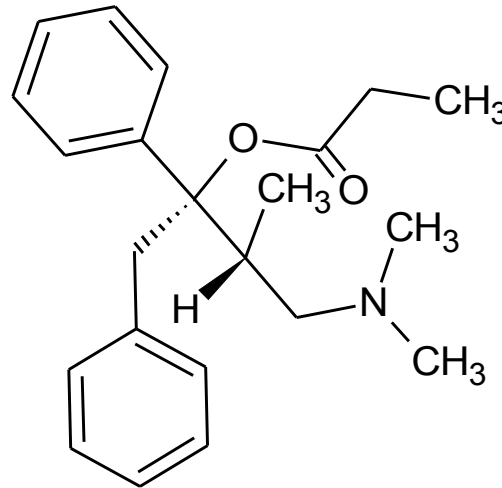
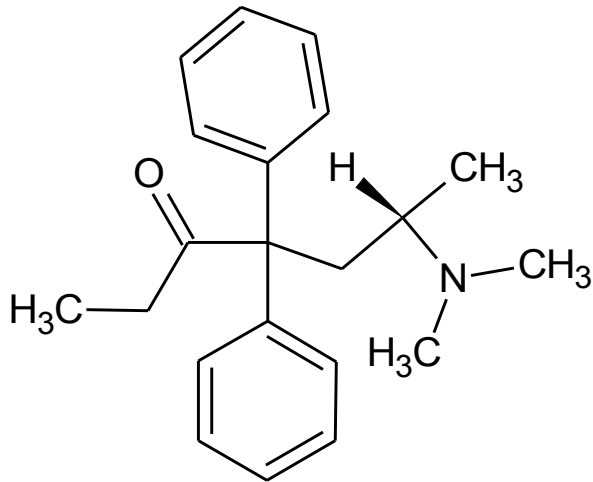
Ultiva[®] inj.

•so called opioid anaesthetics (combined with propofol, ketamine)



Phenylpropylamine derivatives

- can be derived from 4-phenylpiperidines by formal deleting of one methylene group of piperidine ring
- structurally related to fluorene-9-carboxylic acid
- the most simplified structures still to have opioid receptor activity
- activity comparable to morphine
- efficient p.o.
- less AE than morphine



methadone

- withdrawal symptoms less severe and more gradual as compared to morphine \Rightarrow opioid-withdrawal programs
- therapeutical

Methadon Zentiva $\text{\textcircled{R}}$ oral solution

dextropropoxyphene

- substitution of one phenyl with benzyl \Rightarrow 2nd chiral centre
- (+)-(2*S*, 3*R*)-analgesic; 1/10 of methadone activity
- Darvon $\text{\textcircled{R}}$ (USA)

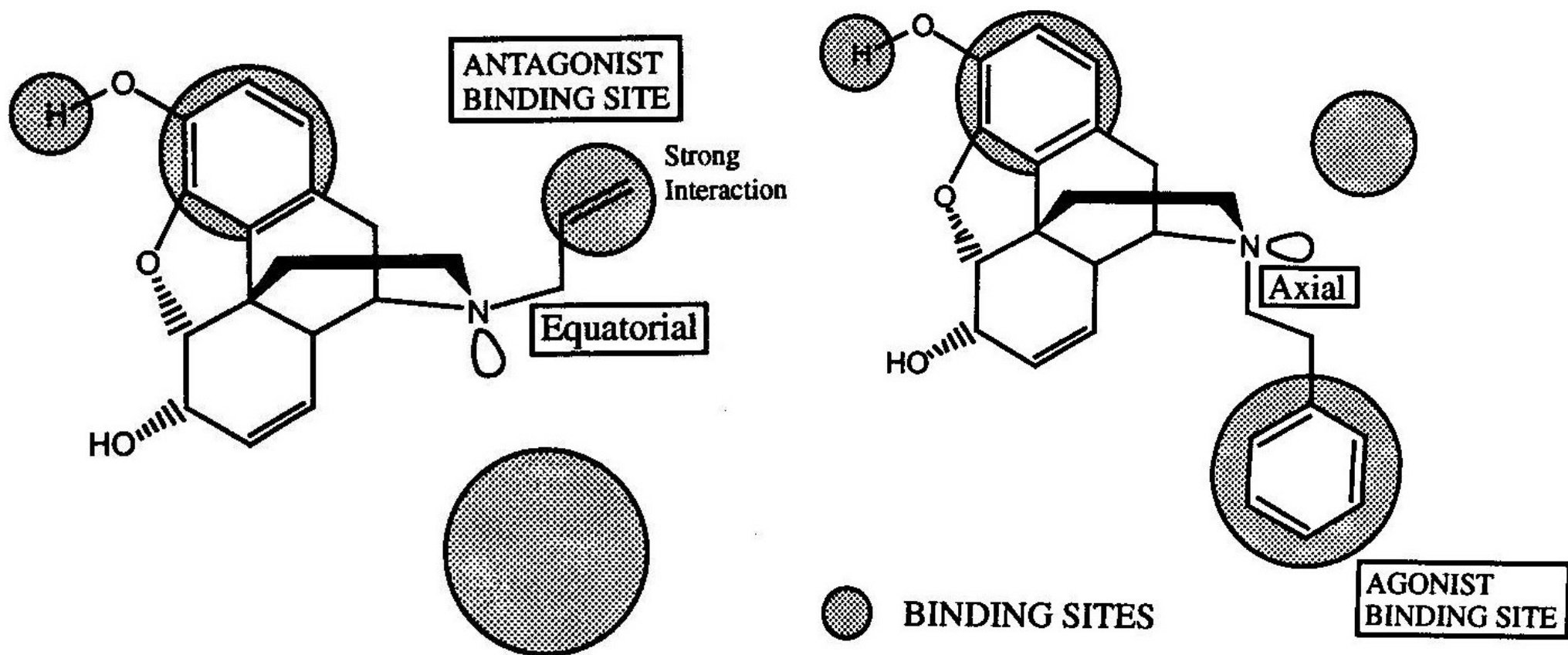
levopropoxyphene

- (-)-(2*R*, 3*S*)-antitussive
- Novrad $\text{\textcircled{R}}$ (USA)

Structure-activity relationships (SAR)

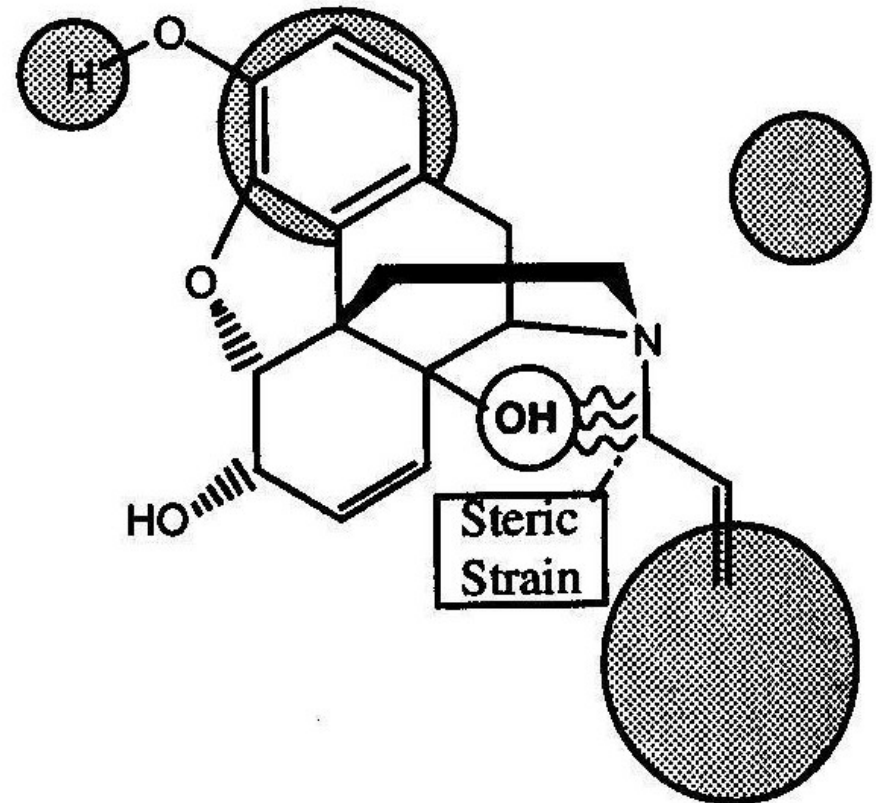
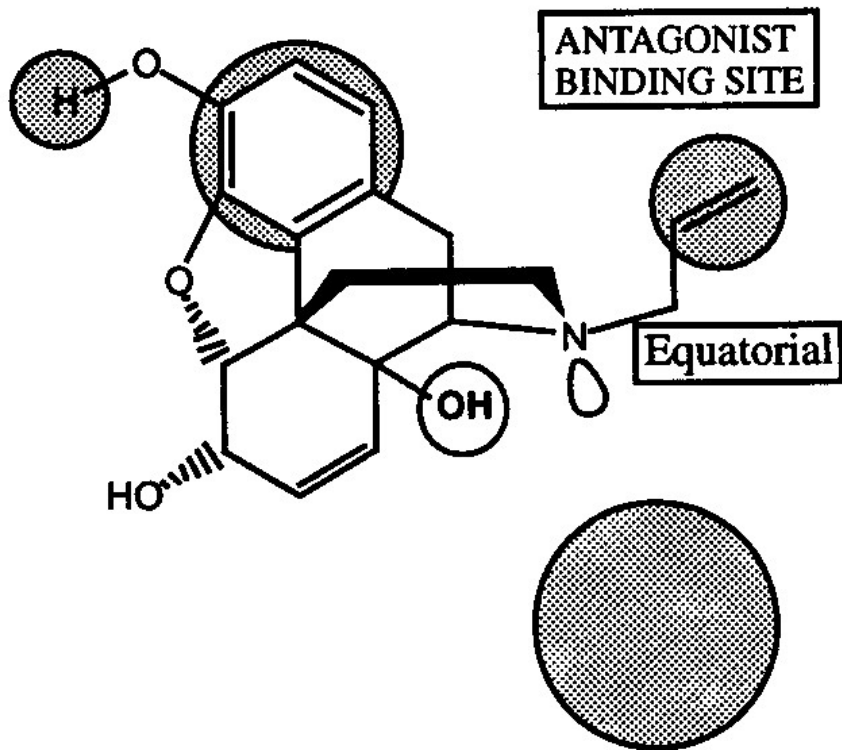
- an aromatic ring and a basic nitrogen atom are necessary for action, a phenolic group is not(\Leftrightarrow the rings B, C, D and E of the morphine skeleton are not necessary for analgesic action)
- quarternary (tetrasubstituted) C(4) of piperidine derivatives is necessary in the frame of this group, with exception of fentanyl
- substitution of methyl at nitrogen in D ring of morphine: to **allyl** leads to antagonists (equatorial position), to **phenethyl** leads to agonists; explanation by presence of 2 different hydrophobic binding sites
- OH group at C(14) supports the placement of a substituent into equatorial position; this moiety as a steric hindrance orients into equatorial position also other substituents than allyl (e.g. cyclopropylmethyl)

N-substituted morphine derivatives acting as agonists and antagonists – a model of interaction with a receptor

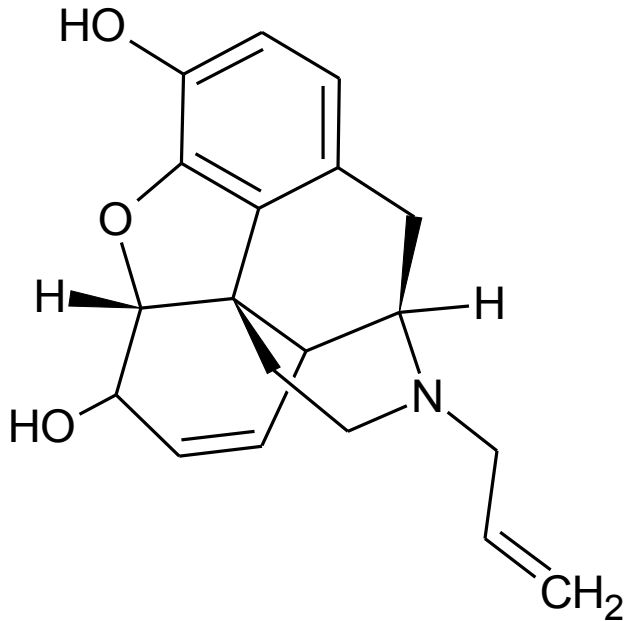


Morphine receptor antagonists with -OH group at C(14)

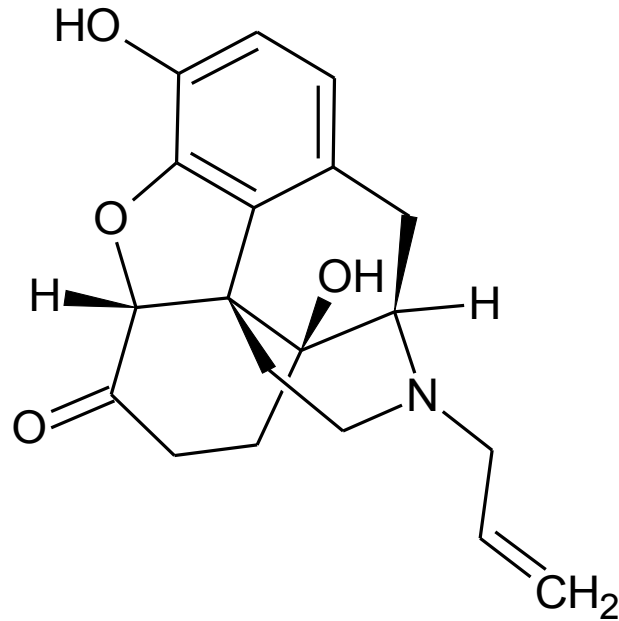
- OH group is a steric hindrance which supports to dominance of the equatorial position of allyl



Morphine receptor antagonists



nalorphine



naloxone

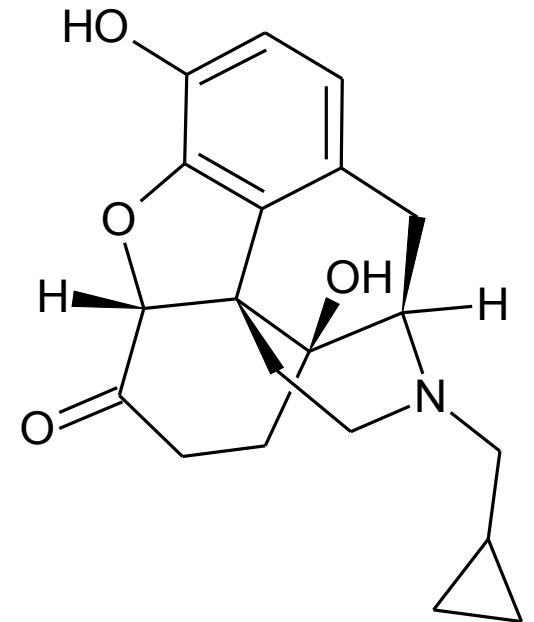
Naloxoni hydrochloridum dihydricum PhEur

- *i.v.* administration only; extensively

metabolised in liver

Naloxone WZF ® Polfa

- antidotes in opiates overdose



naltrexone

Naltrexoni hydrochloridum PhEur

- *p.o.*

- useful also in treatment of alcoholism (blocks binding of endo opioids)

Revia ® por tbl flm