



## Drugs used in diseases characterized by bronchial obstruction



## **Bronchial asthma**



chronic inflammatory disease of airways affecting 300 million people all across the globe prevalence in CZ: 8 %, in children over 10 %

### **Characteristics:**

bronchial hyper-reactivity obstruction (often reversible) inflammation

## **Symptoms:**

shortness of breath (bronchoconstriction, mucous plug, oedema, airway remodeling due to the inflammation)

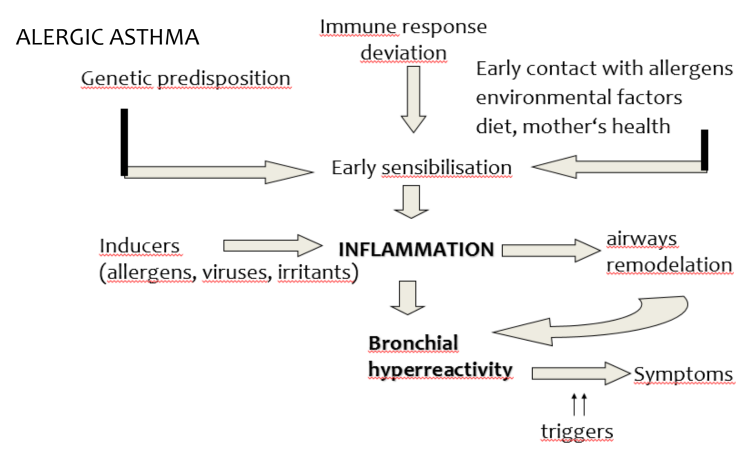
difficult and prolonged **expiration** → wheezing, whistling

cough (especially at night or in early morning)



## **Bronchial asthma**



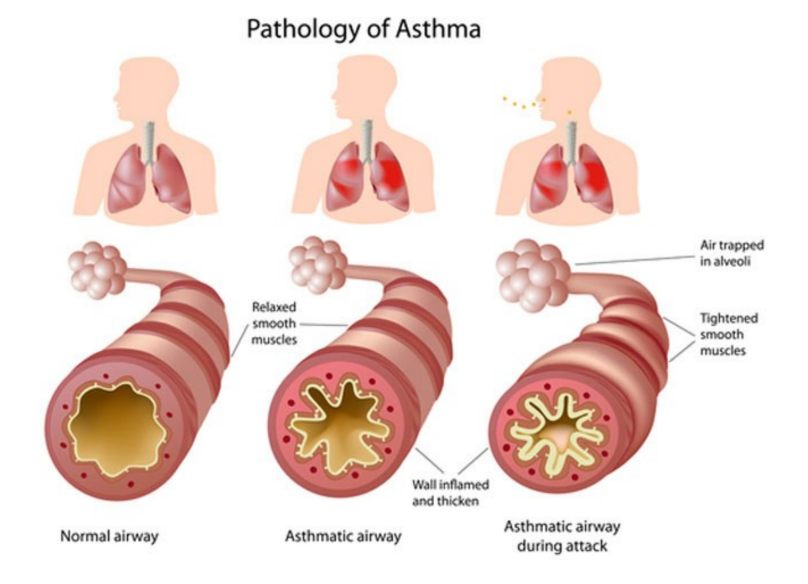


#### **NON-ALERGIC ASTHMA**

- allergy not present
- excercire-induced, aspirin-sesitive, infectious, work-related, endogenous









## Diagnose



Anamnesis – personal, familiar

Clinical examinations - auscultation, signs of atopy, eosinophilia,

PEF – Peak Expiratory Flow

FEV 1 – Forced Expired Volume

Laboratory tests- eosinophilia, IgE

Allergy testing



## Classification with regard to seriousness



Intermittent – sign up to once a week, night symptoms up to twice a month, pulmonary function normal

Mild persistent – signs no more than once daily, night symptoms up to twice a month, PEF at least 80 %

Moderate persistent— signs once a day and are not permanent, night sign no more than once a week, PEF 60-80 %

**Severe persistent**– permanent signs, daily, obstruction, PEF ≤ 60 %



## Managment of asthma



the disease itself cannot be fully treated, the goal is to keep asthma under control

#### **Goals:**

minimalize both acute and chronic symptoms reduction of exacerbations (lessen SABA administration) improvement of the quality of life (physical activity) avoid adverse effects of the treatment



## Chronic obstructive pulmonary disease (COPD)



affecting 600 million people all across the globe prevalence: 8 %

risk factors: smoking, polluted air, dust and chemical vapors at workplace, genetic predisposition

#### **Characteristics:**

chronic inflammation caused and maintained by long-term exposure to harmful agents (irritating gases and particles) poorly reversible, progressing bronchial obstruction production of mucus

### **Symptoms:**

cough (usually whole day, hardly ever only during night)

expectoration

shortness of breath



## Managment of COPD



we can only slow the progression reduction of risk factors is necessary (mainly top quit smoking)

### **Goals:**

symptom reduction

improvement in physical condition and overall health state

prevention of complications and exacerbations



## **Administration**



oral, parenteral (injections, infusions)

#### inhalation

- local administration, high drug concentration at the site of action
  - fast onset of the effect
- minimal penetration to systemic circulation  $\rightarrow \downarrow$  risk of side effects



## Drugs used in diseases characterized by bronchial obstruction

BRONCHODILATATORS



- $\beta_2$  sympathomimetics
- parasympatholytics
- glucocorticoids
- methylxanthines
- roflumilast (COPD only)
- antileukotrienes
- imunoprofhylactics
- monoclonal antibodies

asthma only

- noselective sympathomimetics (epinephrine, life-saving medication)
- adjuvant medication (antitussics, drugs facilitating expectoration)



## $\beta_2$ sympathomimetics



**MoA:** selective  $\beta_2$  stimulants

- inhibition of mediator release from mast cells + stimulation of ciliary beat frequency
- diagnostics post-bronchodilator test (salbutamol)
- mostly inhaled, may be also given orally (mainly in kids)
- not completely selective in their binding to β receptors long-term use = down-regulation of receptors



## $\beta_2$ sympathomimetics



Indication: asthma, COPD

**AE:** nervousness, tremor, cephalgia, palpitation, hypokalemia (mainly when given orally)

CI: hypertension, dysrhythmia, pregnancy



## $\beta_2$ sympatomimetics



Short-acting = SABA (also rapid-acting = RABA)
fast onset of effect, which lasts 4 – 6 hours, inhalation
salbutamol

fenoterol

Long-acting = LABA

effect lasts for up to 12 hours, inhaled or administered orally

salmeterol

clenbuterol

formoterol (RABA)

indakaterol (U-LABA)

vilanterol (U-LABA)



## Parasympatholytics



**MoA:** competitive antagonism of M receptors

- in a form of inhalation

- can be combined with  $\beta_2$ -sympathomimetics or glucocorticoids

**Indication: COPD**, asthma

**AE:** if entering the systemic circulation (low risk, they contain quaternary nitrogen in their structure) – anticholinergic effects

CI: glaucoma, prostate hypertrophy, pregnancy



## **Parasympatholytics**



## ipratropium

- used in asthma as well – in patients resistent to β2 sympathomimetic treatment (approx. 1/6 of patients) short acting (SAMA)

aclidinium (LAMA)

tiotropium (U-LAMA)

glykopyrronium-bromide (U-LAMA)

umeclidinium (U-LAMA)

COPD only



## Glucocorticoids



**MoA:** inhibition of phospholipase A2 by lipocortin

### **Effects I:**

↓ cytokine, PG a LT secretion

↓ lipolytic and proteolytic enzyme secretion

↓ endothelial permeability

block of cell migration

↓ bronchial hyperreactivity,



## Glucocorticoids



#### **Effects II:**

reduction of edema prevention of chronic irreversible changes (hypertrophy and hyperplasia of bronchial smooth muscles, subendothelial fibrosis and thickening of mucous basal membrane)

increase in sensitivity of  $\beta_2$  adrenergic receptors to  $\beta_2$ - SM



## MoA at the cellular level



glucocorticoid + cytoplasma receptor



production of specific mRNA



production of some proteins (lipocortins)



## MoA at the cellular level



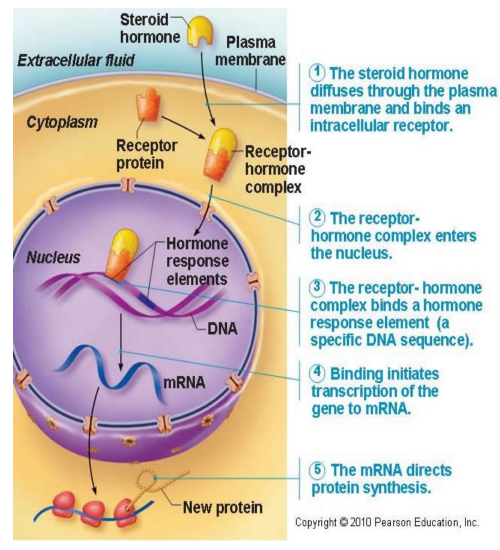
After entering the cell they bind to specific receptors in cytoplasm causing change of conformation = activation of receptors

Complexes of corticoid + receptor are transported to cell nucleus and bind to DNA elements.

The result is increased transcription of genes either inducing or inhibiting synthesis of other proteins

GLC receptors are present in all tissues!!!

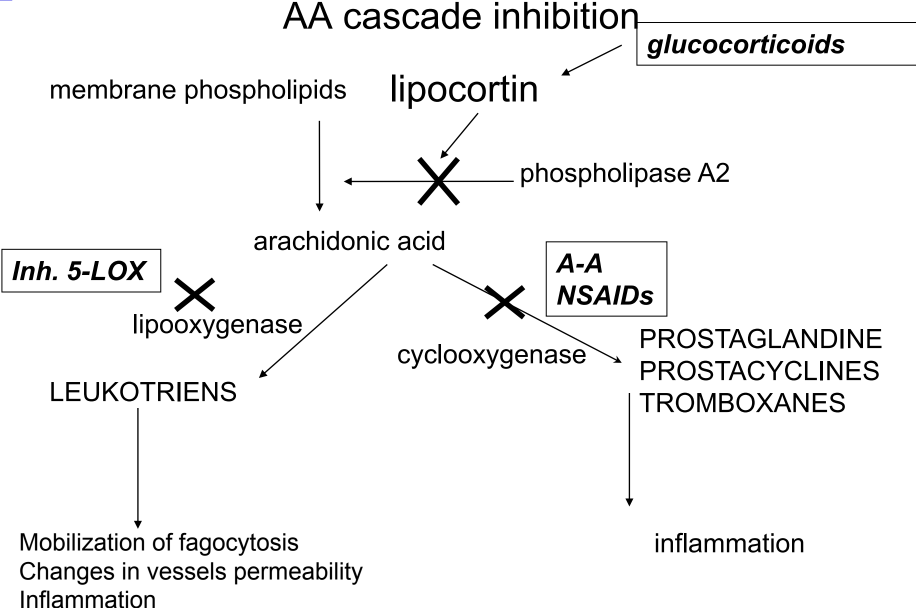
Proteins called **lipocortins** are able to suppress phospholipase A





## **Antiinflammatory effect of GC**







## Glucocorticoids



#### given by inhalation

lower risk of systemic adverse effects
AE: affected vocal cords – croaky voice, oral candidiasis (thrush)

beclomethasone budesonide fluticasone ciclesonide mometasone

#### systemic administration

orally, via injection – acute conditions, doses are gradually decreased, in severe persistent asthma – if nothing else is effective

prednisone triamcinolone hydrocortisone (injection)



## Methylxanthines



**MoA:** phosphodiesterase 1 – 4 inhibitors adenosine receptors antagonists

sustained-release drug forms

#### **Effects:**

- bronchodilatation
- cardiostimulation (+chrono, +inotropic eff. )
- diuretic eff.
- CNS and respiratory center stimulation
- stimulation of hydrochloric acid secretion



## Methylxanthines



#### **Effects:**

- substrates of CYP450 – be cautious if patient is a smoker!

CI: pregnancy, epilepsy, cardiovascular disease

**AE:** tachycardia, palpitations, sleeplessness



## Methylxanthines



## theophylline

- combination therapy with  $\beta_2$  SM is convenient
- becoming obsolent, therapeutic drug monitoring needed
  - variable pharmacokinetics, low therapeutic index

## aminophylline

- a complex of theophylline and ethylendiamine (better solubility)

- COPD, emphysema



## roflumilast



selective long-acting inhibitor of phosphodiesterase 4

reduces the inflammation in bronchi in COPD



## **Antileukotrienes**



**MoA:** antagonism of LT-receptors / inhibition of lipoxygenase

#### LT receptor antagonists:

treatment of persisting asthma, allows lowering of glucocorticoid dose 1-2x a day, orally

#### montelukast

#### **Inhibitors of LOX:**

need for frequent application not registered in CZ (**zileuton** – USA)



## Imunoprophylactics (mast cells stabilizers)



**MoA:** stabilisation of mast cell membrane  $\rightarrow \downarrow$  Ca<sup>2+</sup> influx  $\rightarrow \downarrow$  degranulation of mast cells and thereby  $\downarrow$  histamine release influence on lymphocyte function

prevention of asthma attack, they **do not affect already present bronchospasm** 

**Use:** as preventive, long-term, maintenance therapy – mild and moderate asthma when combined with other antiasthmatics, they allow lowering of their dose

CI: pregancy (1. trimester)

nedokromil, ketotifen (H1 antihistamine), cromoglycate



## Monoclonal antibodies



#### Anti-IgE

#### omalizumab

antibodies against a part of IgE, which binds to mast cells

Indication: severe persistent allergic asthma, which cannot be otherwise controlled

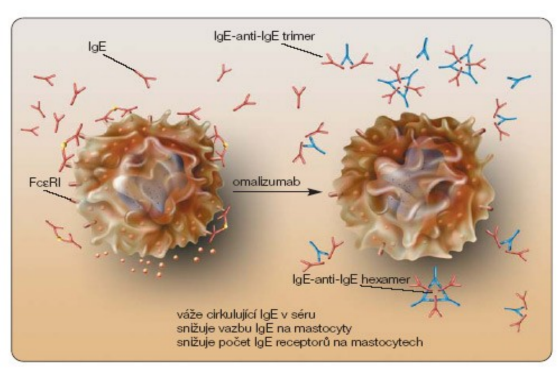
administered subcutaneously in specialized centers only



## Anti-IgE



### omalizumab



Obr. 3 Mechanismus působení omalizumabu



## **Monoclonal antibodies**



## Anti-IL-5

mepolizumab, reslizumab

add-on treatment for severe refractory eosinophilic asthma in adult patients



## Other options



## **Bronchial thermoplasty**

 bronchoskopic procedure, during which a therapeutic radiofrequency energy is delivered to the airway wall, resulting in reduction of smooth mucle cells

## Allergen immunotherapy

induces tolerance to the triggering allergen



## **Devices for inhaled medications**



**MDI** = metered dose inhalers drugs as solutions, propellants

**BAI** = breath-actuated inhalers

**DPI** = dry powder inhalers spinhaler, diskhaler, turbohaler

**nebulizers** (liquid → aerosol)



## **Devices for inhaled medications**



spacers for children and elderly

patient must be educated how to use their inhaler

→ up to 41 % of patients use incorrect technique

inhalers often combine two drugs (bronchodilator + glucocorticoid or two bronchodilators)

## MUNI MED









# Adjuvant medication in diseases characterized by bronchial obstruction and another drugs affecting respiratory system



#### antitussives

drugs facilitating expectoration

H₁ antihistamines (mainly II. a III. generation)