MUNI MED reral principles of poisoning management. Specific antidotes in poisoning therapy.

General principles of drug addiction





- Interconnection of both disciplines
- They study the effects of chemicals on biological systems

Pharmacology - therapeutically useful effects, drugs

Toxicology - adverse, harmful (toxic) effects, poisons and toxins

Paracelsus (1493-1548):

"All substances are poisonous; there is none which is not a poison. The right dose differentiates a poison and a remedy"





Causes of poisoning

- 1. drugs 52%
- 2. industrial products 30% (chemicals for cleaning, organic solvents, cosmetics...)
- 3. plants 8%
- 4. pure bulk chemicals -5%
- 5. funghi 2%
- 6. animal poisons (snakebite) -1%
- 7. others -1%



General principles of acute poisoning treatment

Treatment has to be provided as quickly as possible but always with judgment so that therapeutical procedures do not cause worsening of the patient's state or even death !!!





General principles of poisoning treatment:

- eliminate the substance from organism as quickly as possible (= decontamination)
- antidote (rapid counteraction for poison by means of specific actions);

"a drug, chelating substance, or a chemical that counteracts (neutralizes) the effects of another drug or a poison"

vital functions + symptomatic treatment





• Gastric lavage and administration of emetic, preferably within 1 hour of intoxication (the first treatments should be done prior to transportation to the hospital)

An average patient arrives only after 3 hours





Induced vomiting

- in p.o. poisoning within 4 hours
- within 8 hours after anticholinergic agents
- within 12 hours of pylorospasm inducing agents (eg, salicylates)
- the patient is conscious, without spasms
- Syrup of ipeca (emetin)- non-reg., apomorphine (s.c.)
- mechanic stimulation of pharynx
- (red-eyed treefrog secretion)

Can not evacuate whole stomach content (max 30-50%)!



DO NOT INDUCE VOMITING IF ACIDS OR ALKALI WERE INGESTED

OTHER CONTRAINDICATIONS OF INDUCED VOMITING:



- Somnolence and loss of consciousness
- Intoxication with foaming agents
- Intoxication with hydrocarbons
- Attacks of spasms
- Alimentary intoxications in small infants





Adsorbents

- With poisons ingested p.o.
- Charcoal (adsorbing carbon = Carbo adsorbens) / diosmectit → large active surface
- $50 100 \, g$ in 5 10% suspension, possibly with stomach tube, then repeatedly $50 \, g$ per 4 hours
- Up to 2.5 g/kg

- +: paracetamol, salicylic acid, diazepam, amphetamine
- methyl/ethylalcohol, Li, strong acids and alkali



Toxic substances that are poorly adsorbable by Charcoal

- acids
- alkali
- chlorates
- chlorids
- cyanides
- nitrates
- ethanol
- ethylenglycol
- isopropanol
- methanol

- fluorides
- iron
- ferrous sulphate
- potassium
- sodium
- detergents







Gastric lavage

- In p.o. intoxications within 4 hours
- The patient is conscious, without spasms
- when unconscious, ONLY in lying position and intubated
- warm water (37°C), saline(preparation: 2 teaspoons of salt per 1 litre water), 300 ml
- Sample for toxicological analysis
- In the end (the last lavage) add adsorbent (30 g of activated carbon) or a laxative (Na₂SO₄)

1. Elimination of unabsorbed toxic substances from organism - PEG - laxative, GIT dialysis

- PEG polyethylene glycol in ionic solutions
- 4 liters / 2 hours
- until the evacuated rectal content is clear Indications (toxic and lethal doses):
- rugs bound poorly by charcoal: iron, lithium

Retarded tablets: theophylline, calcium blockers - verapamil, diltiazem!



Increasing the intestinal passage

The patient is conscious, with no spasms

- Administration of big doses of strong and quick-acting laxatives
- Sodium sulphate (20 30 g with a large volume of water)
- Mannitol (ca 50g per 1 litre water; 0.5 1 litre is administered p.o.)
- Castor oil (20 30 ml)



• CI in poisons soluble in fats!!! (castor oil ↑bile secretion and resorption of fats)



Total intestinal lavage

- Large-volume solution (25 ml/kg)
- Through stomach tube, until clean solution flows off
- Without resorption, does not cause diarrhoea
- It only rinses the intestine
- polyethylenglycol + NaSO₄, NaCl





Forced osmotic diuresis

- Infusion of saccharide solutions (20% mannitol; possible combination with furosemide), physiological solution
- Up to several litres / day
- CI: brain and lung oedema, heart failure, anuria





Forced alkali diuresis

- Speeds up elimination of slightly acidic poisons
- Alkalinisation of urine and blood (pH 7.5 9.0)
- NaHCO₃ solutions
- I: salicylates, barbiturates, sulphonamides, antipsychotic drugs,...
- CI: pulmonary oedema, shock, serious impairment of kidneys





Forced acidic diuresis

Speeds up elimination of slightly alkalinic poisons

- Acidification of blood and urine
- 5% Glc solutions with ammonium chloride in i.v. infusion
- I: amphetamines, quinine, quinidine, nicotine, morphine,...
- CI: serious impairment of kidneys



Peritoneal dialysis

Haemodialysis

Haemoperfusion



2. Neutralization of poison through administration of antidote



- Antidote a substance that neutralises the effect of poison
 - specific (using antagonistic effects of pharmaceuticals antidotes that can counteract the effects of poison either partly or completely)
 - Non-specific (adsorption activated medicinal carbon = carbo adsorbens carbo activatus carbo medicinalis)
 RATIO OF CARBON : TOXIC SUBSTANCE = 10 : 1
 (usually 50g / 3 4h; most often intoxications with medicines, chemicals)
 - It is necessary to administer antidote as quickly as possible
 - Dosage according to plasmatic level of toxin



Specific Antidotes



https://www.annemergmed.com/article/S0196-0644(17)30657-1/fulltext







- Check vital signs
- Intubation
- Entry into bloodstream
- Support of CVS (inotropics, vasopressors)
- Therapy of spasms



Toxicological Information Centre



Website of the Toxikologické informační střediskoTIS).

Acute poisoning - what to do?

Dial +420 224 91 92 93 or 224 91 54 02

To receive advice on first aid and what to do next.

Prepare:

- precise information on the accident
- full name
- birth identification number
- health insurance company
- healthcare professional also their IČP (organization identification number)
 In order to facilitate the consultation, the doctors are asked to calculate (provided it can be ascertained) the quantity of medication (active substance) that intoxicated the patient. Also please try to estimate or find out the body weight of the patient.



Toxicological Information Centre

- A 24/7 nationwide telephone medical information service to consult cases of acute human and animal intoxications
- For both laypersons and doctors
- The goal of the TIS is to decrease the number and severity of intoxications and to favourably effect the course of accidents. The Centre provides information on the chemical composition of commercial products and on the therapy of acute intoxications with these products
- It does not deal with:

the influence of chemical compounds on foetus cancerogeneity adverse effects of medicinal drugs impact of chemical compounds on the environment



Intoxication with medicines





Intoxication with medicines

Most often: sedatives, hypnotics, analgesics

Causes of death:

- Injury to CNS psychotropics
- Injury to CVS cardio*glycosides antiasthmatic drugs*
- Liver injury paracetamol, nimesulide, protease inhibitors,





General principles of drug addiction



Addiction = compulsive drug use despite harmful consequences



is characterized by an inability to stop using a drug (failure to meet work, social, or family obligations; tolerance and withdrawal).

accompanied by unnatural cravings that prompt the compulsive behaviors.

It is a primary, **chronic**, **neurobiologic disease** with genetic, psychosocial and environmental factors that influence its development and manifestations.

It is characterized by behaviours that include one or more of the following:

loss of control over drug use

continued use despite harm

compulsive use and craving



International Statistical Classification of Diseases and Related Health Problems 10th Revision:



Mental and behavioural disorders due to psychoactive substance use (F10-F19)

- F10: Mental and behavioural disorders due to use of alcohol
- F11: Mental and behavioural disorders due to use of opioids
- F12: Mental and behavioural disorders due to use of cannabinoids
- F13: Mental and behavioural disorders due to use of sedatives or hypnotics
- F14: Mental and behavioural disorders due to use of cocaine



International Statistical Classification of Diseases and Related Health Problems 10th Revision:

- F15: Mental and behavioural disorders due to use of other stimulants, including caffeine
- F16: Mental and behavioural disorders due to use of hallucinogens
- F17: Mental and behavioural disorders due to use of tobacco
- F18: Mental and behavioural disorders due to use of volatile solvents
- F19: Mental and behavioural disorders due to multiple drug use and use of other psychoactive substances



Historic context of drug use



Opium known already in neolithic age (8 000 – 5 000 years B.C.)

Coca and resin from hemp – known thousands years

Drugs were first used for their therapeutic purposes, secondary for they narcotic purposes

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caffeine (1820),
nicotine (1828),
cocaine (1859),
ephedrine (1887)
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Summary of frequently abused substances



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alcohol
   nicotine
      cannabinoids (hemp drugs)
         opioids
            benzodiazepines
               "classical" psychostimulant drugs
                   MDMA (exctasy)
                     "new" synthetic substances
                          hallucinogens
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19th century: beginning of commercial narcotics production

(e.g. morphine since 1828, cocaine 1862, heroine 1898)

Legal consumption of drugs was ended by opium conventions:

1909 Shanghai,

1912 Haag

1925 Geneva

Illegal way: French Connection (France), Cosa Nostra (USA)

After WWII:

Single Convention on Narcotic Drugs of 1961 - an international treaty to prohibit production and supply of specific (nominally narcotic) drugs and of drugs with similar effects



Council of the Government for Drug Policy Coordination - Annual registr (2012):

CZ – the most frequently abused drugs:

Psychostimulant drugs (particularly methamphetamine, syn. pervitin)

Hemp drugs (particularly <u>tetrahydrocannabinol – THC</u>).





Tolerance: a decrease in the effect of a drug as a consequence of repeated exposure (the effectiveness can decrease with continued use).

Mechanisms of Tolerance:

Pharmacokinetic Tolerance (enzyme induction effect)

It occurs because of a decreased quantity of the substance reaching the site it affects.

This may be caused by an increase in induction of the enzymes required for degradation of the drug e.g. CYP450 enzymes.

This is most commonly seen with substances such as ethanol.

This type of tolerance is most evident with oral ingestion, because other routes of drug administration bypass first-pass metabolism.





- Pharmacodynamic Tolerance (NT depletion, receptor plasticity)
 - It occurs when the cellular response to a substance is reduced with repeated use.
- This may be caused by a reduced receptor response to receptor agonists (receptor desensitization), a reduction in receptor density (usually associated with receptor agonists), or other mechanisms leading to changes in action potential firing rate.

Dependence: a maladaptive pattern of substance use, leading to clinically significant tolerance, impairment, or distress; an adaptive state associated with a withdrawal syndrome upon cessation of repeated exposure to a stimulus (e.g., drug intake).





Dependence develops when the neurons adapt to the repeated drug exposure and only function normally in the presence of the drug.

When the drug is withdrawn, several physiologic reactions occur. These can be mild (caffeine) or even life threatening (alcohol).

This is known as the **withdrawal syndrome**.





Physical dependence x psychological dependence

Physical dependence (physiologic dependence) referrers to the adverse physical symptoms and signs that result from the withdrawal of the drug.

It results from many of the same mechanisms that produce tolerance.

As with tolerance, homeostatic set-points are altered to compensate for the presence of the drug.

If drug use is discontinued, the altered set-points produce **effects opposite** to those manifested in the presence of the drug.





Physical dependence x psychological dependence

Psychological dependence

Psychological dependence is a change in emotional state that occurs after using a substance or engaging in a behaviour over a period of time.

i.e. dependency on specific psychological phenomena provoked by the drug (e.g. euphoria)

This change in emotional state is a result of changes in brain chemicals.

It can cause **craving**, motivation to seek out the substance or behavior, irritability, anxiety, or general dissatisfaction when withdrawing from the substance or activity.

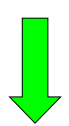




Inverse tolerance (sensitization): the drug becomes more effective with repeated doses.



Tolerance



Decreased response to substance effects. A higher dose is required to achieve the same effect.



(Robinson & Berridge, 1993)



•<u>Increased</u> response following repeated drug administration

Substance is usually given in <u>shorter intervals</u> or continuously

Intermitent drug administration





Inverse tolerance (sensitization):



There are two hypothesis to explain mechanism of sensitization to psychostimulants:

1) Intermittent exposure to a drug will cause intermittent dopamine release. This will lead to decreased sensitivity or density of pre-synaptic dopamine autoreceptors.

These receptors are responsible for negative feed back \rightarrow increased dopamine release \rightarrow increase stimulatory effects of dopamine.

Important rather for development of sensitization than expression.

2) Long-term intermittent exposure to a drug provokes intermittent release of high amounts of dopamine → gradual depletion of dopamine in cytoplasm.

This results in increased sensitivity of D_1 postsynaptic receptor (because they are not stimulated by their natural ligand).

Thus, after challenge dose administration (that acts through activation of the same postsynaptic receptors), an augmented behavioural response can be expected.



Exposure to alcohol and other drugs (AODs):

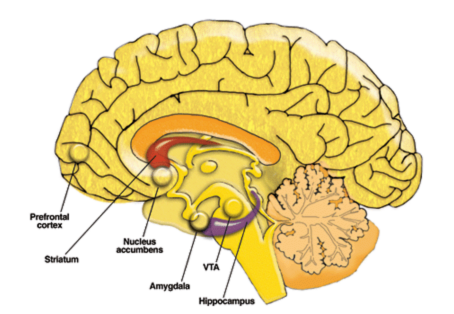


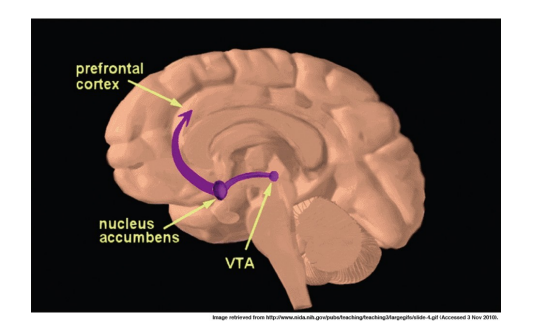
Plastic changes associated with AOD use - release of the neurotransmitter dopamine from cells in ventral tegmental area (VTA) induced by addictive drugs.

The VTA is one of the components of the mesolimbic dopamine system – **REWARD PATHWAY**.

Neurons whose cell bodies are located in the VTA, extend long axons most prominently to the nucleus accumbens (NAc) and the prefrontal cortex

Dopamine release in the mesolimbic system is critical for the drive to ingest AODs.







The mesocorticolimbic dopamine system as an initial target of addictive drugs.

The VTA, at the origin of the mesocorticolimbic system, is composed of dopamine projection neurons that are under inhibitory control of GABA interneurons



The main targets are the NAc and the mPFC.

Addictive drugs cause an increase in mesocorticolimbic dopamine through:

- 1) direct activation of dopamine neurons (e.g., nicotine);
- 2) indirect disinhibition of dopamine neurons (opioids, cannabinoids, benzodiazepines);
 - 3) interference with dopamine reuptake (cocaine, ecstasy, and amphetamines).

