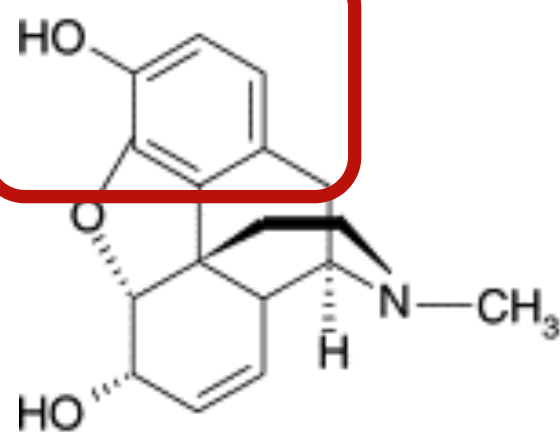
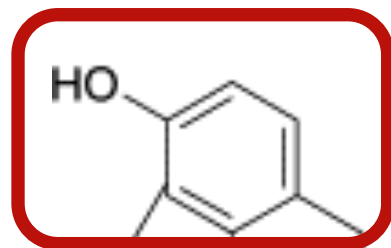


# OPIOIDS



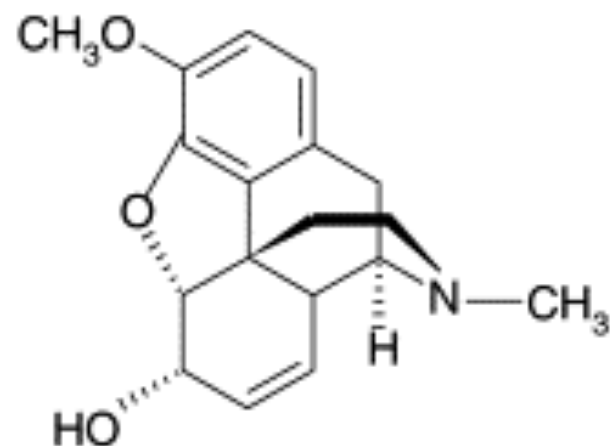
# ALKALOIDS IN OPIUM

Heroin 2 -COCH<sub>3</sub>



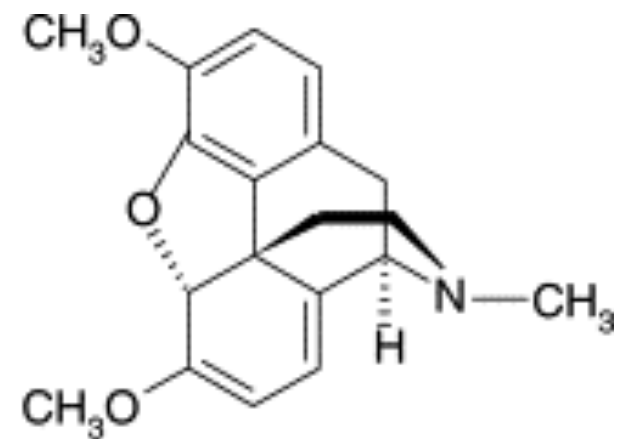
morphine

10%



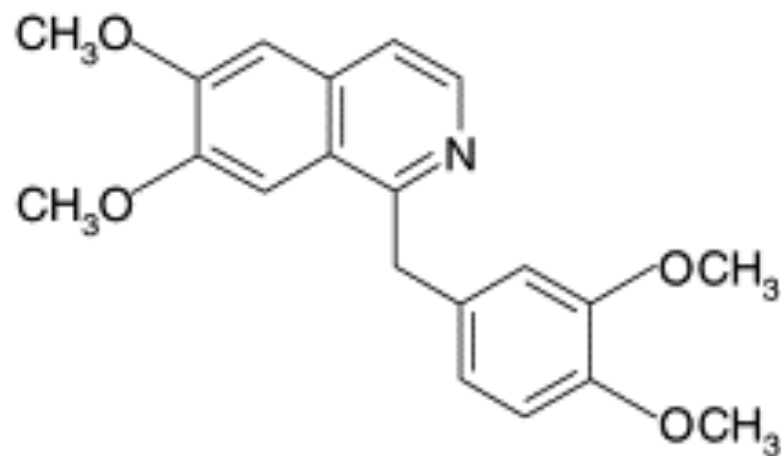
codeine

0,5%



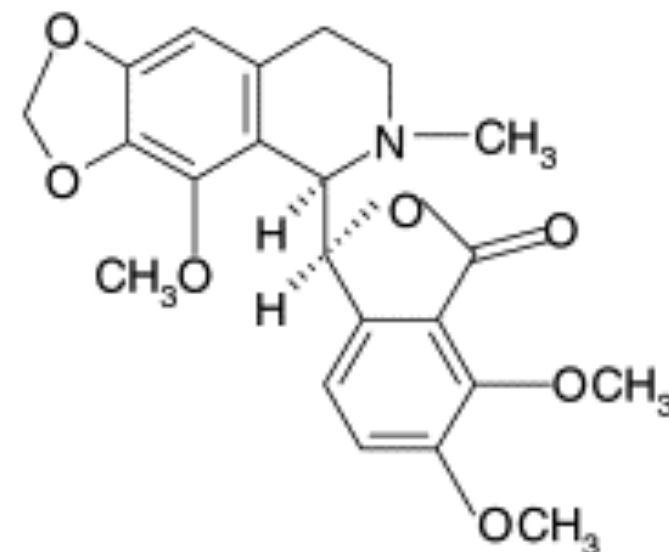
thebaine

0,2 %



papaverine

1 %



noscapine

6 %

% percentage in the opium juice

# PHARMACOLOGICAL ACTIONS OF MORPHINE

## **Central Nervous System**

Analgesia

Euphoria

Sedation

(Dysphoria and hallucinations)

Pupillary constriction

Nausea and vomiting

Respiratory depression

Depression of cough reflex

Tolerance and dependence

## **Gastrointestinal tract**

Reduced motility and increased tone with:

Constipation

Contraction of biliary sphincter

## **Other actions**

Histamine release with:

Urticaria and itching

Broncho constriction

Hypotension and bradycardia

Immunosuppressant effects

# OPIOIDS RECEPTORS AND THEIR LIGANDS

In the 1950s:

proposal of the presence of specific receptors for opioids

In the 1970s:

Proposal of the presence of three different receptors:

- mu receptors (from Morphine) MOR
- kappa receptors (from Ketocyclazocine) KOR
- delta receptors (from Deferent vessels) DOR

Isolation and characterization of endogenous ligands (endorphins):

- Beta-endorphins
- Dynorphins
- Enkephalins

In the 1990s:

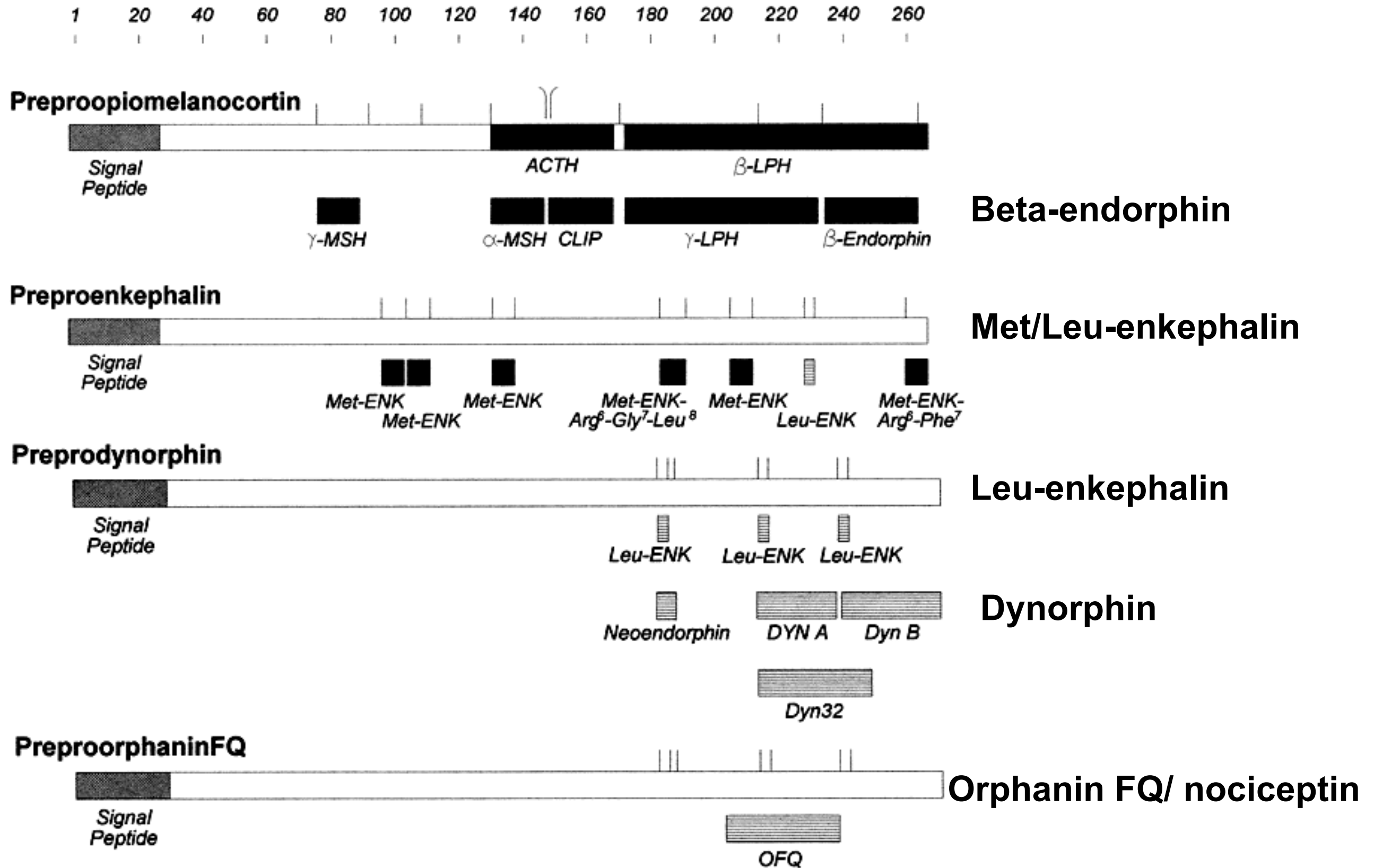
- Cloning of MOR, DOR and KOR GPCRs
- Identification of Orphanin FQ/ nociceptin receptor (no affinity towards naloxone)



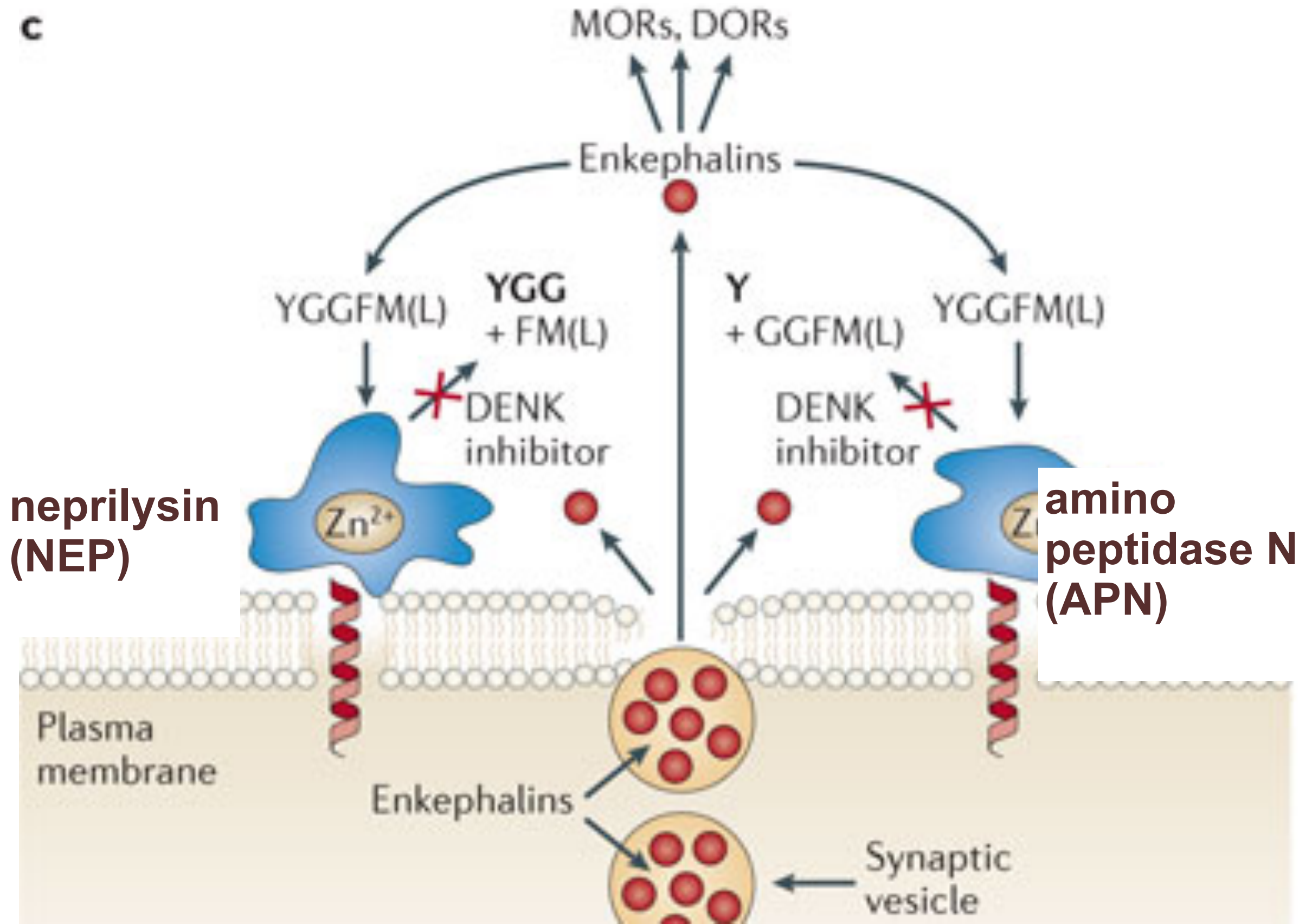
# ENDOGENOUS OPIOID PEPTIDES

<b>OFQ/N</b>	Phe-Gly-Gly-Phe-Thr-Gly-Ala-Arg-Lys-Ser-Ala-Arg-Lys-Leu-Ala-Asn-Gln
<b>OFQ/N(1-11)</b>	Phe-Gly-Gly-Phe-Thr-Gly-Ala-Arg-Lys-Ser-Ala
<b>OFQ/N(1-7)</b>	Phe-Gly-Gly-Phe-Thr-Gly-Ala
<b>OFQ2</b>	Phe-Ser-Glu-Phe-Met-Arg-Gln-Tyr-Leu-Val-Leu-Ser-Met-Gln-Ser-Ser-Gln
<b>ppOFQ/N<sub>160-187</sub></b> (mouse)	Phe-Ser-Glu-Phe-Met-Arg-Gln-Tyr-Leu-Val-Leu-Ser-Met-Gln-Ser-Ser-Gln
<b>Nocistatin</b> (human)	Arg-Arg-Arg-Thr-Leu-His-Gln-Asn-Gly-Asn-Val Met-Pro-Arg-Val-Arg-Ser-Leu-Phe-Gln-Glu-Gln-Glu-Glu-Pro-Glu-Pro-Gly-Met-Glu-Glu-Ala-Gly-Glu-Met-Glu-Gln-Lys-Gln-Leu-Gln
<b>Dynorphin A</b>	Tyr-Gly-Gly-Phe-Leu-Arg-Arg-Ile-Arg-Pro-Lys-Leu-Lys-Trp-Asp-Asn-Gln
<b>[Leu<sup>5</sup>]enkephalin</b>	Tyr-Gly-Gly-Phe-Leu
<b>[Met<sup>5</sup>]enkephalin</b>	Tyr-Gly-Gly-Phe-Met
<b>β-Endorphin</b>	Tyr-Gly-Gly-Phe-Met-Thr-Ser-Glu-Lys-Ser-Gln-Thr-Pro-Leu-Val-Thr-Leu-Phe-Lys-Asn-Ala-Ile-Ile-Lys-Asn-Ala-His-Lys-Lys-Gly-Gln
<b>Endomorphin 1</b>	Tyr-Pro-Trp-Phe-NH <sub>2</sub>
<b>Endomorphin 2</b>	Tyr-Pro-Phe-Phe-NH <sub>2</sub>

# ENDOGENOUS OPIOID PEPTIDES: SYNTHESIS



# METABOLISM OF ENDOGENOUS PEPTIDES



# SELECTIVITY OF OPIOID LIGANDS

	MOR	DOR	KOR	NOR
<b>ENDOGENOUS OPIOIDS</b>				
Beta-endorphin	+++	+++	+	-
Leu-enkephalin	(++)	+++	+	-
Met-enkephalin	++	+++	+	-
Dynorphin	+	+	+++	-
Orphanin FQ/nociceptin	-	-	-	+++
<b>RECEPTOR SELECTIVE DRUGS</b>				
Agonists				
DAMGO	+++	-	-	-
DPDPE	-	++	-	-
Enadoline	-	-	+++	-
Ro64-6198	-	-	-	+++
	-	-	-	-
Antagonists				
CTOP	+++	-	-	-
Naltrindole	-	+++	+	-
Nor-binaltorphimine	+	+	+++	-
SB 612111	-	-	-	+++



## MOR RECEPTORS SELECTIVE LIGANDS

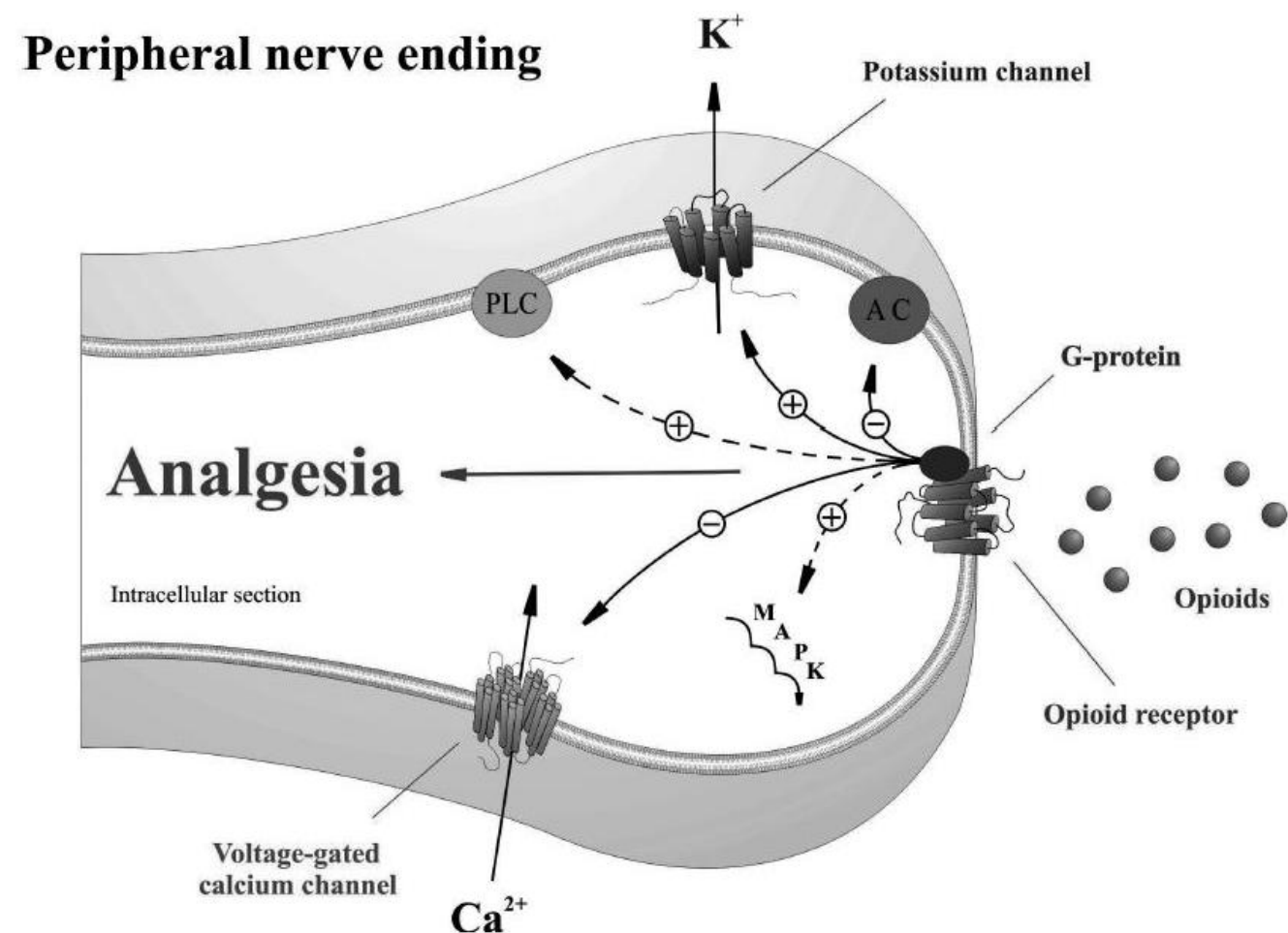
AGONISTS	PARTIAL AGONISTS	ANTAGONISTS
<b>Morphine</b> (T <sub>1/2</sub> = 2 h) Meperidine Levorphanol Fentanyl Tramadol* <b>Methadone</b> (T <sub>1/2</sub> = 14-40 h)	<b>Buprenorphin</b> Nalbuphine Nalorphine	<b>Naloxone</b> Naltrexone

\* also inhibition of NA and 5-HT uptake

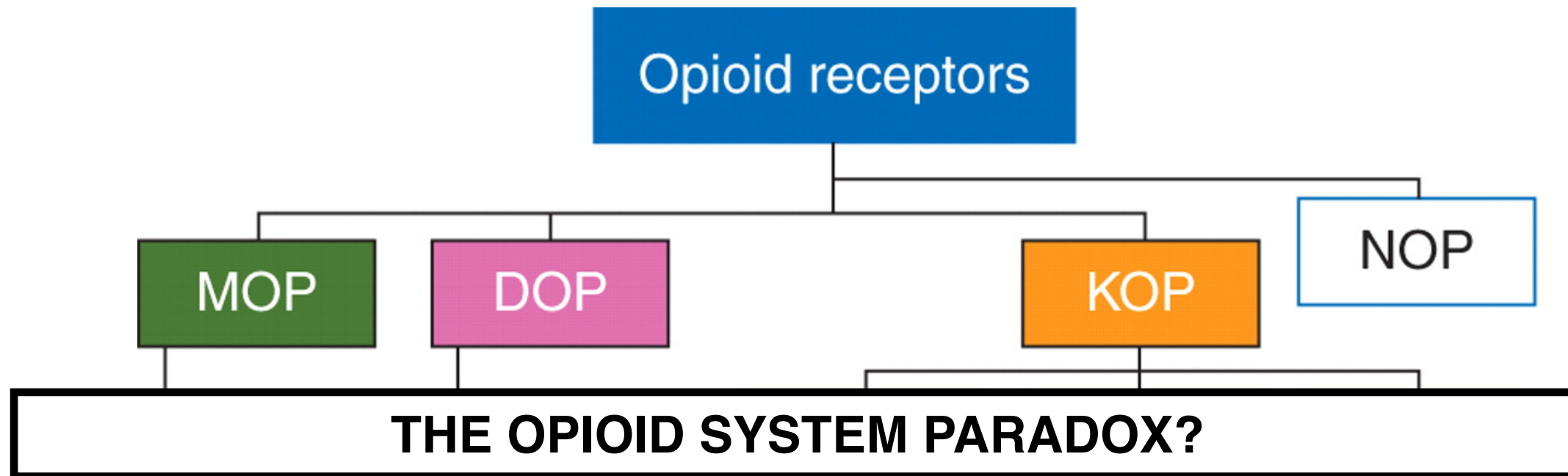
# MECHANISM OF ACTION

All four types of opioid receptors are Gi/o-protein coupled receptors

1. Adenynyl cyclase inhibition
2. N and P/Q voltage-dependent calcium channel inhibition
3. Activation of GIRK (G protein-inhibited rectifying K<sup>+</sup> channels)
4. Activation of MAP kinase pathway



# CLASSIFICATION OF THE OPIOID RECEPTOR FAMILY



A large number of endogenous ligands (at least 11) converge on only a small number of opioid receptors (4 genes)

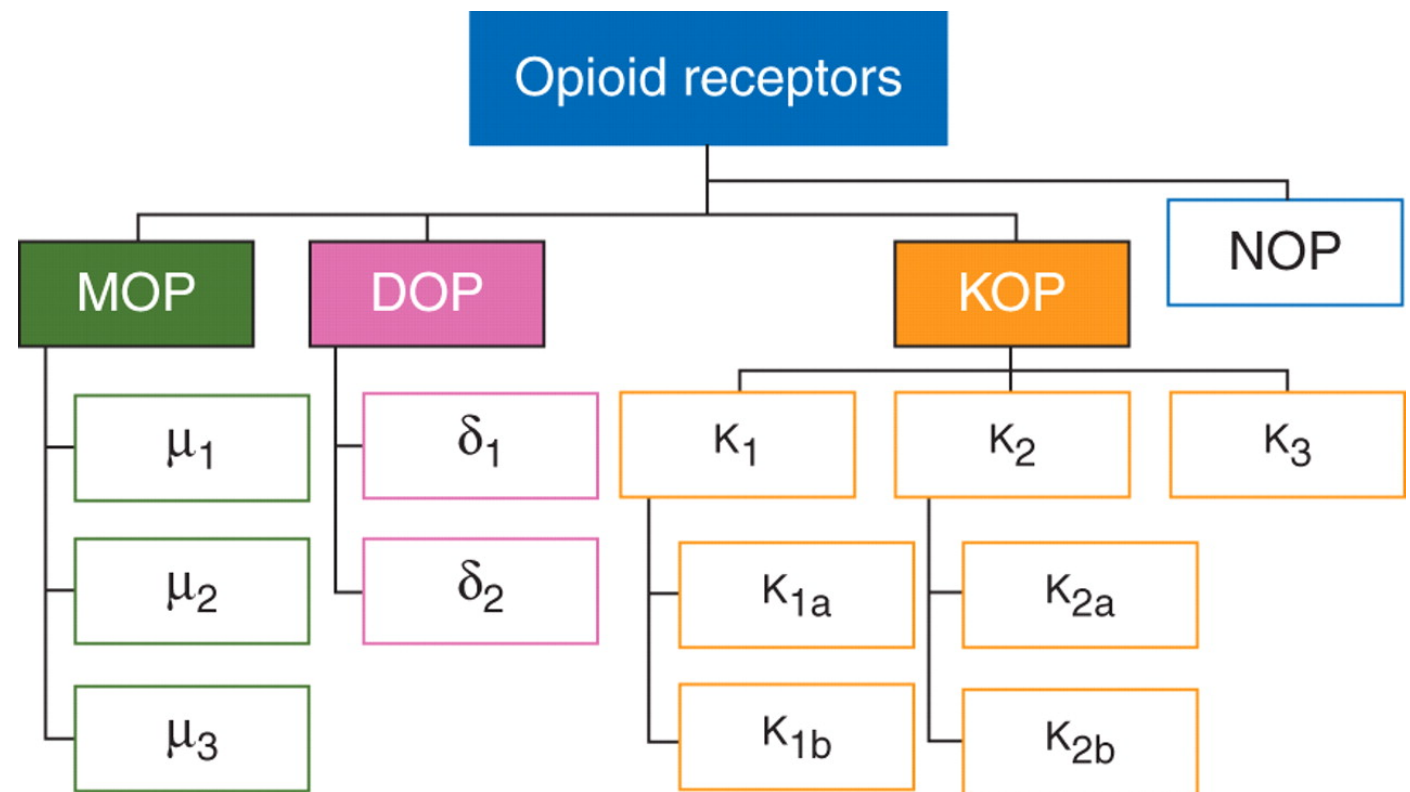
Endogenous ligands display poor selectivity towards opioid receptors (with the exception of Dynorphin A for KOR)

Several pharmacological evidence suggest the existence of multiple receptor subtypes

MOP antagonists (e.g. Naloxonazine) block morphine-induced analgesia but not alter respiratory depression, constipation or itching  
On the other hand, knockout of MOP receptor inhibits all the MOP receptor associated activities

# PHARMACOLOGICAL CLASSIFICATION OF THE OPIOID RECEPTOR FAMILY

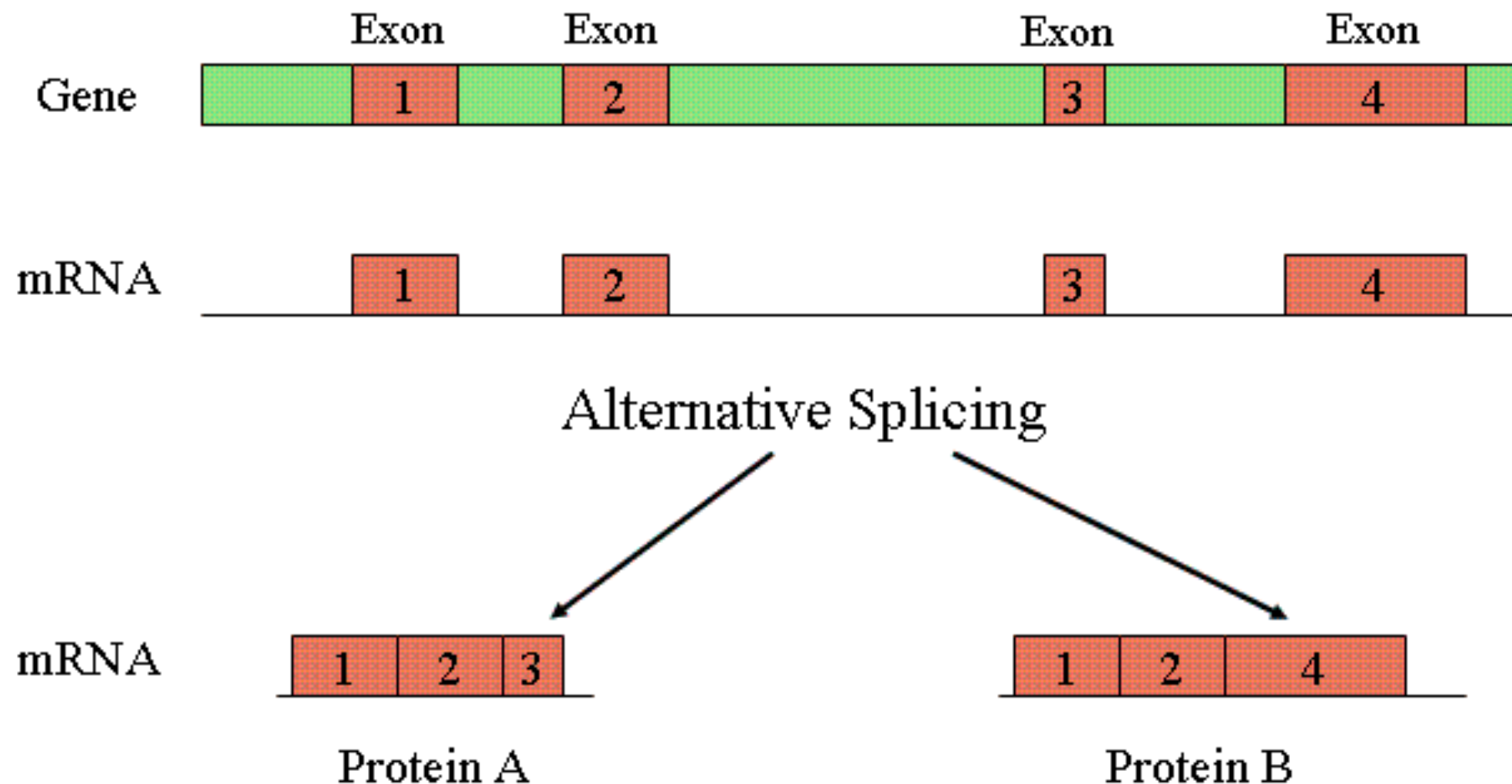
**DO OPIOID  
RECEPTOR  
SUBTYPES EXIST?**



*Three alternative possible mechanisms:*

- Alternative splicing of a common gene product
- Functional selectivity (biased agonism)
- Omo - and/or hetero-dimerization

# 1. ALTERNATIVE SPLICING



Exon 1 is necessary  
for the analgesic  
activity of morphine

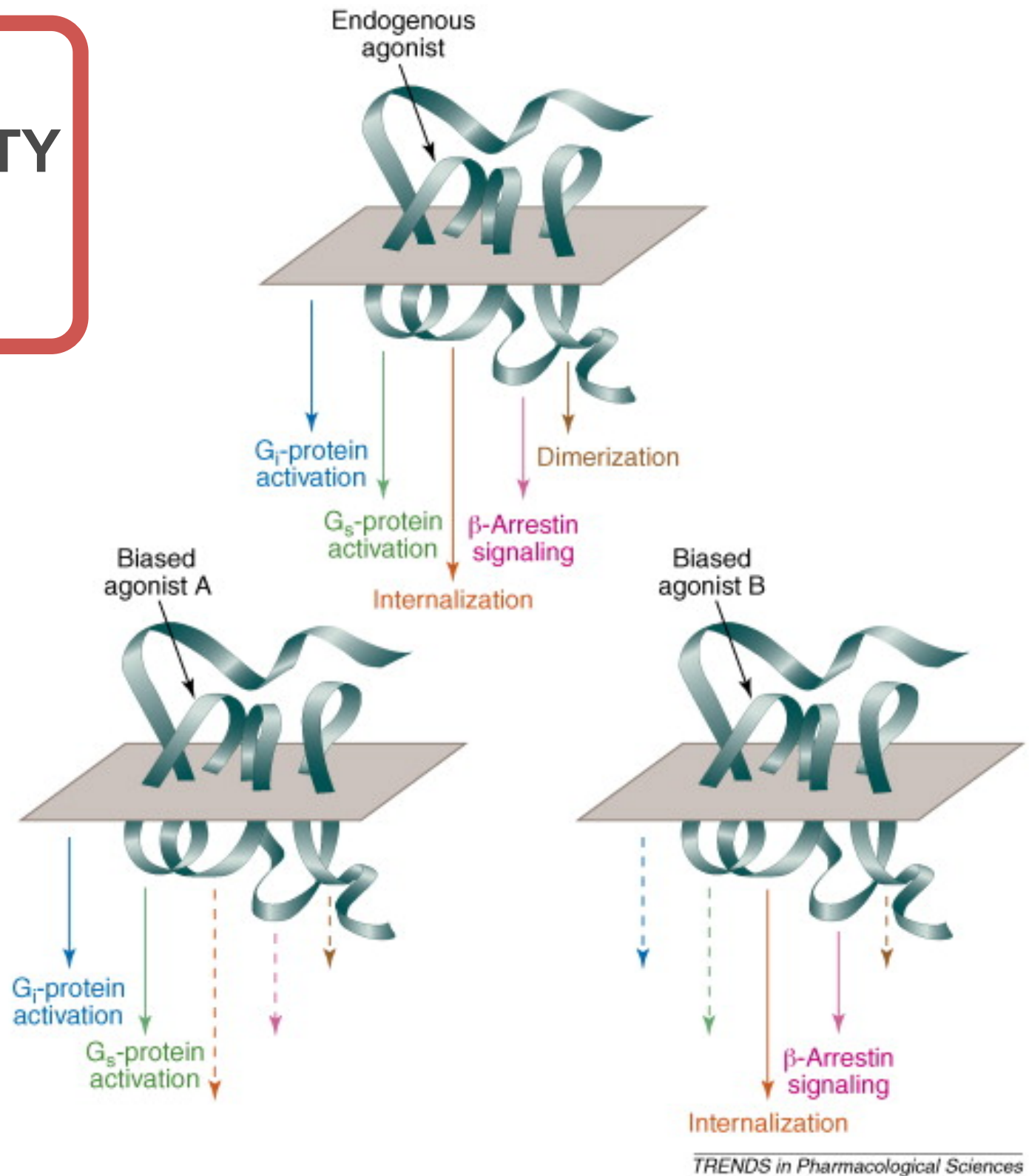


Exon 2 is necessary  
for the analgesic  
activity of fentanyl

PROTEIN ISOFORMS

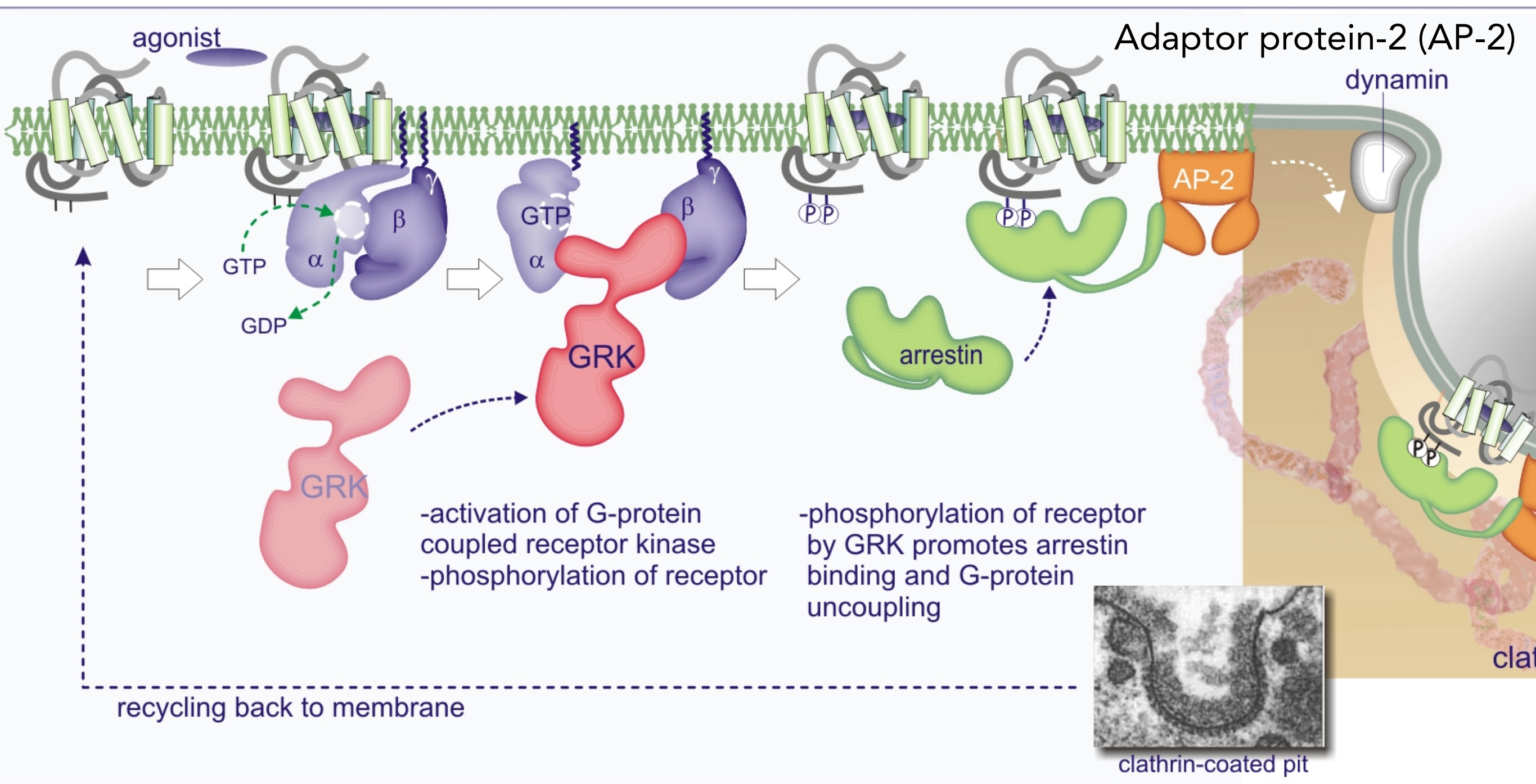


## 2. FUNCTIONAL SELECTIVITY (ligand-selective efficacy)



Different ligands influence which G protein associates with the receptor thus promoting distinct coupling efficiencies (distinct intracellular pathways)

# DESENSITIZATION of GPCRs: molecular mechanisms



# Role of G-protein coupled receptor kinase (GRK) and arrestins

## Turning off the signal: desensitization of GPCRs function

Receptor **desensitization** is a reduced response of a receptor that follows a prolonged exposure to an agonist and it is due to uncoupling of a receptor from G proteins

Desensitization also results from receptor **internalization**, the removal of receptors from a plasma membrane by endocytosis (**downregulation**)

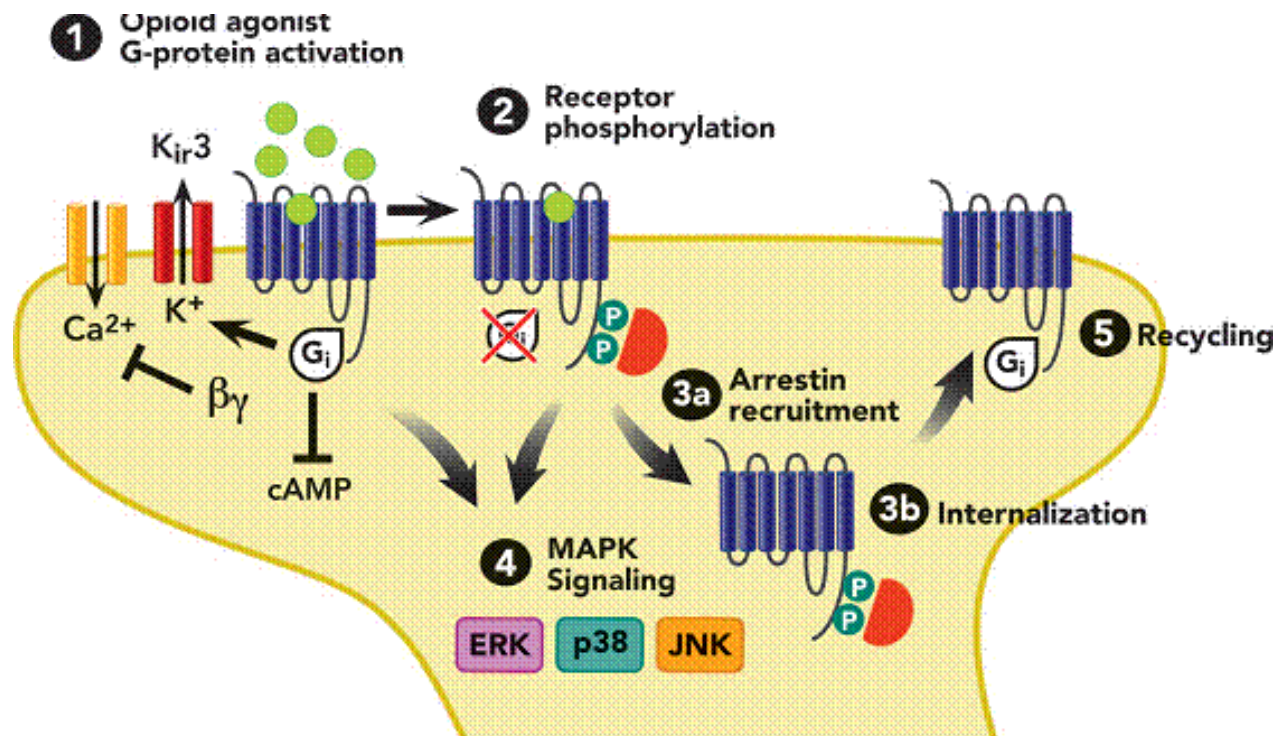
Internalization can be followed by receptor recycling (**resensitization**) or lysosomal degradation

Desensitization can cause (pharmacodynamic) **tolerance**, the need to increase the drug dose to obtain the required effect



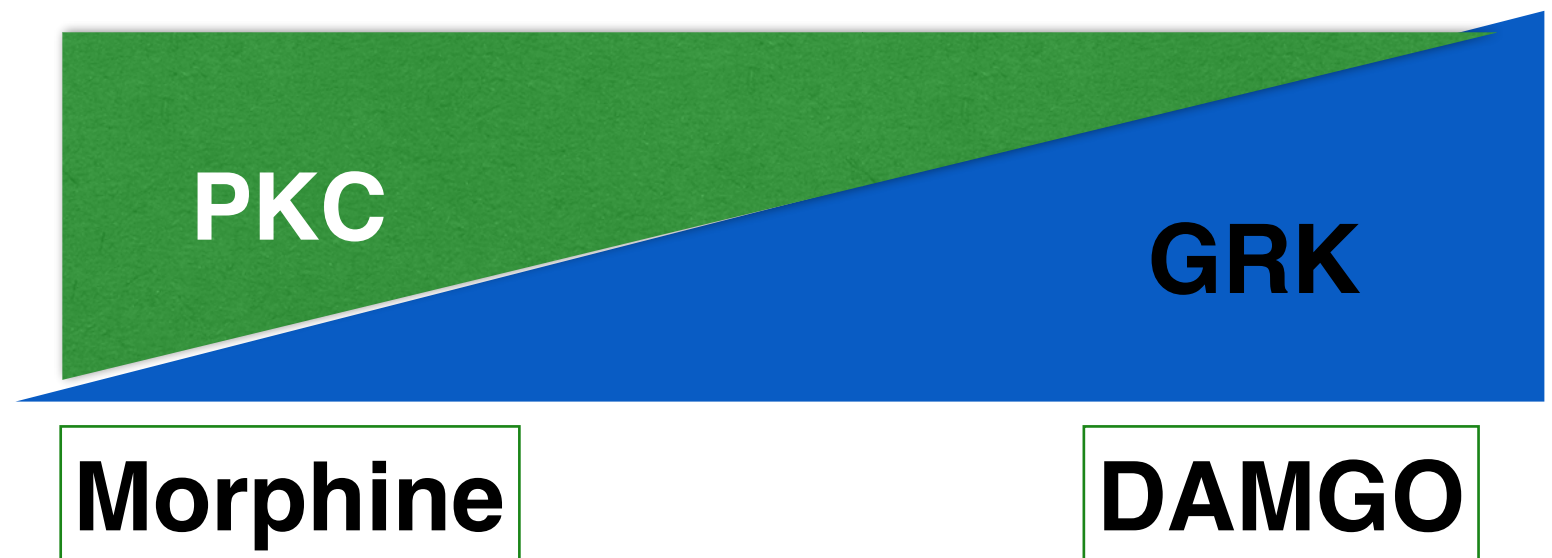
# FUNCTIONAL SELECTIVITY

Selective ligands of opioid receptors can direct the receptor to favor one or more signaling events

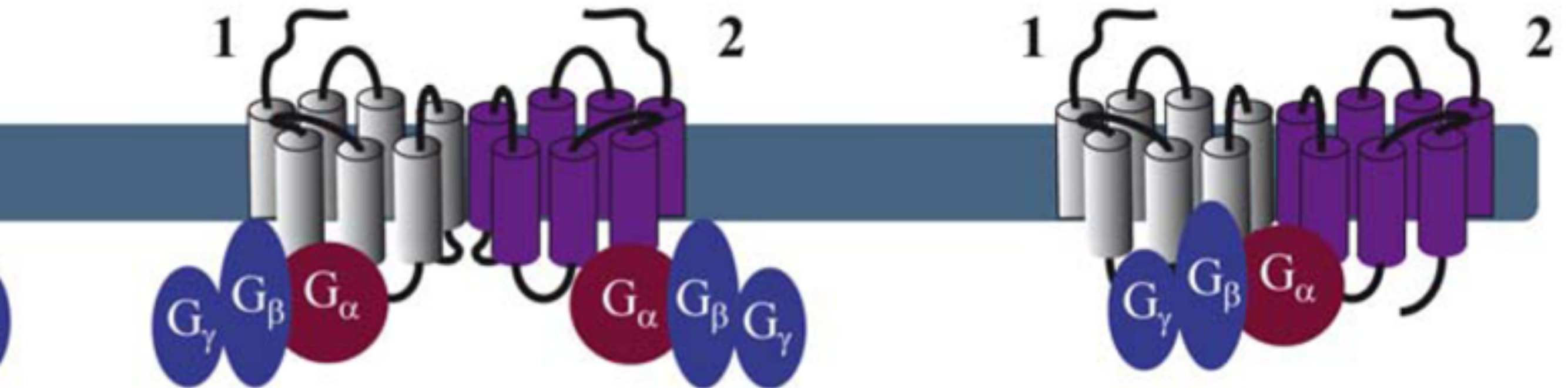


Morphine does not promote MOR receptor internalization and causes tolerance at high degree

In contrast, DAMGO causes robust internalization and low tolerance degree



### 3. GPCR DIMERIZATION



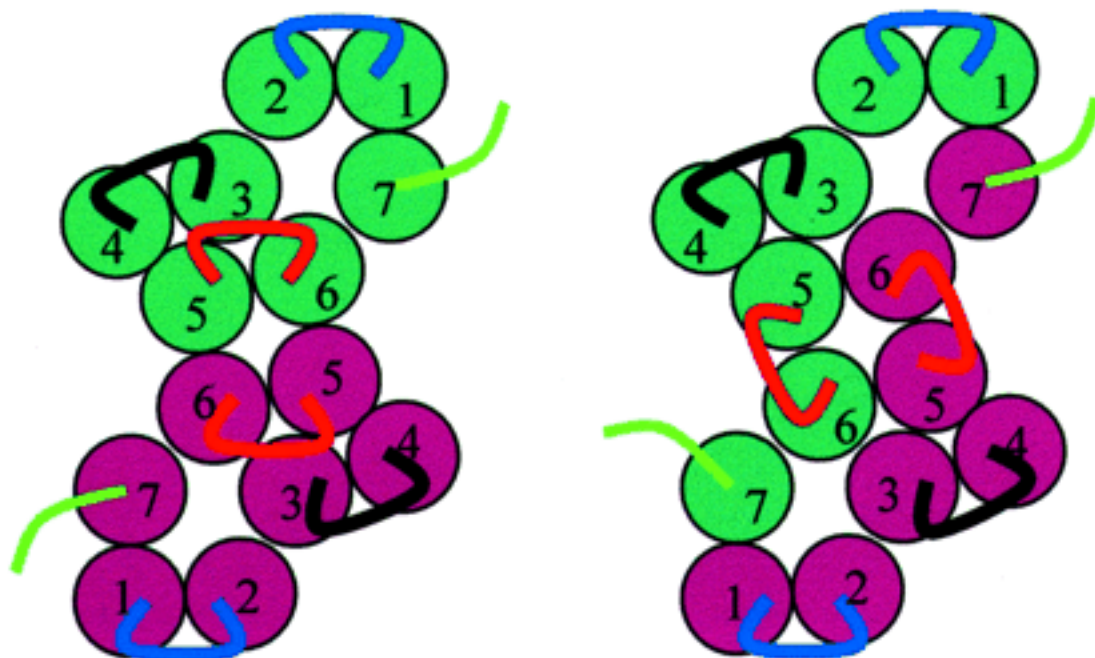
Potential GPCR dimer interfaces

Contact dimers

Domain dimer interfaces

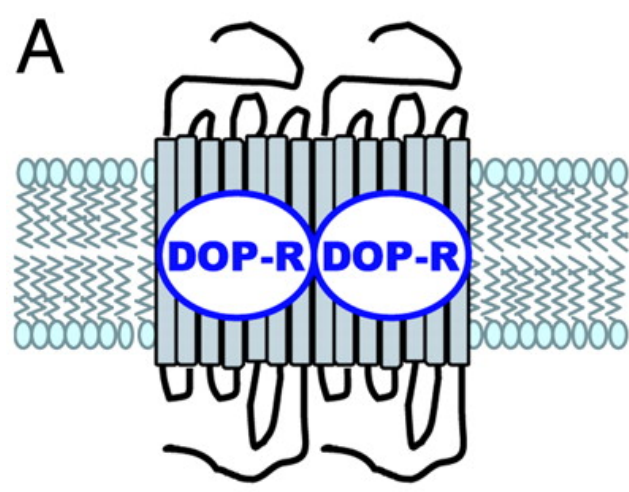
Dimerization affects  
signal transmission and  
desensitization

and can explain the  
differences in efficacy  
and in abuse potential of  
different ligands

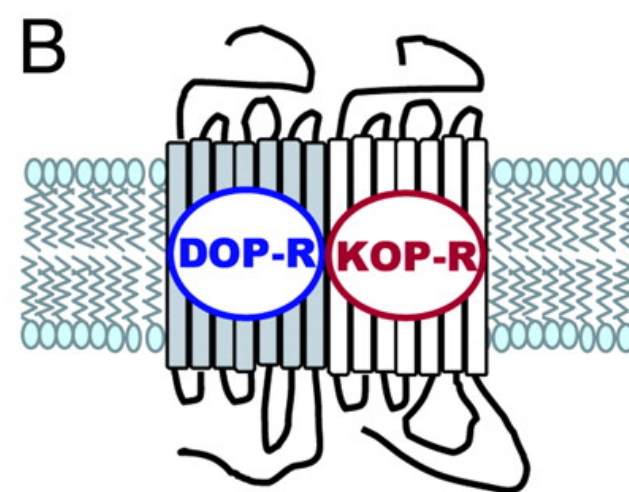




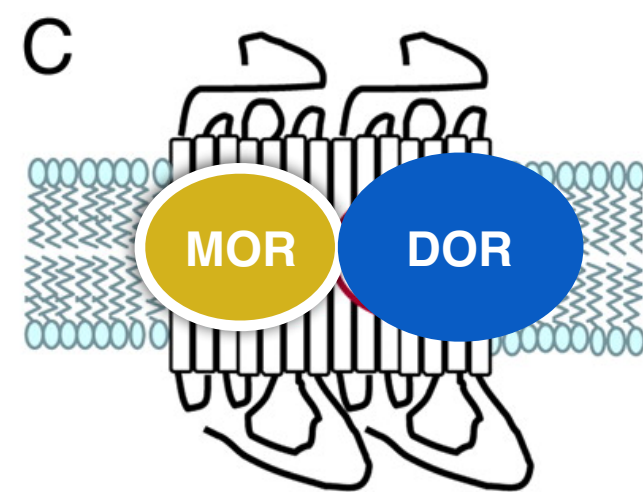
# OMO- and HETERO-DIMERIZATION between Opioid receptor subtypes



No response:  
Agonists stimulation causes  
the dimer dissociation

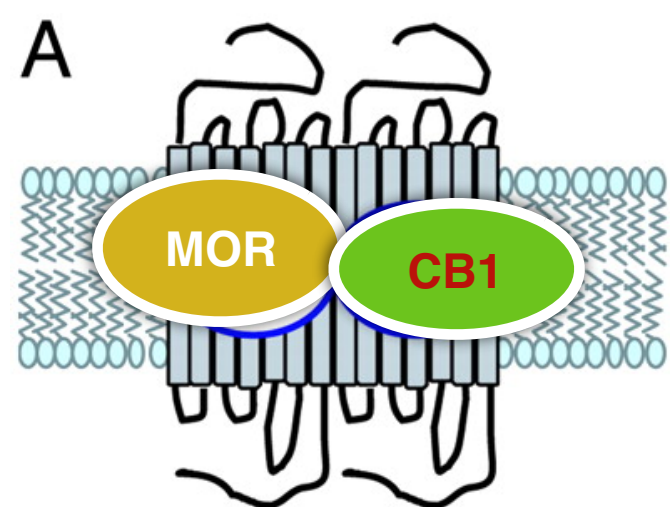


Strong response:  
Reduced internalization

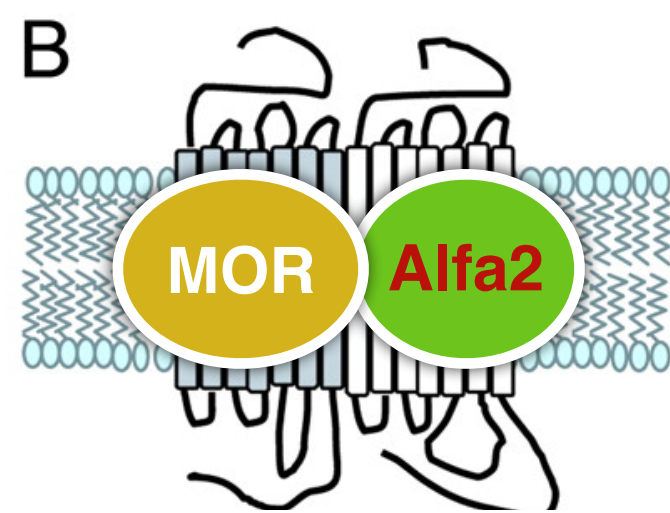


Different signal properties: in  
DOR absence, MOR  
dependent tolerance and  
dependence are reduced

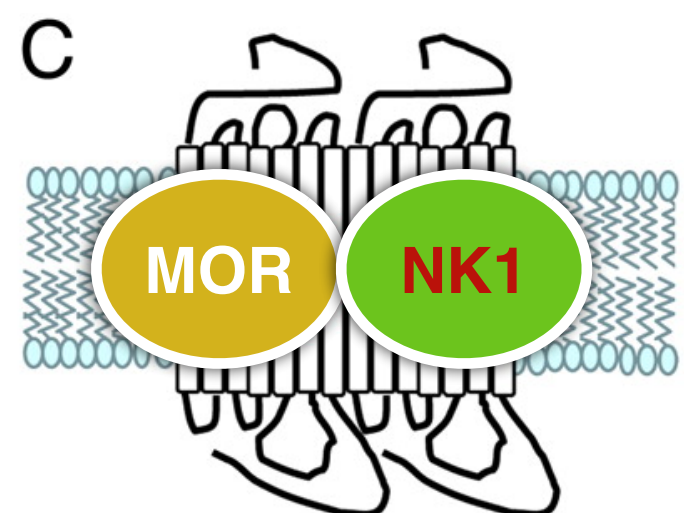
OMO- and HETERO-DIMERIZATION between GPCR



Enhances the potency  
of morphine



Potentiated phosphorylation  
of MAPK induced by  
morphine



SubP causes  
internalization

# FUNCTIONAL EFFECTS ASSOCIATED WITH THE MAIN TYPES OF OPIOID RECEPTOR

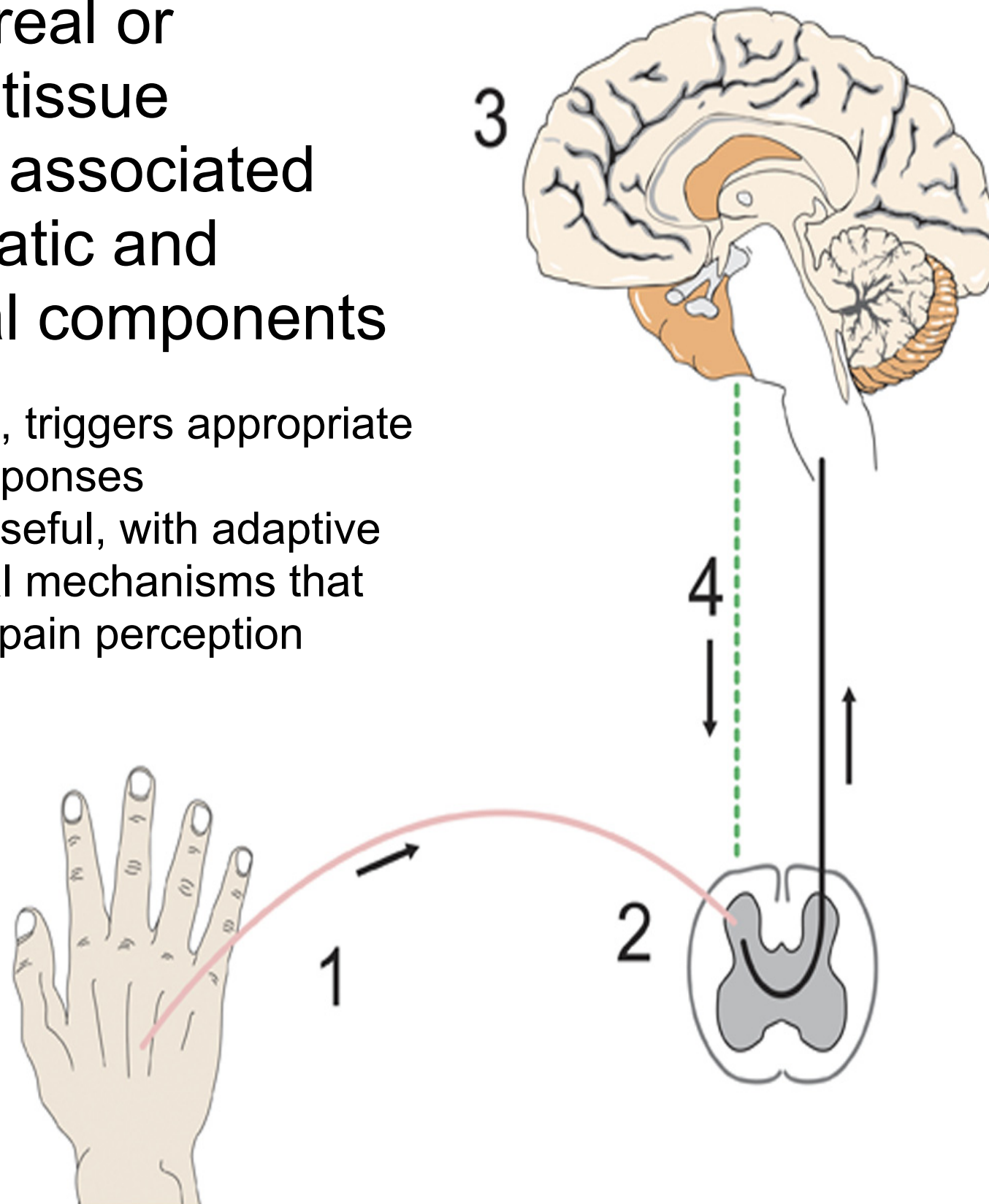
	MOP	DOP	KOP	NOP
Analgesia				
supraspinal	+++	-	-	antag
spinal	++	++	+	++
peripheral	++	-	++	-
Respiratory depression	+++	++	-	-
Pupil constriction	++	-	+	-
Reduced gastrointestinal motility	++	++	+	-
Euphoria	+++	-	-	-
Dysphoria and allucinations	-	-	+++	-
Sedation	++	-	++	-
Tolerance and dependence	+++	-	-	-

# PAIN PATHWAYS

Pain: a sensorial and emotional experience due to a real or potential tissue damage, associated with somatic and emotional components

**Acute:** useful, triggers appropriate protective responses

**Chronic:** unuseful, with adaptive and emotional mechanisms that can increase pain perception



Afferent nerves stimulated by noxious stimuli (1)

activate spinal neurones (2)

that take the information to the supraspinal centers (3)

The SNC modulates the overall response through efferent control systems (4)

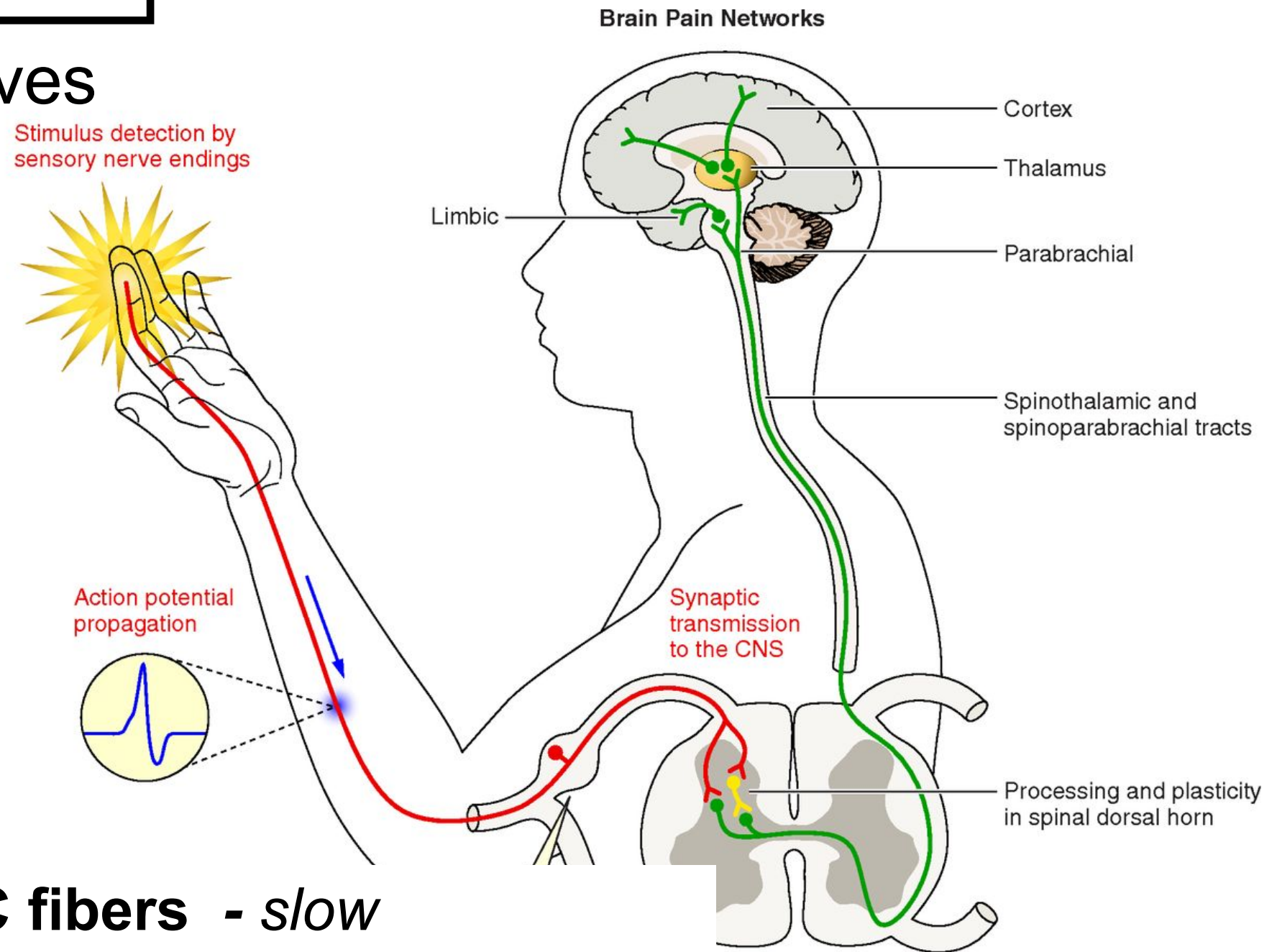


# 1st ORDER NEURONS

Primary sensory nerves  
Nociceptors

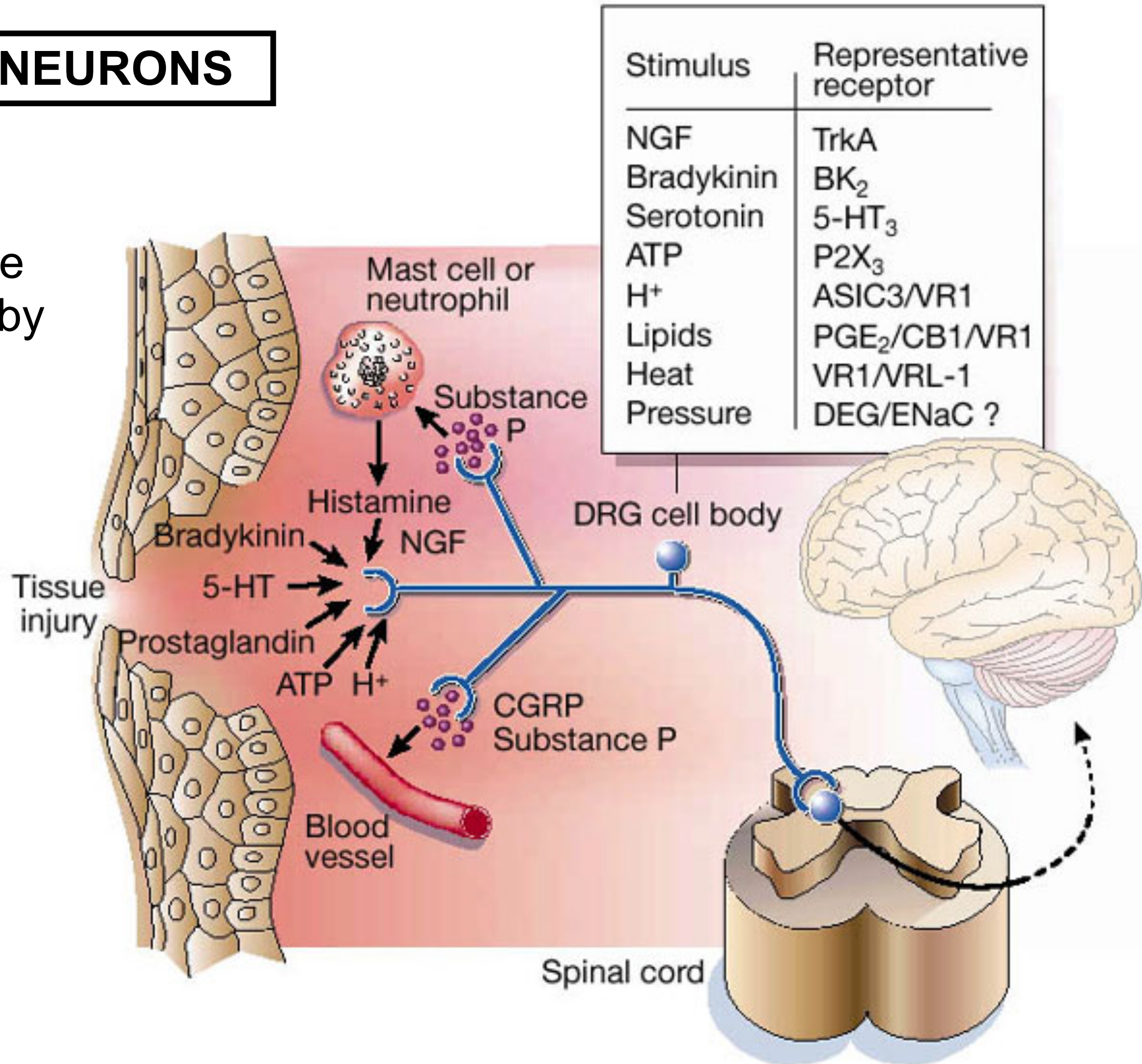
***A $\delta$  fibers*** - fast  
conductance speed:  
small myelinated  
sensitive to  
mechanical noxious  
stimuli  
localized pain

***C fibers*** - slow  
conductance speed:  
small unmyelinated  
sensitive to thermal  
changes, chemicals,  
pressure  
diffuse and strong pain



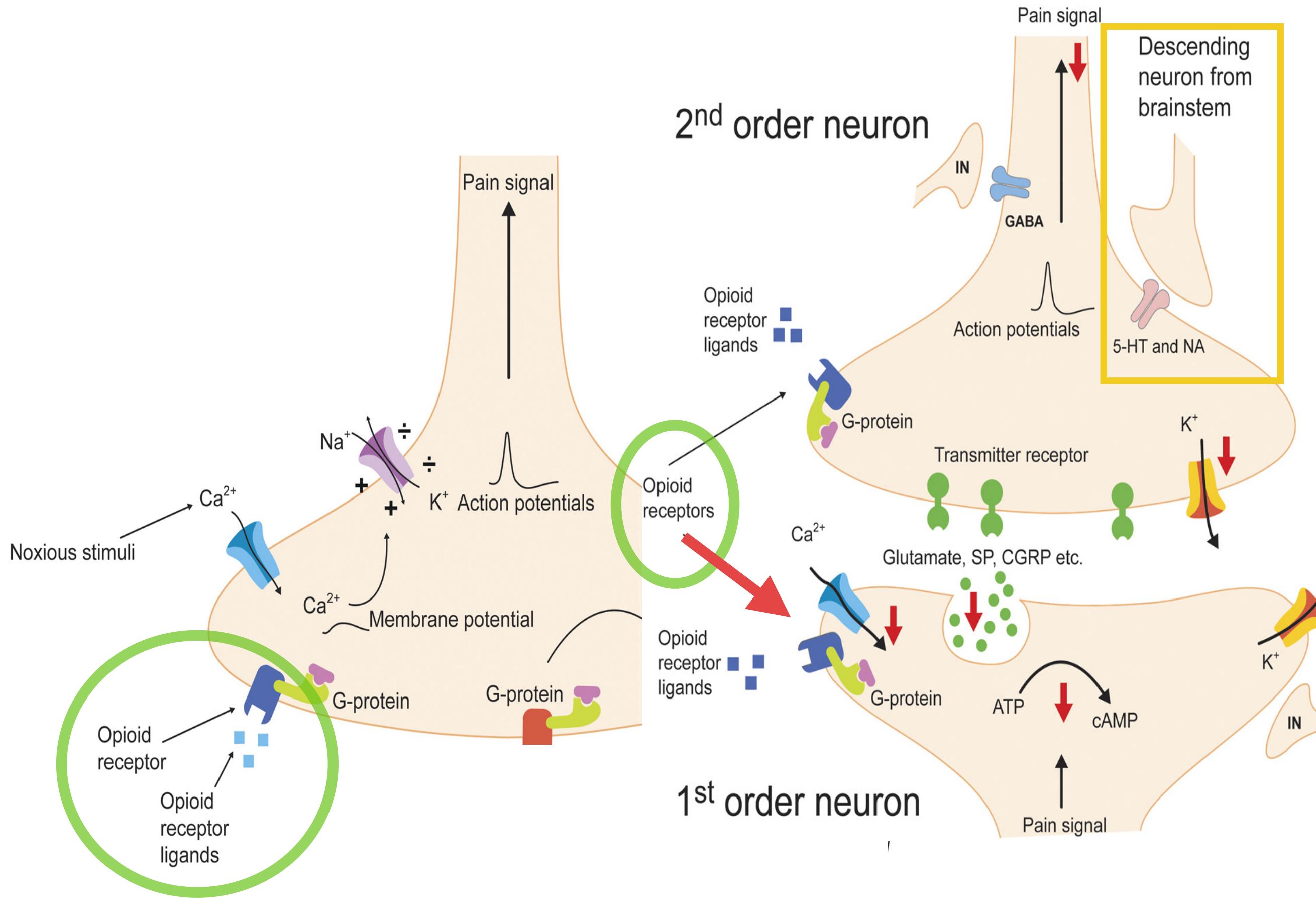
# 1st ORDER NEURONS

Nociceptors are activated also by mediators released after tissue injury



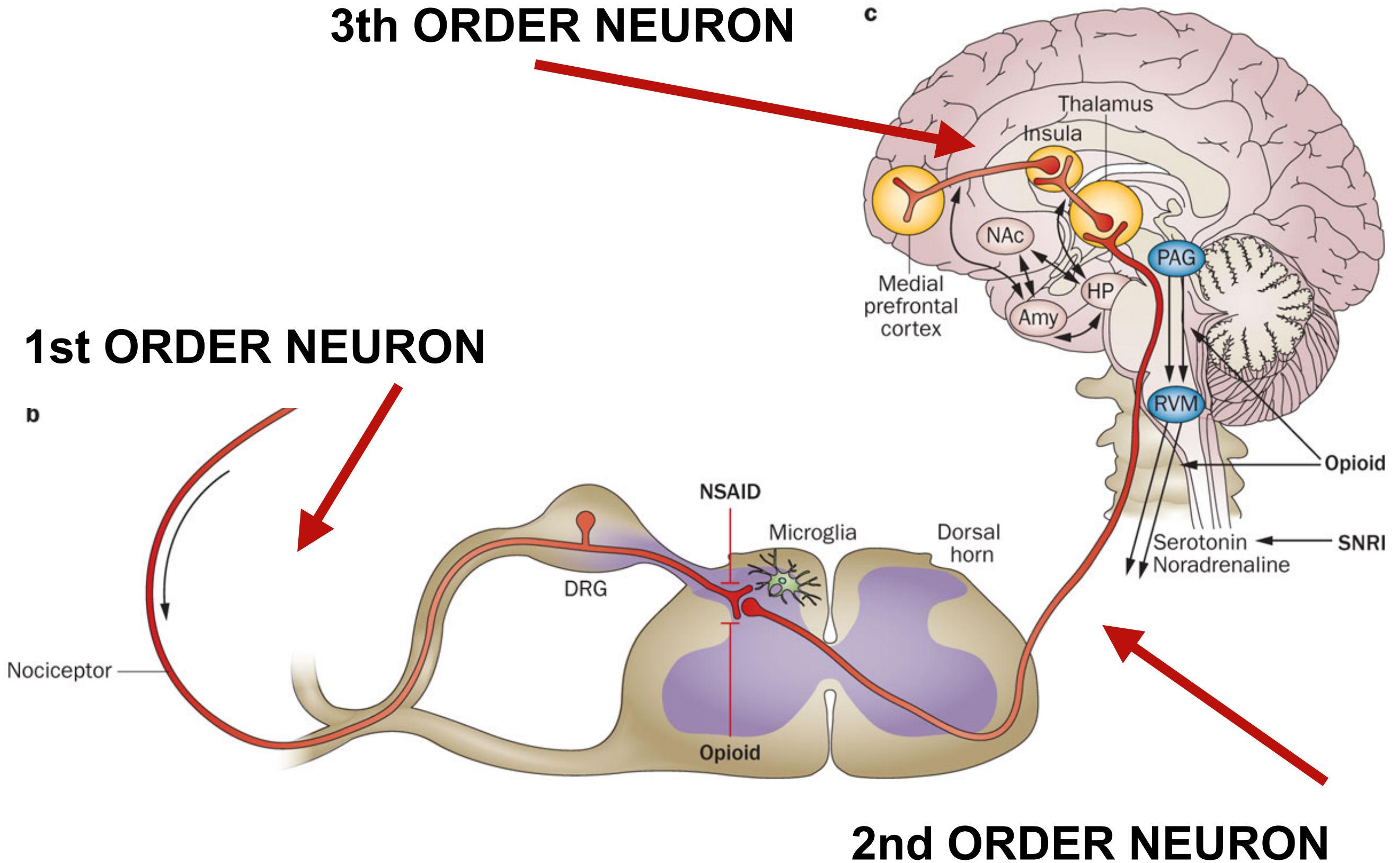


# PAIN PATHWAYS: 1st and 2nd order neurons



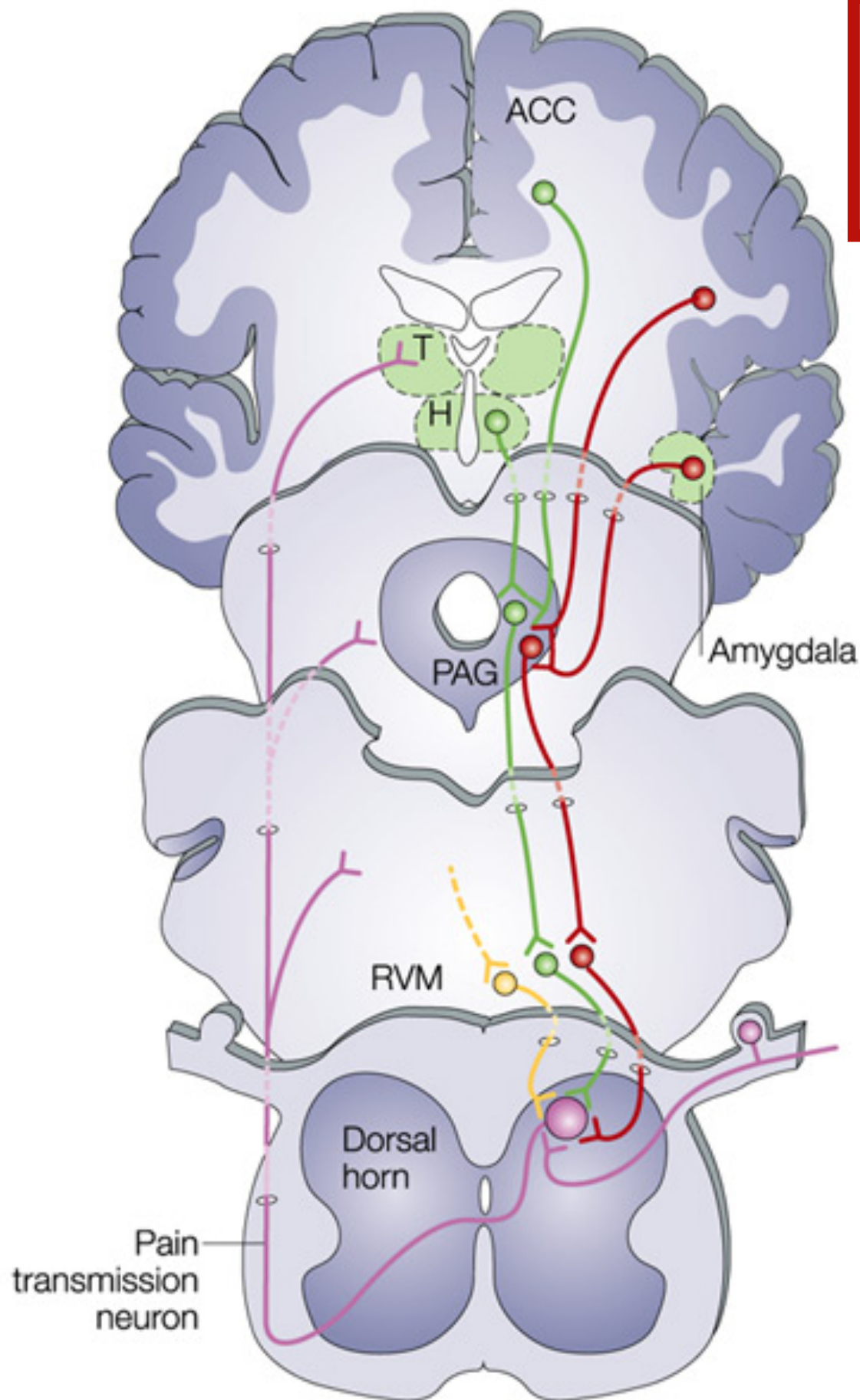
**3th ORDER NEURON**

**1st ORDER NEURON**





# THE TOP-DOWN PATHWAY: the efferent control systems



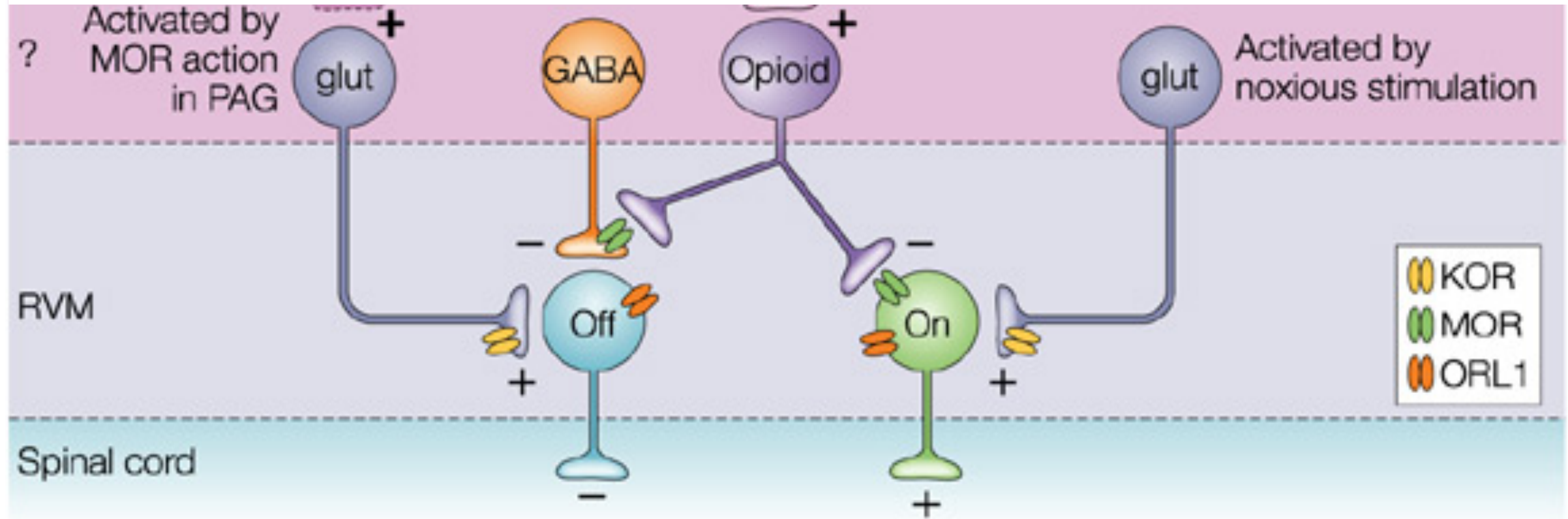
facilitatory (ON cells, red)  
Yes, pain!

inhibitory (OFF cells, green)  
serotonergic (yellow)  
No pain....

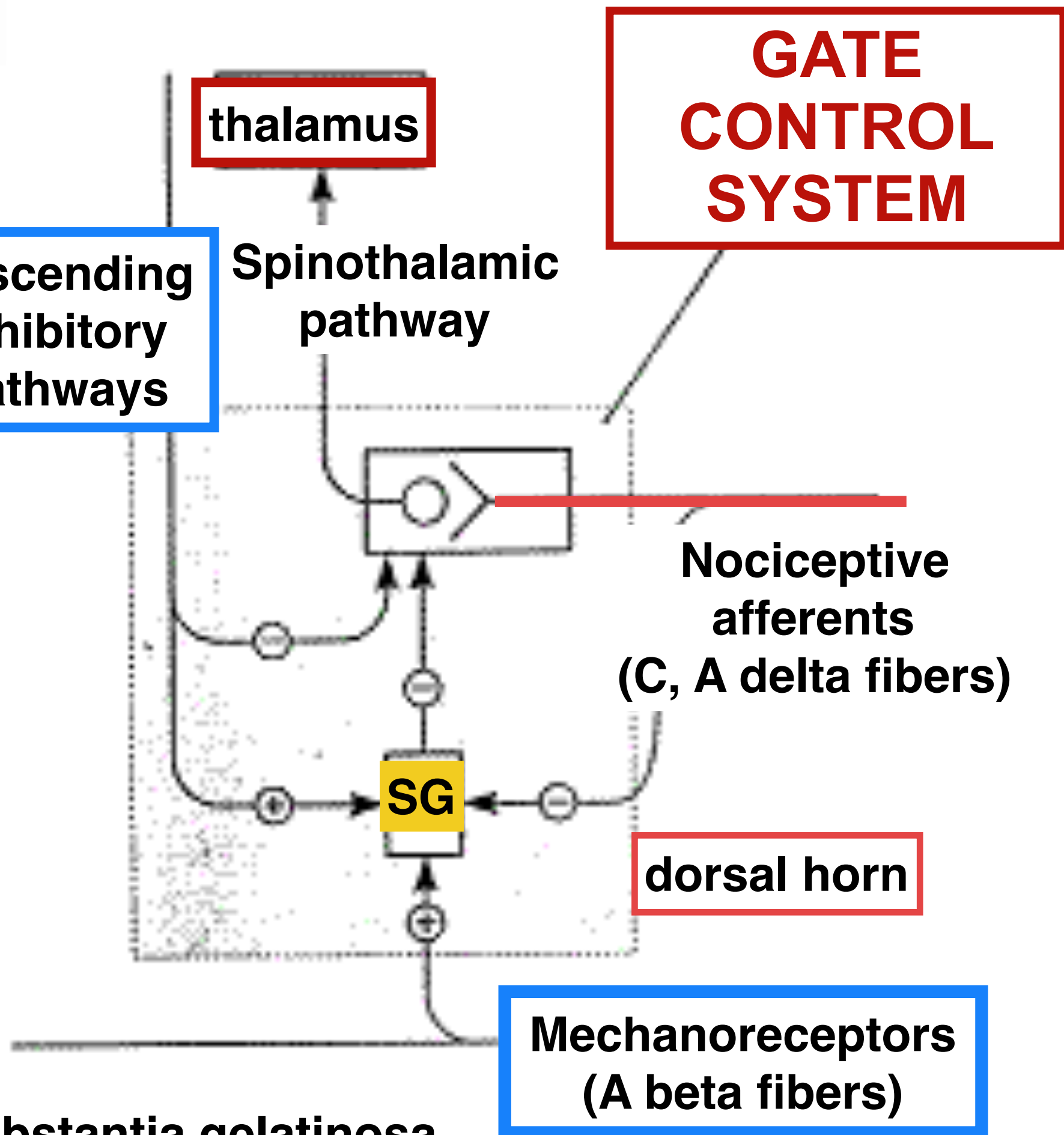
anterior cingulate cortex (ACC)  
hypothalamus (H)  
thalamus (T)  
periaqueductal grey (PAG)  
rostral ventromedial medulla (RVM)

## Within the RVM

MOR agonists produce anti-nociceptive effects by inhibiting ON cells and disinhibiting OFF cells

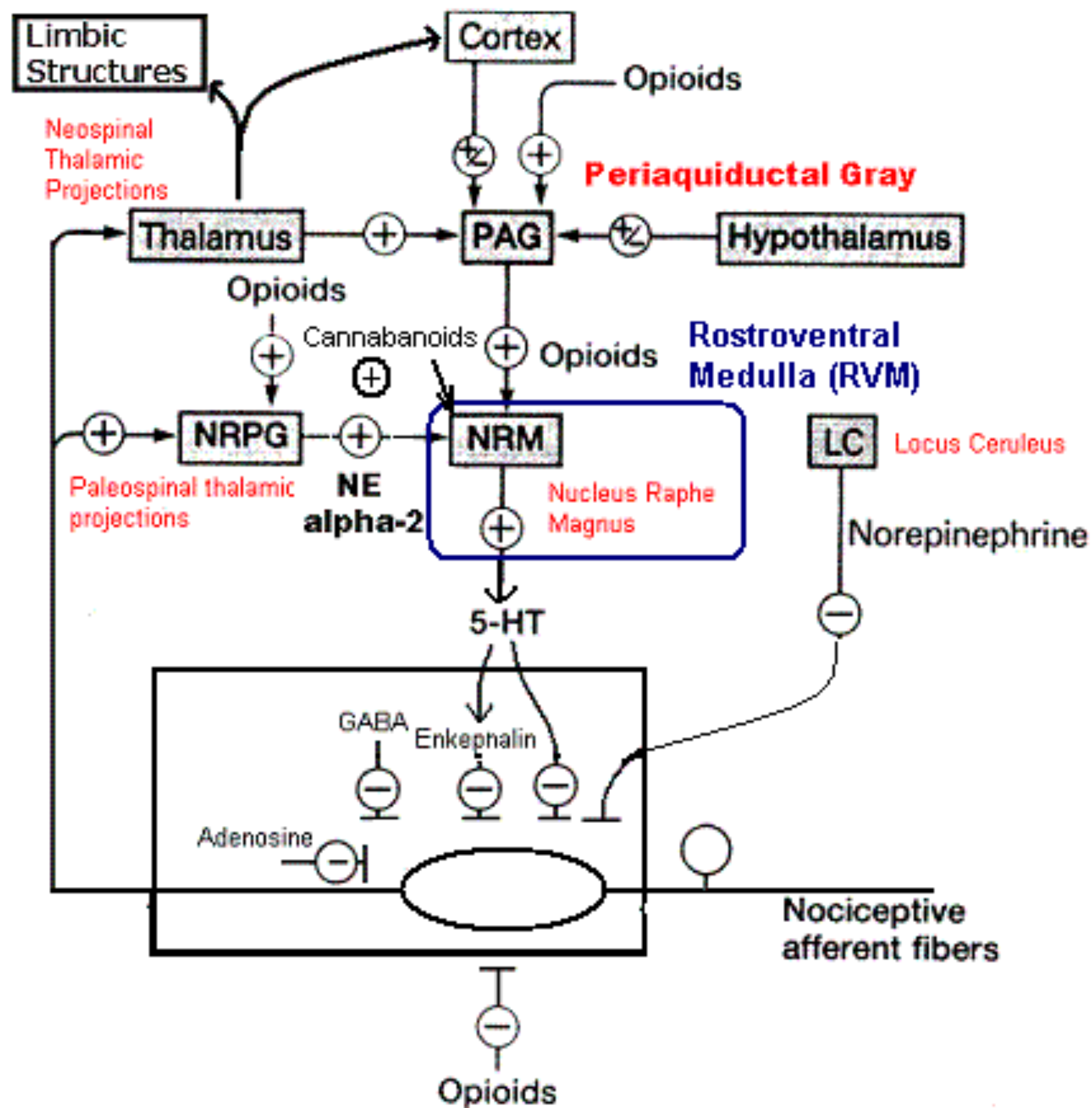


No pain.... Yes, pain!





## Pain pathways: an overview



## Opioid-synthesizing neurons

## Opioid receptors

Cerebral cortex

Thalamus

Nucleus accumbens

Amygdala

Hippocampal formation

Striatum

Hypothalamus

Arcuate nucleus

Periaqueductal gray

Ventral tegmental area

Locus ceruleus

Nucleus ambiguus

Nucleus of the solitary tract

Ventrolateral medullary reticular formation  
(including ventral respiratory groups)

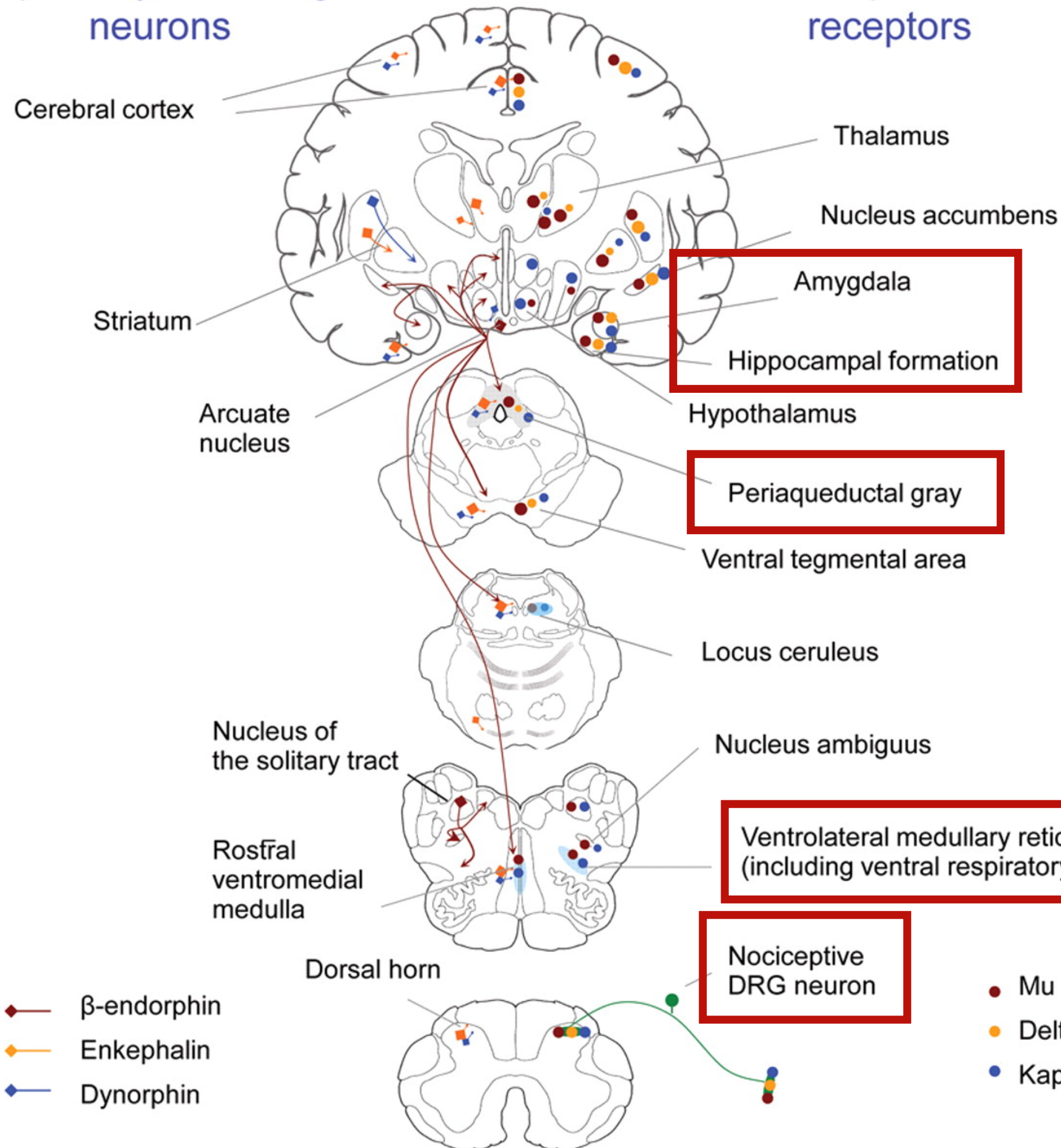
Ros̄tral  
ventromedial  
medulla

Nociceptive  
DRG neuron

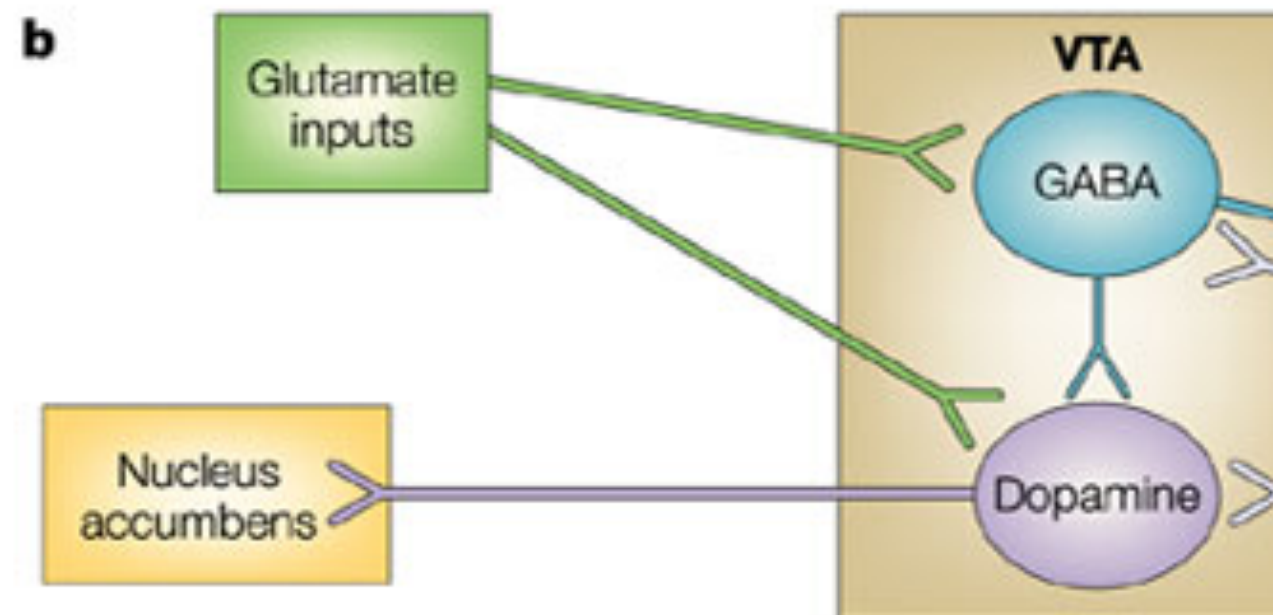
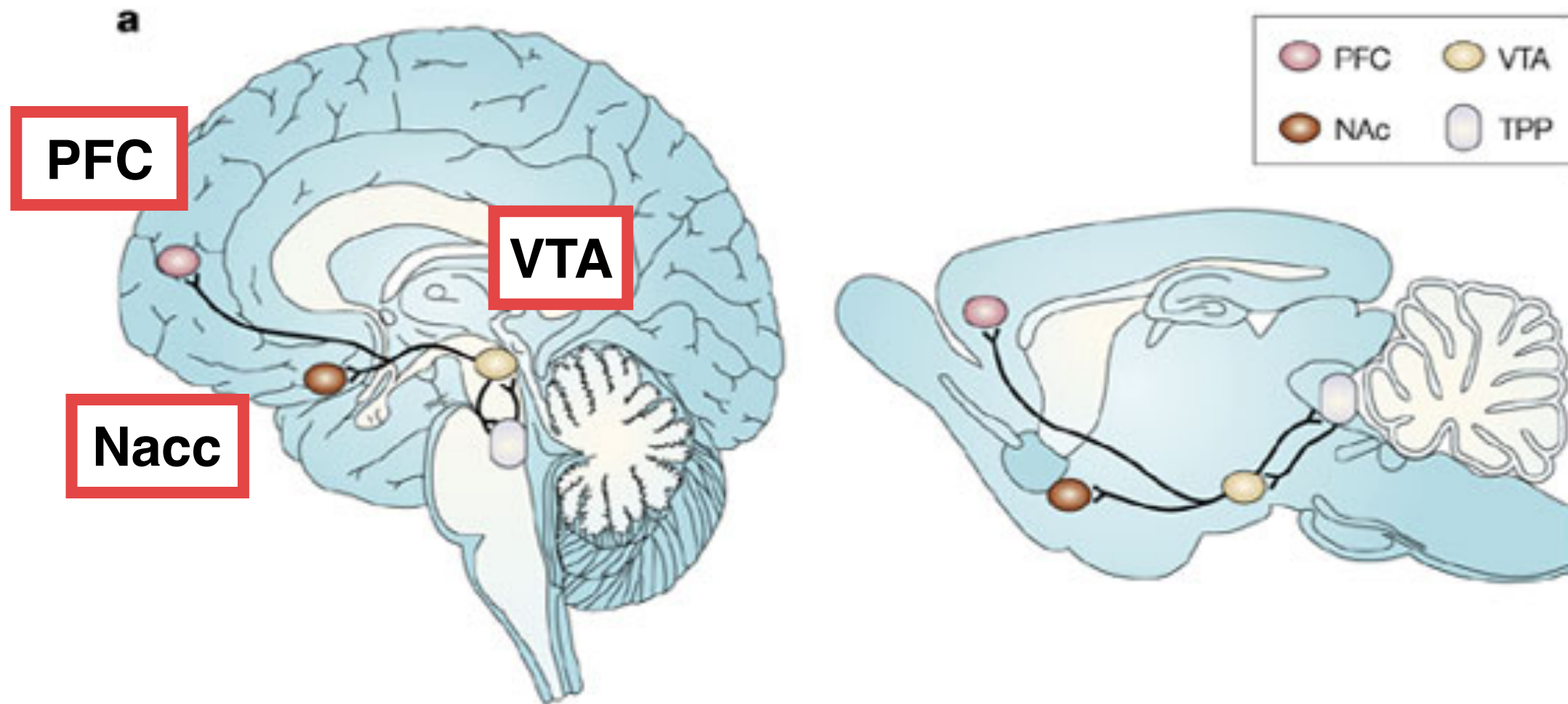
Dorsal horn

- ◆  $\beta$ -endorphin
- ◆ Enkephalin
- ◆ Dynorphin

- Mu
- Delta
- Kappa



# THE MESOLIMBIC DOPAMINE PATHWAY: THE REWARD CIRCUIT AND THE BASIS FOR DRUG ABUSE



VTA ventral tegmental area

Nacc nucleus accumbens

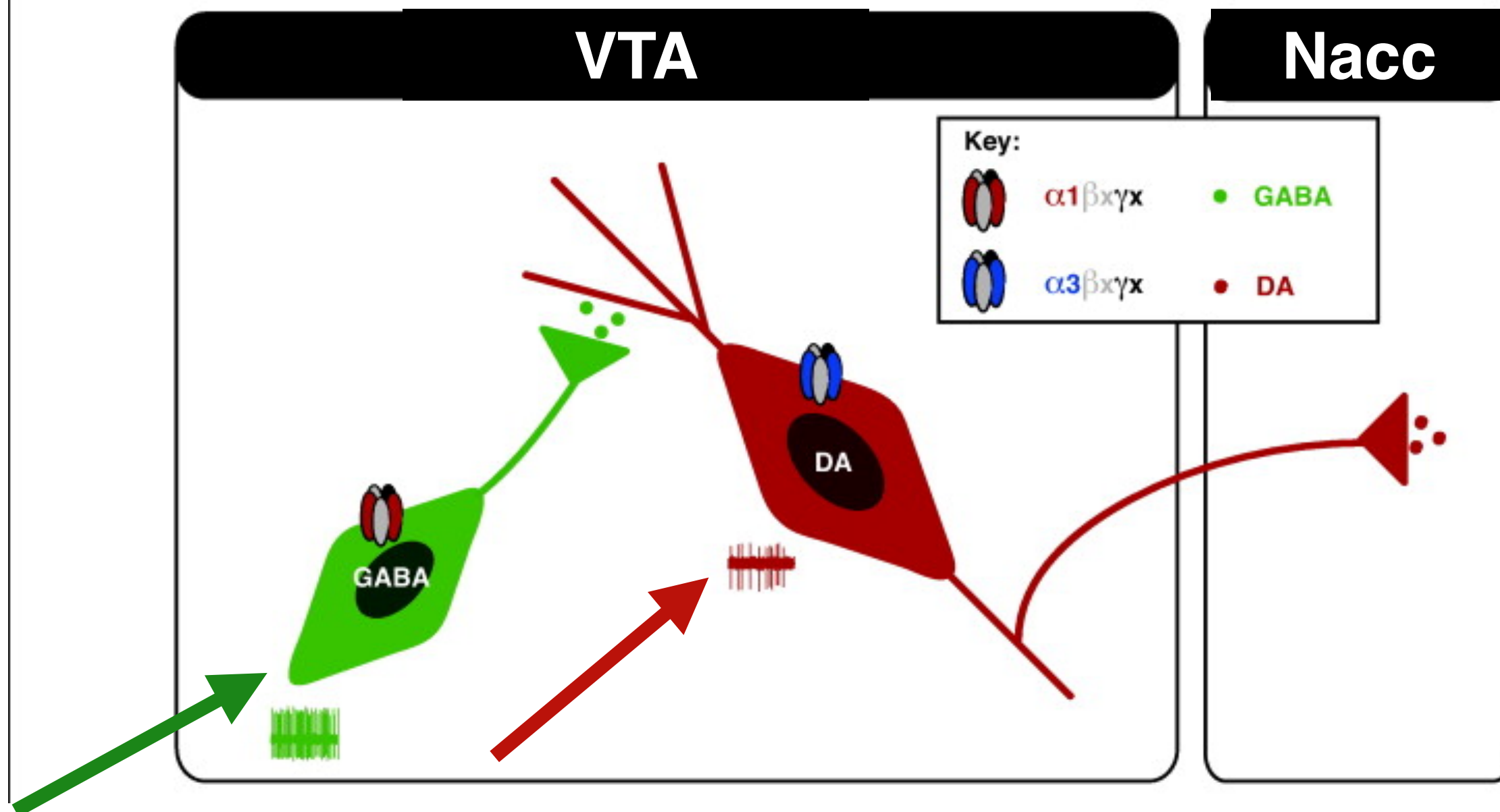
PFC prefrontal cortex



# THE MESOLIMBIC DOPAMINE PATHWAY: THE REWARD CIRCUIT AND THE BASIS FOR DRUG ABUSE

The dopaminergic neurons activity in the VTA is negatively controlled by the basal activity of GABAergic neurones

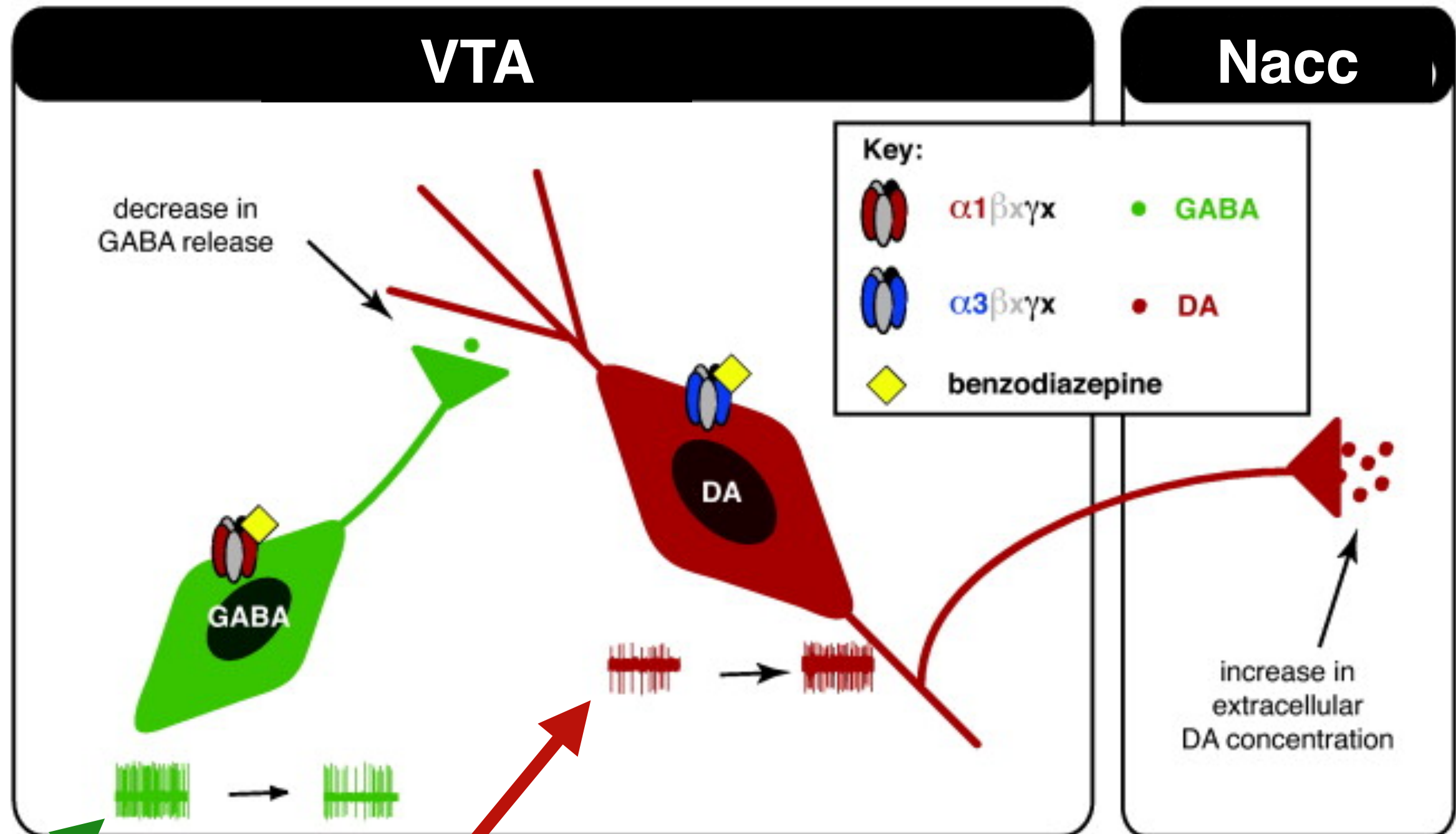
(a) No benzodiazepine:

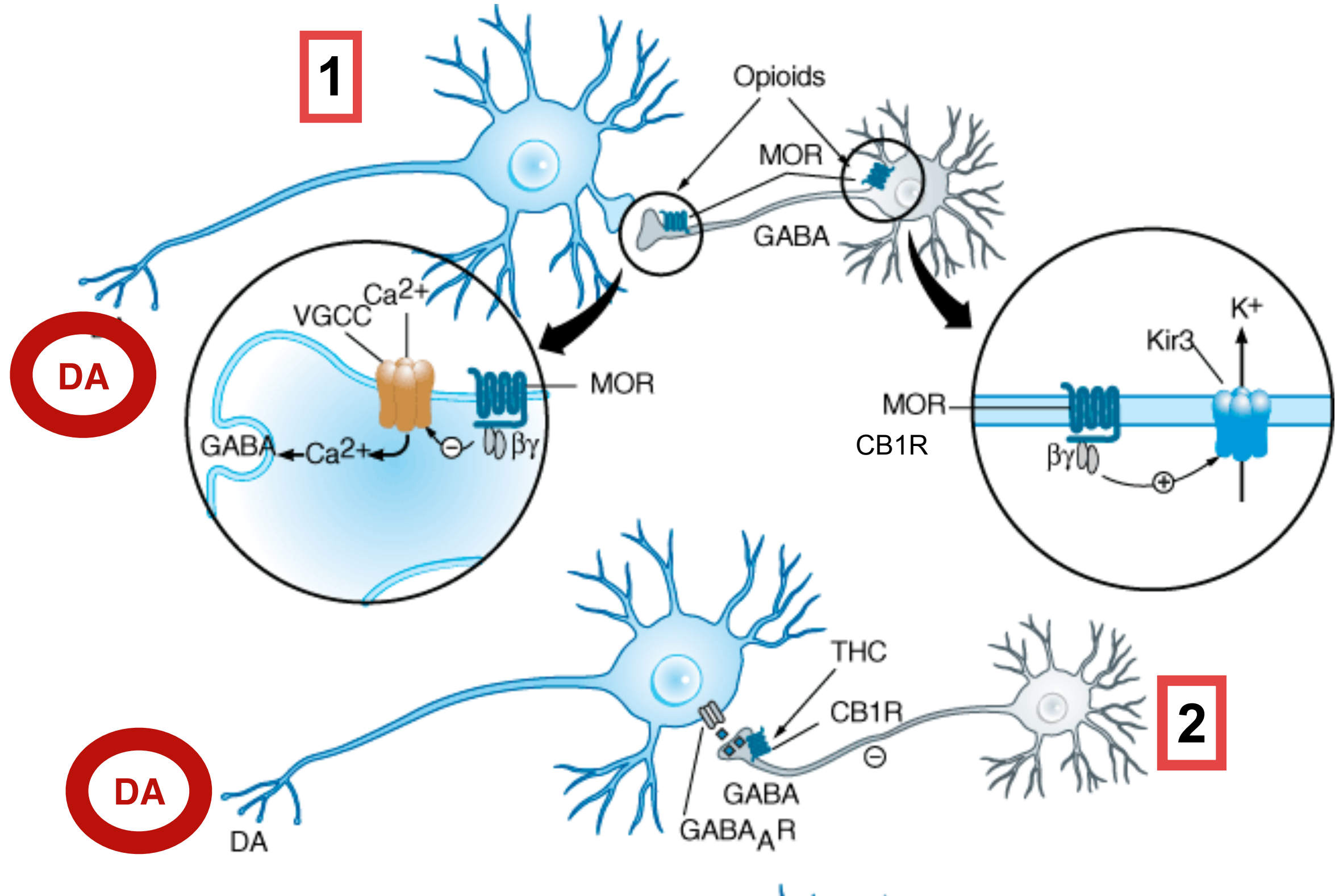




# All drugs that cause disinhibition of dopaminergic neurons are potentially drug of abuse

## Disinhibition mechanism of dopaminergic neurons in the VTA by benzodiazepines





**Disinhibition of dopaminergic neurons in the VTA by opioid (1) and cannabinoid (2) receptors expressed on GABAergic neurons**