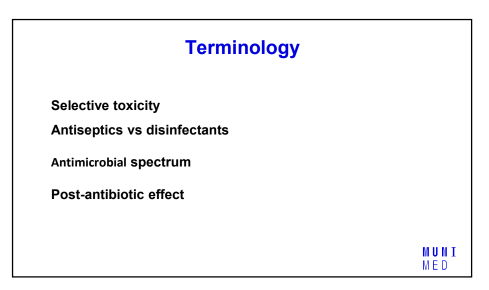
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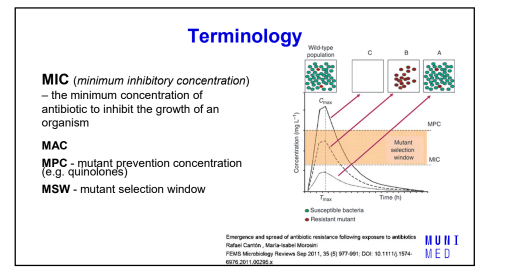
Antibacterial drugs

Department of Pharmacology, 2020

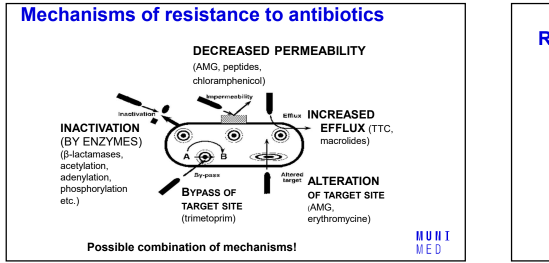
Notes for Pharmacology II practicals

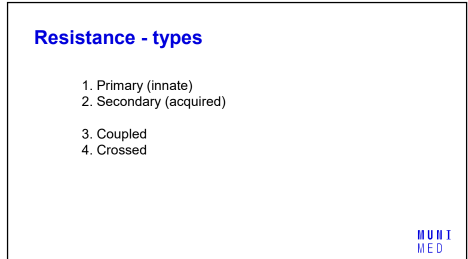
This study material is exclusively for students of general medicine and stomatology in Pharmacology II course. It contains only basic notes of discussed topics, which should be completed with more details and actual information during practical courses to make a complete material for test or exam studies. Which means that without your own notes from the lesson this presentation IS NOT SUFFICIENT for proper preparation for neither tests in practicals nor the final exam.



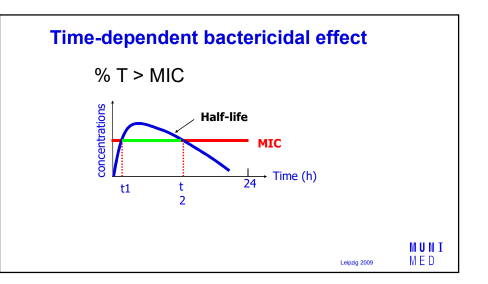


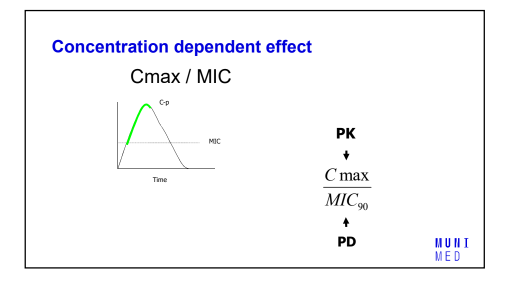
Terminology **Resistance to antibiotics** chromosomal determinants ٠ extrachromosomal determinants: genes for resistance to ٠ antibiotics ("r genes") – R plasmids Drug resistance can be spread: from person to person by bacteria ٠ from bacterium to bacterium by plasmids ٠ from plasmid to plasmid (or chromosomes) by transposons ٠ MUNI MED

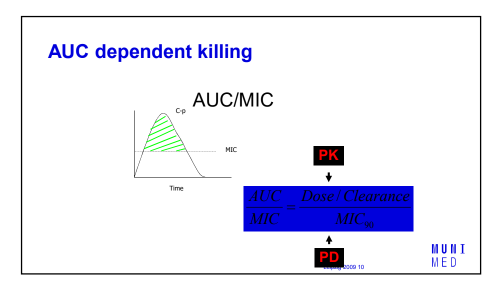


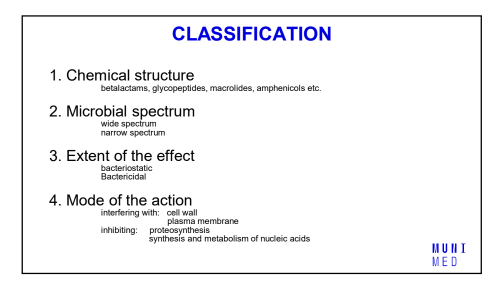


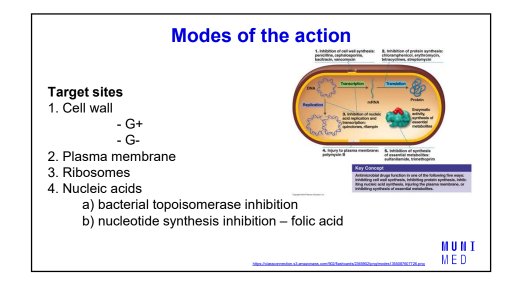
Terminology			
Pharmacodynamic index	Definition	Effect	Examples
T>MIC	Once the concentration of an antibiotic is above the MIC (typically 3-5 times greater than the MIC), there is not an increased rate of killing with increasing concentrations of antibiotic	Time dependent	beta-lactams
Cmax/MIC	As the concentration of an antibiotic increases, its rate of killing increases	Concentration dependent	aminoglycosides
AUC 0-24/MIC	The rate of bacterial killing is both related to the amount of time above the MIC and the total exposure of antibiotic to the organism	AUC dependent	glycopeptides

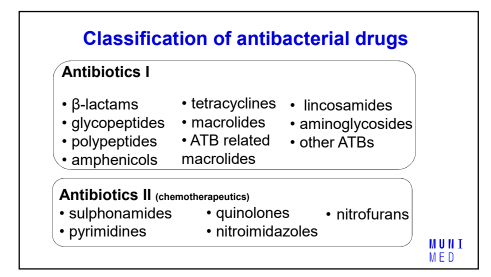


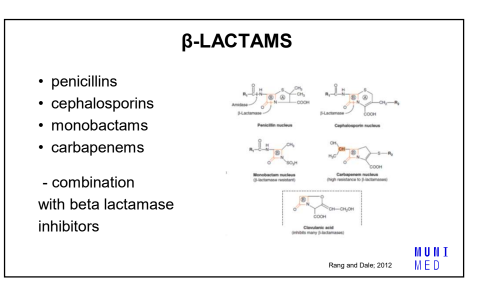


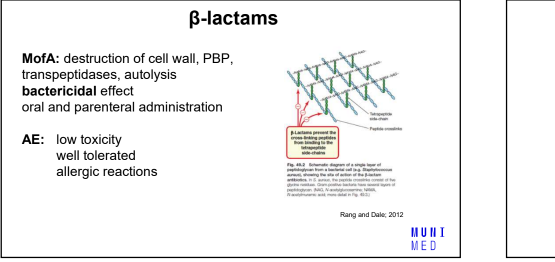


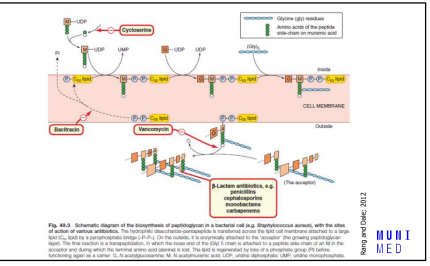






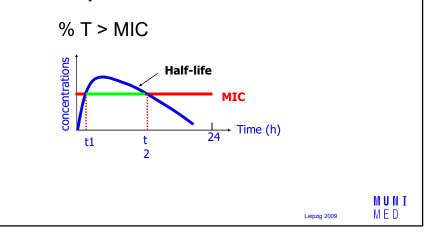


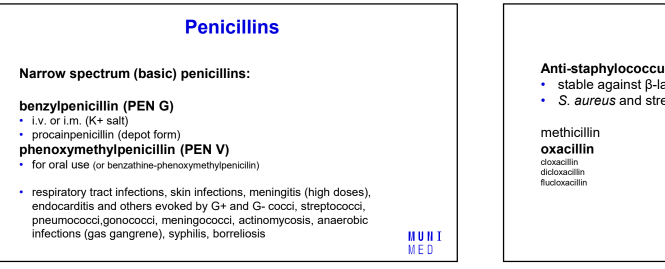


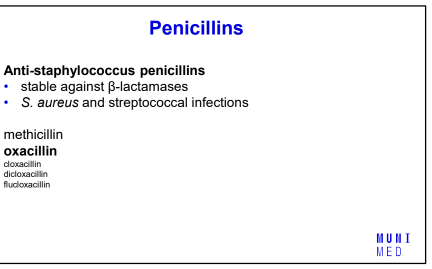


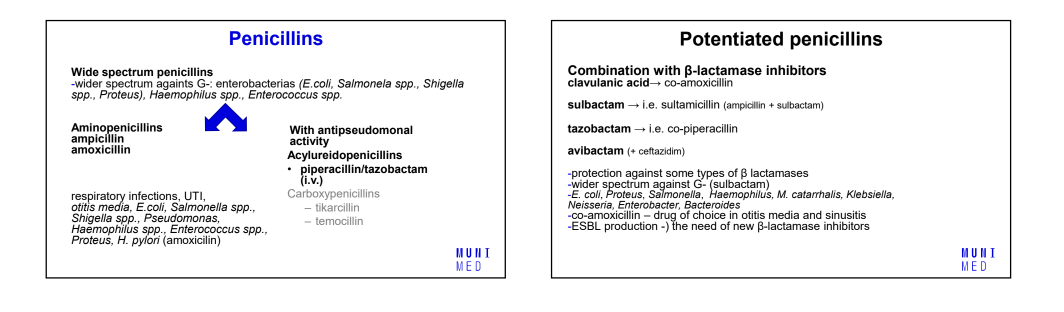
	PENICILLINS	
natural or semisynthet	tic	
Classification:	narrow spectrum	
	anti-staphylococcus	
	wide spectrum	
PK: i.v., i.m., p.o.		
-well distributed to boo	dy fluids, passing into joints, bile, saliva, milk and	
across placenta		
- lipid-insolubile, do no		
-short t _{1/2} , renal excret	tion	
	codynamic driver of effectiveness	
-T>MIC main pharmad -dosing every 6-8h, SI		
		MILNT

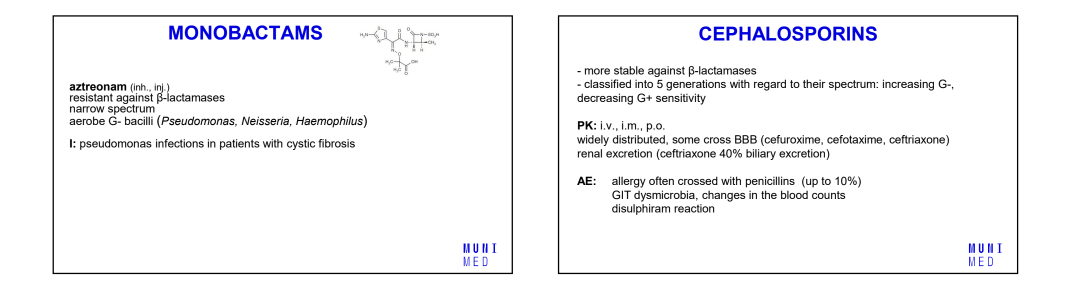
Time-dependent bactericidal effect











Cephalosporins

Ist generation cefazolin cefadroxil (p.o.) G+ cocci (*staphylococci, streptococci*), *E. coli, Proteus, Klebsiella, Neisserie* other G- are usually resistant (e.g. haemophilus) I: S. aureus infections, prophylaxis in surgery, tonsil pharyngitis, bronchitis, sinusitis, urinary infections

IInd generation cefuroxime (cefuroxime axetil p.o.) cefprozil

wider spectrum against G+ i G- : H. influ., enterobacterias, Neisseria, Proteus, E. coli, Klebsiella, Moraxella catarrhalis, anaerobes and B. fragilis. I: tonsil pharyngitis, bronchitis, sinusitis, urinary infections, borreliosis

Cephalosporins IIIrd. p.o. ceftriaxon cefixim i.v.: cefotaxim ceftazidim cefoperazon (+ sulbaktam) · enterobacterias, partially pseudomonades more stable against β -lactamases, higher efficacy (the best for G-) ٠ all i.v. agents cross BBB!!!! I: meningitis, UTI, respiratory infections, infections of skin, bones, joints; septicemia MUNI MED



GLYCOPEPTIDES

vancomycin, i.v. teicoplanin, i.v.

MofA: cell wall synthesis inhibition – binding to pentapeptide precursor; bactericidal

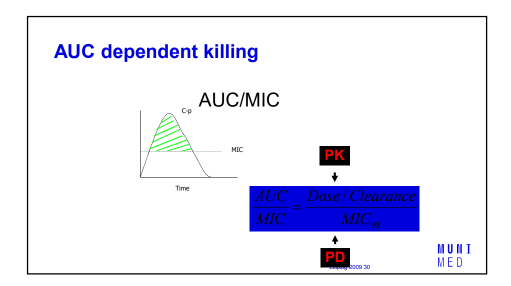
resistance, VRE; synergic effect with aminoglycosides TDM - vancomycin

PK: i.v. infusion, min. mtb., renal excretion
 I: reserve ATB for the serious, resistant G+ infections (MRSA) – endocarditis, osteomyelitis, pneumonia; local (p.o.) intestinal infections (not absorbed from gut)
 AE: rashes (red man syndrome), ototoxicity, nephrotoxicity

LIPOPEPTIDES

daptomycin - only G+ (MRSA - skin, endocarditis); in combination therapy in G+-

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LIPOGLYCOPEPTIDES dalbavancin dalbavancin oitavancin • similar antimicrobial spectrum with vancomycine, higher activity against G+ • dalbavancin - extremely long plasma half-life (14 days) • perspective therapy (skin infections, OPAT regimen - Outpatient Parenteral Antimicrobial Therapy)

POLYPEPTIDES

colistin (colistimethate; polymyxin E), i.v., inh. polymyxin B

MofA: disrupts the plasma membrane by its detergent activity

I: aerobic, multiresistant G- (*Ps. aeruginosa, Haemophilus, Klebsiella*) local application (oph., ORL, GYN, gut decontamination, cystic fibrosis) or infusion/injection

AE: nephrotoxicity, ototoxicity, neurotoxicity! return to use of colistin in nosocomial infections

II D

TETRACYCLINES

doxycykline, p.o.

tigecycline (glycylcyclin), i.v. - Clostridium difficile therapy! minocycline, tetracycline

MofA: proteosynthesis inhibition – reversible binding to 30S ribosomal subunit; bacteriostatic

primary resistant staph., strept. + pneumococci!

PK: doxycycline absorption p.o., (non-absorbable complexes with cations in GIT), lipophilic, widely distributed, high conc. in bile, enterohepatic recirculation

AE: disrupts tooth enamel and bone matrix – interfere with growth \rightarrow CI in children and in pregnancy, lactation, phototoxicity, dysmicrobia – GIT disturbances, vaginal dysmicrobia, suprainfection, hepatotoxicity

I: respiratory and urinary tract infections, ORL, therapy of biliary tract inf., borreliosis, syphilis, gonorrhea, ureaplasma, leptospirosis, chlamydiosis, mycoplasmosis, acne (minocycline)

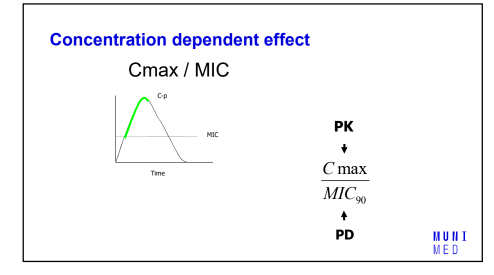
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gentamicin, amikacin (i.v.) isepamicin, netilmicin, tobramycin (inh.) kanamycin (oph.), neomycin (oph., drm., vag.)

MofA: proteosynthesis inhibition, irreversible binding to 30S ribosomal subunit (bactericidal effect), not in anaerobic bact.
post antibiotic effect and concentration-dependent effect
PK: parent. (highly polar molecules), not cross BBB, T_{1/2} 2-3hod, renal excretion (>50% unchanged)
AE: nephrotoxicity, ototoxicity, ↑↑ doses - neurotoxicity
I: sepsis, serious uroinfections (pyelonephritis), lower respiratory infections (in

combination), orthopedic and surgical infections (postoperative) syst. toxicity (TDM!) - not drugs of choice, comb. therapy (β-lactams)



Administration of aminoglycosides

- in combination therapy
- in one daily dose
- concentration dependent effect+ post antibiotic effect
- more daily doses
- synergic effect in comb. with β-lactams (exceptionally glycopeptides)
- in bacterial endocarditis caused G+ cocci (enterococci, staphylococci)

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Macrolides

- AE:
 - GIT intolerance diarrhea, anorexia, nausea, vomiting, cholestatic jaundice
 - · allergies
 - suprainfections
 - prolong. QT int.
- drug interactions
 - CYP inhibitors
 - increase in blood levels of statins, antiepileptic drugs, BZD, antidepressants, monoclonal antibodies, , immunosuppressant drugs (cyclosporine, tacrolimus), warfarin
 - decrease in effects of clopidogrel, betalactams, lincosamides

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Macrolides

clarithromycin, i.v., p.o.

- both upper and lower respiratory infections, *Mycobacterium leprae, otitis media*, skin and soft tissues
- in combination therapy Helicobacter pylori
- · not in pregnant women (interference with angiogenesis)
- prolongs QT interval
- high risk of drug interactions

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Macrolides

azithromycin, p.o.

best penetration to most tissues less drug interactions long T_{1/2} post-antibiotic effect DO NOT use in common infections, tonsillitis etc....

roxithromycin, p.o.

safe in pregnant women (with allergy to betalactams)

spiramycine, p.o.

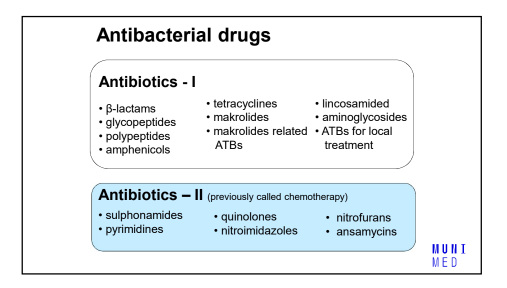
drug of choice in **congenital toxoplasmosis** safe in patients treated with theophylline

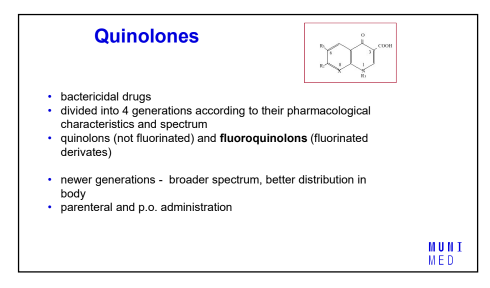
MUNI Med

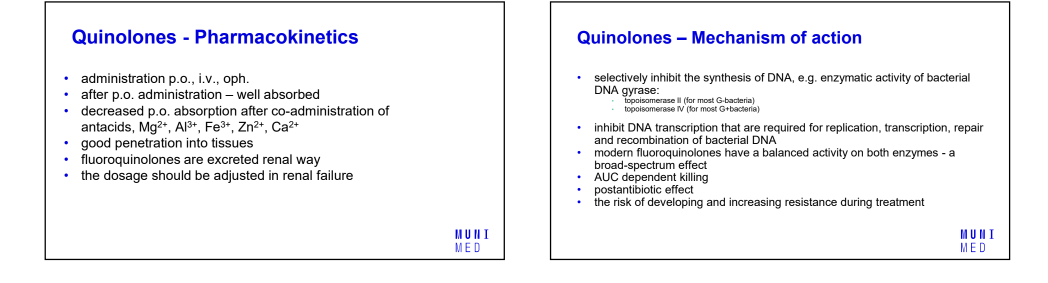
ATB related to macrolides Oxazolidinones Iinezolid i.v, p.o. • novel MofA (inhibition of proteosynthesis – blocks formation of 70S ribosome) • G+ (MRSA, VRE, nosocomial/community pneumonia, *Cl. difficile*) • non-selective MAO inhibitor – interactions • serotonin syndrome

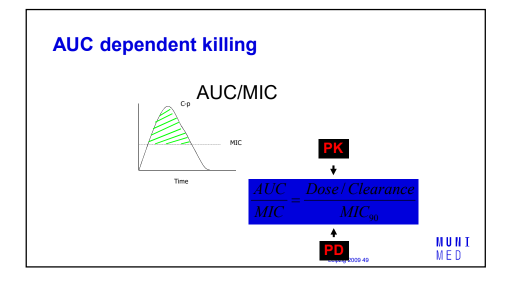
ATB related to macrolides cl Μ Streptogramins rik quinupristin dalfopristin G+ (MRSA, VRE) P pla Ketolides A telithromycin -C pneumonia, bronchitis, sinusitis, tonsillitis/pharyngitis in infections resistant to I: beta lactam and macrolide therapy OS solithromycin – MRSA, gonococci er ali MUNI MED

LINCOSAMIDES	
lindamycin, p.o., i.v., i.m., loc.	
lofA: proteosynthesis inhibition – reversible binding to 50S bosomal subunit	
'K : p.o. and parent., well penetrates to teeth and bones, lacenta, milk, not cross BBB	
LE: allergy, pseudomembranous colitis crossed-resistance with macrolide	
respiratory infections, skin and soft tissues infections, steomyelitis, dental, intraabdominal, gyn., pneumonia, malaria, ndocarditis prophylactic use, gynecologic infections (loc.),	
Iternative treatment of beta lactams hypersensitivity	MUNI Med







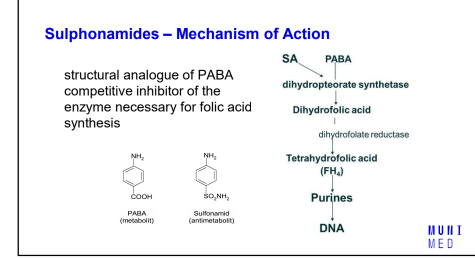


Sulphonamides

Wide-spectrum:

G+ and G- bacteria, streptococci, hemophilia, actinomycetes, nocardiosis, Pneumocystis jiroveci, chlamydia, Toxoplasma gondii, Neisseria meningitides ineffective in Pseudomonas, Proteus - resistance !!!

Bacteriostatic, in combination - bactericidal



Sulphonamides Long acting effect (8-10 hours) sulphamethoxazole in combination with trimethoprim (cotrimoxazol, SMZ-TMP) Local use: sulphasalazine microflora metabolizes it to sulphapyridine (SA) and 5aminosalicylic acid (anti-inflammatory) – inflammatory bowel disease silver salt of sulfadiazine (local skin treatment) sulphacetamide (oph.)

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Sulphonamides - Pharmacokinetics

- parenteral and p.o. administration, local use
- good absorption >70%
- great penetration into tissues and cells
- hepatic metabolism via acetylation and glucuronidation
- high binding to plasma proteins displacement of other drugs and increase of their free fraction
- Drug interactions!!!
 - p.o. anticoagulants, methotrexate, sulphonylureas
- penetrate to the placenta and partly HEB
- renal excretion

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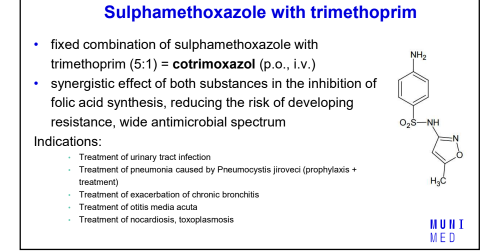
Sulphonamides - Adverse Effects

- GIT disorders
- Allergic skin reactions rash (Stevens-Johnson's and Lyell's syndrome), photosensitivity, drug fever (5-10 days after initiation of treatment) even with topical application
- Hematotoxicity hematopoietic disorders, bone marrow suppression, anemia, leucopenia, thrombocytopenia
- · Deficiency of folate megaloblastic anemia
- Interstitial nephritis risk of precipitation in the urinary tract acid pH of the urine (avoid of acidic foods, vitamin C, acetylsalicylic acid ...)

KI:

gravidity and breastfeeding

newborns (until 2 months) with immature enzymatic system (hyperbilirubinemia) MUII I



Sulphasalazine anti-inflammatory drug with an immunosuppressive effect derivate of aminosalicylic acid Indication: Treatment of ulcerative colitis, Crohn's disease, rheumatoid arthritis (DMARDs) after p.o. administration 30% of dose is absorbed 70% is degraded by intestinal bacteria in the colon: sulphapyridine inhibits the action of NK cells and transforms lymphocytes AE - nausea, vomiting, abdominal pain, drowsiness, anuria, crystalluria and / or hematuria, convulsions mesalasine (5-aminosalicylic salt) inhibits cyclooxygenase and lipoxygenase in the intestinal wall, thereby preventing the formation of prostaglandins, leukotrienes and other inflammatory mediators MUNI MED

Silver salt of sulphadiazine

Local use – cream, impregnated bandage

Indications:

 prophylaxis and treatment of infected skin lesions, wounds, abrasions and burns, leg ulcers and bed sores

CI:

- preterm infants and infants up to one month of age
- pregnant and nursing women

Sulphacetamide

Local use – eye drops Indications:

- Treatment of eye infection and inflammation
- Prophylactically after injuries and burns of eye

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Sulphonamides - Indications

- UTI (SMZ-TMP)
- respiratory infection (Pneumocystis pneumonia) (SMZ-TMP)
- otitis media acuta (SMZ-TMP)
- malaria, nocardiosis (sulphadoxine)
- Local treatment of eye infection (sulphacetamide)
- Local treatment of skin burns, dekubitus (silver salt of sulphadiazine)
- ulcerative colitis, Crohn's disease (sulphasalazine s 5aminosalicylic acid)

Quinolones – Indications in general

Treatment of infections urogenital system (UTI) Treatment of respiratory infections Treatment of infections skin, bones, joints, soft tissues, gonorrhoea

Fluoroquinolones are back-up drugs, indicated only in situations where other antibiotics are inactive in vitro or inappropriate for treatment because of toxicity or side effects.

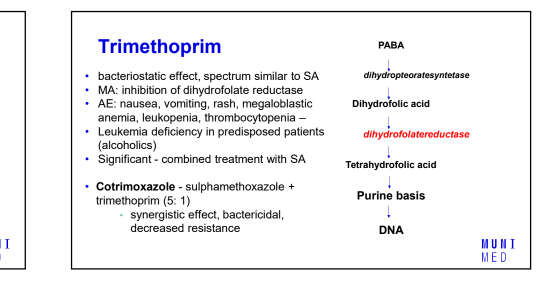
Usually administered in combination with other ATBs

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Quinolones – Adverse Effects often, but mild (nausea, vomiting, neurotoxicity, cramps, vertigo, headache) GIT disorders (5%) - nausea, vomiting • CNS toxicity (1-4%) - headache, vertigo, spasm, convulsion, depression (elderly patients) Prolongation of QT interval, malignant arrhythmia allergy (1-2%), photo toxicity hepatotoxicity IT – antacids, theophylline, caffeine, warfarin, cyclosporine tendinitis/tendinopathy, rupture of Achilles tendon arthropathy in animal models (in children with cartilage damage not shown except for arthralgia (1.3%) in patients with CF) KI: newborns and children (inhibition of bone cartilage growth), 1. trimester of pregnancy, breastfeeding epilepsy MUNI MED

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Generation	Drug	Indication
I.	nalidixic acid, oxolinic acid	Drugs with limited effect on G- (urinary ATBs)
	norfloxacine ofloxacine	Treatment of UTI
II.	ciprofloxacine	Treatment of respiratory, UTI, GIT infections, bones, joints, soft tissue, skin infections enterobacteria, <i>P. aeruginosa</i> , neisseria, haemophilus, legionella, <i>Neisseria meningitidis</i> , Anthrax
lev	levofloxacine	Drugs with higher activity on G + (pneumococcus), respiratory ATB
III.	sparfloxacine, gatifloxacine, tosufloxacine, pazufloxacine	Drugs more effective against G+ (pneumococcus), respiratory ATBs
IV.	trovafloxacine, gemifloxacine, sitafloxacine,	Drugs more effective against anaerobes, same spectrum as III. generation of cephalosporines



Nitroimidazoles

primarily bactericidal effects on anaerobes and protozoa

Mechanism of action: inhibition of DNA replication

Indications:

- treatment of peptic ulcers Helicobacter pylori eradication
- in combination with other antibiotics peritonitis
- amoebic dysentery intestinal disease
- trichomoniasis caused by *Trichomonas vaginalis* in women it is manifested by vaginal discharge, men show inflammation of the urethra, both partners should be treated simultaneously

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Nitroimidazoles

- Pharmacokinetics:
- 80% absorption after p.o. administration
- good penetration into tissues and cerebrospinal fluid, through the placenta into breast milk (KI)
- renal excretion
- AE:
- GIT disorders nausea, vomiting, diarrhea
- CNS disorders (dizziness, insomnia, depression)
- dark colored urine
- · long-term administration neutropenia, leukopenia (blood count)

metronidazole (disulfiram effect)

- ornidazole
- tinidazole

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 bacteriostatic, at higher concentrations bactericidal ATBs G + and G-bacteria, protozoa Mechanism of action: non-specific inhibition of bacterial enzymes release of superoxides and other oxygen compound Indications: prophylaxis and treatment UTI (nitrofurantoin, p.o.) gynecological infections, including trichomoniasis (nifuratel, p.o., topical treatment) intestinal infection 	Nitrofurans	
prophylaxis and treatment UTI (nitrofurantoin, p.o.) gynecological infections, including trichomoniasis (nifuratel, p.o., topical treatment) intestinal infection	G + and G-bacteria, protozoa Mechanism of action: • non-specific inhibition of bacterial enzymes	
	prophylaxis and treatment UTI (nitrofurantoin , p.o.) gynecological infections, including trichomoniasis (nifuratel , p.o., topical treatment)	MUNI Med

Nitrofurans

AE:

- allergy GIT disorders

- Gri disorders
 hepatotoxicity
 hematopoietic disorders megaloblastic and haemolytic anemia
 neurotoxicity
 pneumonia
 KI:

- pregnant, breastfeeding children

Ansamycines

 inhibit bacterial RNA polymerase, bactericidal effect 	
Indications: Treatment of pulmonary tuberculosis, G +, G-bacteria Mycobacterium sp. easy resistance - always in combination! drug interactions: inductors of CYP 450	
 AE: GIT disorders (nausea, vomiting, increase in liver enzymes, jaundice) Hematopoietic disorders (leukopenia, thrombocytopenia, anemia) arthralgia, myalgia 	
rifabutin (i.v., p.o., local use) rifamixine (non-absorbable form) – p.o., local use	MUNI Med

mupirocin MofA: proteosynthesis inhibition I: impetigo, folliculitis, furunculosis bacitracin + neomycin oph, drm., nas. fusidic acid inhibits synthesis of proteins in cell wall	
oph, drm., nas. fusidic acid inhibits synthesis of proteins in cell wall	
inhibits synthesis of proteins in cell wall	
against G+ - staphylococci I: impetigo, superficial folliculitis, skin wounds with infection; with betamethasone atopic dermatitis and contact dermatitis	
retapamulin (fusafungin)	MUNI Med

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ATB combinations	
Advantages:	
1. Spectrum widening	
2. Decrease of resistance development risk	
3. Decrease of adverse reaction probability	
4. Increase in ATB efficacy	
Unsuitable combinations	
drugs with similar AE (nephrotoxicity, hepatotoxicity, …)	мимт
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ATBs in dentistry

Use

- prevention for risk patients (due to ADA)
 - artificial heart valves
 - · a history of ineffective endocarditis
 - · a cardiac transplant with developed valve problem
 - · some of congenital heart conditions

· in some types of stomatosurgeries

 for all dental procedures that involve manipulation of gingival tissue or the periapical region of the teeth, or perforation of the oral mucosa

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