

OPIOID ANALGESICS (ANALGESICS – ANODYNES)

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Pain

Definition:

"subjective unpleasant sensory or emotional experience accompanied by real or potential damage of tissues, with motoric and vegetative responses "



Pain – types and classification

A) by duration

B) according to pathophysiology



Pain – types and classification

A) According to length of experience

- 1) acute: sign of and disease, danger or damage to organism...
- 2) chronic: more than 3 months / unusually long for a given disease or disorder



Pain – types and classification

A) According to length of experience

- 1) acute: physiological sensory perception,
 - tissue damage,
 - mobilizes defensive forces of the organism in order to remove the inducing cause of the pain
- 2) chronic: pathological,

pain may persists even after the removal of the causes → difficult to determine whether the pain arose as a result of persistent pathological activity in the nerve endings in the periphery, or is the source of the CNS



Pain – types and classification B) According to pathophysiology

- nociceptive irritation of nociceptors
 Therapy: "analgesic ladder" according WHO (see below; not used for aggressive procedure in the treatment for cancer or breakthrough pain)
- 2) neurological and neuropathic pain
 Therapy: antidepressants and **anticonvulsants**(in combination with opioids or some muscule relaxants; neuroprotective vitamins thiamine; antimigraine drugs from the group of the so-called triptans; antipsychotics = neuroleptics)



Pain – types and classification B) According to pathophysiology

3) psychogenic pain somatization, hypochondric and somatoform disorder

Therapy: psychopharmac drugs (antidepressants – TCA, SSRI, anxiolytics, antipsychotics)



■ neuralgia

sharp, paroxysmal pain, affects peripheral or cranial nerves (often the trigeminus, facialis) → after traumatological damage, compression, viral infects (herpetic), metabolic (DM)

■ pain in the chronic compression of peripheral nerves and nerve roots

hernia of the intervertebral discs, compression of the nerve in the spinal cord → pain + paresthesia, pain acquires a hot character



■ischemic pain

due to disorders of blood circulation in the myocardium, smooth or skeletal muscle

■ migraine

migraine is characterized by attacks of pulsating, mostly unilateral headache lasting typically 4-72 hours with nausea, possible vomiting, photofobia and phonofobia, suffering from 12% of the adult population



■ phantom pain

surgically or traumatically removed parts of the human body, most commonly the lower limb or other parts of the body (ablation of the breast, as well as after the removal of the visceral organs - the colon);

apply pathophysiological influences peripheral, central and psychogenic;

■ breakthrough pain

sudden, transient, mostly short-term worsening of pain in patients who have well-controlled baseline pain;

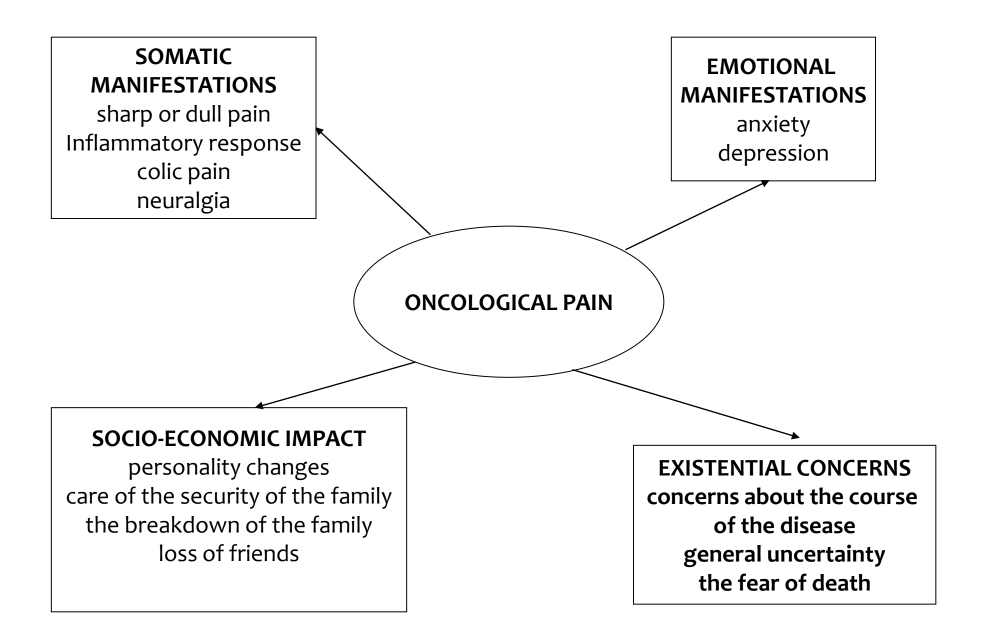
usually in patients treated with opioids for cancer diagnosis; typically in progression of cancer



delivery pain

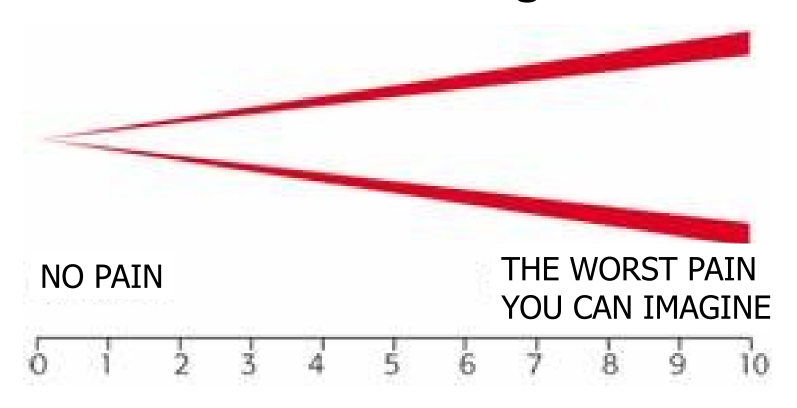
- belongs to the strongest pain reliever, nevertheless, that before the birth are rising thresholds for somatic and visceral pain
- the tissue is developed by excessive pressure, they are strongly being pushed and lacerations occur
- tissues are under influence of bradykinin, H⁺, K⁺, histamine and serotonin
- induction of stress → ↑ cortisol, epinephrine, norepinephrine, dopamine → somatic and psychological reactions





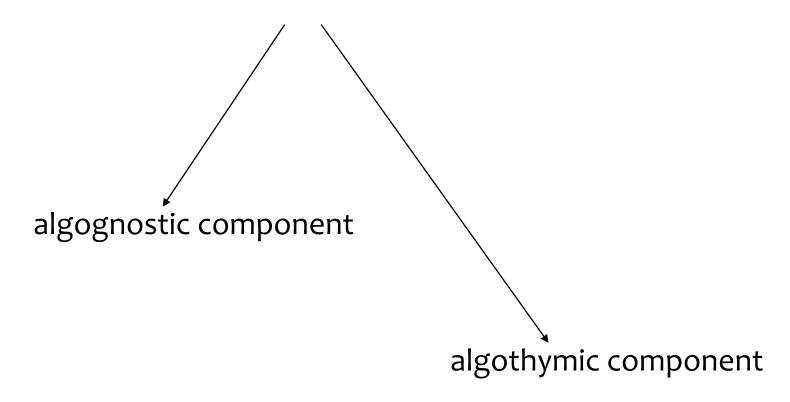


Diagnostics of Pain VAS: Visual analogue scale





Process of pain perception





Pain – causes and mechanism

tissue damage □ production of prostaglandines and other substances □ effects on the free nerve endings □ transduction of signal up to the brain neurons □ PAIN
Mediators of pain
(act on the nociceptors = pain receptors)
"algogenic substances"
bradykinin ——— + ——— ↑PGE (mediators of inflammation), increase sensitivity of nociceptors
histamine
acetylcholine
substance P (pain)



Pain – causes and mechanism

Endogenous pain suppressing (analgesic) substances:

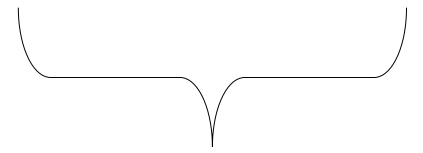
endorphins enkephalins dynorphins



Pain transduction – 3 neuron's tract

MAIN PAIN PATHWAYS

tractus spinothalamicus vs. tractus spinoreticulothalamicus (spinothalamic tract) (spinoreticulothalamic tract)



the tracks leading from the spinal cord (spinal ganglia) to specific areas of the brain (finally to the cerebral cortex) information about pain is received and processed



Pain transduction – 3 neuron's tract

- → spinothalamic tract 3 neuron`s phylogenetically younger pathway
- sharp, well localized pain
- → spinoretikulothalamic tract phylogenetically older polysynaptic system, impulses are transmitted to the higher centres through short axonal pathways
- dull, poorly localized pain
- vegetative response: blood pressure change, tachypnea, mydriasis, diaphoresis, increased muscle tone,...



Pharmacological modulation of pain

Analgesics – anodynes (opioids)

Non-opioid analgesics (analgesics – antipyretics □ NSAIDs)

Local anaesthetics

General anaesthetics

Adjuvant therapy (antidepressants, neuroleptics - antipsychotics, antiepileptics - anticonvulsants, antimigrenics, central/peripheral myorelaxants, corticoids, bisphosphonates, caffeine...)



Pharmacological modulation of pain

<u>Analgesics</u> – suppress perception of pain (increase the pain threshold) <u>selectively</u> without influencing perception of other stimuli

analgesics – anodynes (opioids)

act on spinal and supraspinal level, cause effects on somatic and visceral pain, strong effects on consciousness, act substantially more strongly than non-opioid analgesics

non-opioid analgesics

mostly peripheral effects (some have central effects!), effects on inflammation, weaker effects in general, no effects on visceral pain, no addiction



Analgesics - anodynes

blocking transmission of pain signals between cells of the CNS (in the spinal cord, brain), as well as <u>endogenous</u> <u>opioids</u>:

endorphins, enkephalins, dynorphins

→ binding to opioid receptors (agonists)

Opiates

substances similar structurally to morphine with analgesic effect (natural origin, currently produced synthetically)

Opioids

+ synthetic, semisynthetic and endogenous opioid peptides + exogenous opioid analgesics



Opioid receptors - μ κ δ (σ)

- G-protein coupled
- the interaction of the opioid with receptor → G- protein inhibition → reduction of neurotransmitter release + inhibition of neuronal activity
- •adenylylcyclase inhibition, facilitation of K⁺ channels opening postsynaptically, inhibition of Ca²⁺ channels opening presynaptically

- µ
- **■** K
- **•** δ
- **■**(σ)

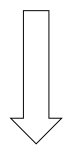


Opioid receptors - μ κ δ (σ)

- µ supraspinal analgesia, euphoria, sedation, miosis, breath depression, addiction, GIT effects
- κ spinal + peripheral analgesia, sedation, dysphoria, miosis, GIT effects, (somatic addiction)
- δ spinal analgesia, breath depression, inhibition of GIT motility
- [σ] dysphoric effect, psychotomimetic effect
 [(hallucinations, perception disturbances), anxiety]



Opioid receptors - μ κ δ (σ)





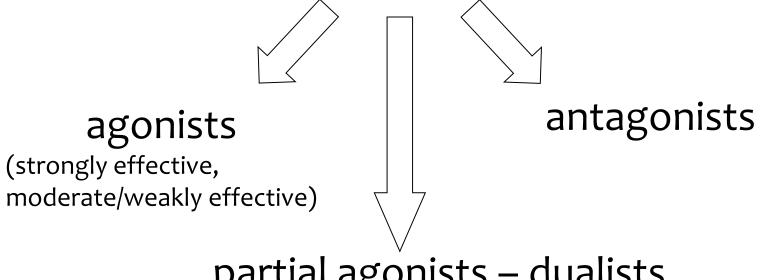
FOR ANALGESIC EFFECT IS CRUCIAL ESPECIALLY ACTIVATION OF RECEPTORS:

μ – supraspinal analgesia

к – spinal + peripheral analgesia



Pharmacological influence on opioid receptors



partial agonists – dualists mixed agonists - antagonists

Atypical opioids



CENTRAL: analgesic

suppression of respiratory center

sedation (+-)

suppression of anxiety

euphoria/dysphoria

antitussive effect

nausea and vomiting

↑ tendency to convulsions/cramps

miosis

↑secretion of ADH,

↓ GnRH, corticotropine, FSH, LH, ACTH, cortisol,

testosterone)



TOLERANCE!!!

(to all effects of opioids except constipation and miosis!)

ADDICTION!!!



PERIPHERAL:

- decrease intestinal motility, slowdown propulsion of GIT content
- increase muscle tone of GIT and urinary bladder
- increase sphincter tone of gall bladder and urinary bladder
- constriction of pyloric sphincter, delayed gastric emptying
- vasodilation, orthostatic hypotension
- histaminoliberation
- inhibition of ciliated epithelium



PERIPHERAL:

- urological tract increase tone of renal pelvis, ureter, m. detrusor and sphincter of bladder...urine retention, especially in postoperative conditions
- uterus \(\psi \) tone and motility, may prolong labor



Pharmacokinetics of analgesics - anodynes

ABSORPTION parenteral

oral (,,first pass effect" !!!)

perrectal

transdermal

sublingual

transmucous (nasal)

DISTRIBUTION

parenchymatous organs

muscles

adipose tissue (lipophilic drugs → e.g. fentanyl)

pass well across BBB → brain (fentanyl,

heroin,..)



Pharmacokinetics of analgesics - anodynes

BIOTRANSFORMATION primarily in liver polar metabolites

- inactive metabolites
- <u>active metabolites</u> (codeine, tramadol, morphine...)

EXCRETION • kidneys - urine

• liver - bile



Opioid agonists

morphine

- 10 % of opium content, together with codeine, thebaine
 - + other phenanthrene alkaloids
- isolated in 1803 (Sertürner)
- high affinity to μ receptors selective μ agonism



Opioid agonists morphine

- > see above effects
- application routes:
- > orally (also p.o. with sustained release)
- > parenterally (i.v., i.m., s.c., epidural,...)
- > perrectally

Indications: chronic cancer pain, pain after surgery, injuries, (pain during acute myocardial infarction → today, given preference to other opioids)



Other strong opioid analgesics

methadone

- less sedation and euphoria than in morphine
- ↑ bioavailibility after oral administration, ↑t ½
- acts on opioid + NMDA receptors
- Use: addiction treatment (heroin) → 2 benefits
 - → change from injection application to oral administration
 - → ↑t ½ decreases plasmatic fluctuation of methadone → less withdrawal symptoms



Other strong opioid analgesics

heroin (= diacetylmorphine)

- not used in clinical medicine (in Czech Republic)
 (but in Great Britain can be therapeutically used!)
- causes severe addiction; abused!
- heroin belongs to the most health and personality devastating substance!!!



Other strong opioid analgesics

fentanyl, sufentanil, remifentanil,...

- pass well across HEB († concentrations in CNS)
- strong, short analgesia (fentanyl 100 x more potent
 - than morphine, sufentanil 1000 x more potent than morphine)
- strong respiratory depression!, ↓ emetogenic
 potency, CAVE → can cause muscle rigidity
- risk of serotonin syndrome in combination with
 5-HTergic drugs



Other strong opioid analgesics

fentanyl, sufentanil, remifentanil,...

- Indications, use:
- in anaesthesiology → neuroleptanalgesia

(= neuroleptic (AP) + opioid)

→ analgosedation (e.g. opioid + BZD)

- therapy of strong pain acute myocardial infarction, cancer pain,...
- fentanyl in TTS (\uparrow duration of action can be used in chronic cancer pain), transmucous (can be used in breakthrough pain)





Other strong opioid analgesics

piritramid

- less respiratory depression than morphine
- less emetogenic potency
- usually well tolerated parenteral administration
- <u>Use:</u> therapy of acute strong pain, e.g. after surgery (PCA), acute myocardial infarction, pain after injuries,...

Others: oxycodone, hydromorfone (not registered in Czech Rep.), **oxymorphone** (not registered in Czech Rep.)

CAVE: all strong opioids are prescribed to forms with blue band ("opiate forms"), very strict accounted and subjected to the rules for handling with narcotics and psychotropic drugs and their precursors!!!



Other strong opioid analgesics pethidine (=meperidine)

- ↓suppression of respiratory center than morphine
- Janalgesic potency than morphine (5-10 x weaker)
- metabolite norpethidine is proconvulsive and causes hallucinations
- administration orally and parenterally

Indications: cancer pain, pain after injuries, pain during acute myocardial infarction, pain after surgery, premedication before general anaesthesia...today not often used (high risk of abuse, hallucinations!)



Opioid agonists (moderate and weak potent)

codeine

- 10 % mtb. to morphine
- antitussive effect in subanalg. doses
- analgesic effect in combinations (paracetamol, ASA)
 analgesic potency: codeine 50mg ~ ASA 1g
- † risk of addiction than strong opioids
- CAVE ↑ risk of addiction of combined (compositive) analgesics
- causes obstipation
- not used in children!!!



Opioid agonists (moderate and weak potent)

dihydrocodeine

- suitable in pains combined with cough (this co-incidency is not necessary for dihydrocodeine indication)
- in Czech Rep. dihydrocodeine in sustained release drug form
 (effect 12 h) → indication for chronic moderate and strong
 pain

Side effects: obstipation, †liver tests, histaminoliberation

CAVE: codeine and dihydrocodeine are prescribed to normal forms - without blue band!



Partial agonists + mixed agonists - antagonists

- · lower affinity to μ receptors, high affinity to κ rec.,
- respectively κ-agonists μ-antagonists or partial μ receptor agonists (buprenorphine)
- less potential for addiction, but exists!
- lower analgesic effect than full agonists
- · less side effects than full agonists



Partial agonists + mixed agonists - antagonists

buprenorphine

- partial μ rcp. agonist
- ↓tolerance in comparism with other opioids
- ↓abuse potential, obstipation and other GIT effects
- ·↑ "first pass effect"! Do not administer orally!!!

· Use:

- 1) strong chronic pain (TTS!)
- 2) substitution therapy of opioid (heroin) addiction → combined with opioid antagonist naloxone in one drug form (sublingual) → in injection application naloxone antagonizes effects of buprenorphine (in sublingual administration naloxone does not act!)



Partial agonists + mixed agonists – antagonists – other representatives

- mixed agonists antagonists
- usually μ -antagonists and κ -agonists (event. also δ -agonists)
- possibility of σ-receptor activation → psychotomimetic and hallucinogenic effects)
- analgesic effects are weaker than full agonists
- today minimal use
- **pentazocin**, **butorfanol** in Czech Rep. not registered



Partial agonists + mixed agonists – antagonists – other representatives

nalbuphine

- for short-term therapy of moderate and strong pain
- unsuitable for long-term therapy
- parenteral administration (i.v., i.m., s.c.)
- causes respiratory depression comparable to morphine, but suppression of respiratory center has drug ceiling effect
- <u>Use:</u> perioperative pain, suppression of pain in obstetrics (BE AWARE: in newborns risk of breathing depression, bradycardia, cyanosis and hypotension → newborn's monitoring necessary!)



Atypical opioids

tramadol

low affinity to μ receptors + norepinephrine and serotonin reuptake inhibition (= atypical mechanism of action, similar to some antidepressants from SSRI group; effect of tramadol can not be fully antagonized by opioid antagonists)

- approximately 1/6 1/10 of morphine analgesic potency
- very suitable combination with paracetamol
- less side effects (minimal respiratory depression)
- risk of serotonin syndrome
- in Czech Rep. very often prescribed analgesic, prescription to normal forms without blue band; more drug forms
- RISK OF ADDICTION!!!

Use: therapy of moderate and strong pain (acute and chronic)



Atypical opioids

tapentadol

- dual mechanism of action
- μ agonist + NRI (+ σ agonist) NEW GROUP - MOR-NRI (μ receptor agonism - noradrenaline reuptake inhibitor)
- more effective than tramadol, analgesia comparable with oxycodone, but less adverse effects
- suitable for the treatment of acute (but also chronic pain e.g. vertebrogenic; also effective in diabetic neuropathy neuropathic pain!!!)
- relatively few adverse effects (compared to classical strong opioids, e.g. oxycodone)
- p.o. administration (also tbl. with sustained release)

CAVE: tapentadol is prescribed to forms with blue band ("opiate forms"), very strict accounted and subjected to the rules for handling with narcotics and psychotropic drugs and their precursors!!!



Antagonists of opioid receptors

naloxone, naltrexone

Indications: treatment of opioid intoxication, treatment of respiratory depression induced by opioids, addiction diagnostics (withdrawal symptoms)

TRIAD: coma, respiratory depression, miosis



Opioid-induced side effects

- respiratory depression (suppression of breathing)
- nausea and vomiting
- sedation, inhibition of cognitive functions
- constipation (solution = oxycodone + naloxone)
- ADDICTION
- be carefull in pro-convulsive states! (e.g. epilepsy proconvulsive action – decrease of the threshold for seizures)
- † intracranial pressure



Intoxication by opioid agonists

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nausea, "flush", tinnitus

apathy, sedation, sleep, miosis

superficial breathing

cyanotic, cold skin, tachycardia

asphyxia

TRIAD: coma, respiratory depression, miosis
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Treatment:

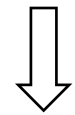
naloxone i.v.

ventilation, vital functions,

parenteral liquids in unconsciousness



Withdrawal symptoms



occur approximately after 3-4 weeks of opioid administration

"craving" ("drogenhunger"), "craving" for the another dose (psychic addiction arises easiest to heroin, oncology patients treated with opioids → < 1% of patients)

unrest, depression

anxiety, weakness, nervousness, mydriasis

lacrimation, ↑ nose secretion, frisson (goosebumps),

↑ perspiration, pain, stenocardia



Rotation of opioids

Switch in case of AE

Sometimes even in equianalgesic dose increase of effect



Other indications of opioids

- antitussive effect
 - can be induced by codeine and dextromethorphan in dry nonproductive cough
- constipative effect
 - can be induced by loperamide and diphenoxylate in functional diarrhea
- premedication before anaesthesia and surgery under general anaesthesia
 - leads to calm the patient and based on the synergism of drugs reduces the total dose of narcotics (thereby increasing the safety of anaesthesia)
 - particularly fentanyl and its derivatives are used
 - combination of opioid analgesic with neuroleptic (fentanyl + droperidol) within neuroleptanalgesia
- replacement (substitution) therapy of addiction to heroin or other opioids methadone, buprenorphine



General rules of pain pharmacotherapy management

- **■**WHO's pain relief ladder
- ¶ Step 1 (VAS 0-4)
 - non-opioid analgesics ± adjuvant treatment
- - pain persists, intensifies, no change in the objective finding
 - weak/moderate opioid analgesics ± non-opioid analgesics ± adjuvant treatment
- - pain persists, intensifies, there is no indication for another treatment
 - strong opioid analgesics ± non-opioid analgesics ± adjuvant treatment ± weak/moderate opioid analgesics

