

Antibacterial chemotherapeutics 2

1. β -lactame antibiotics

1.1 Penicillins

1.2 Cephalosporins

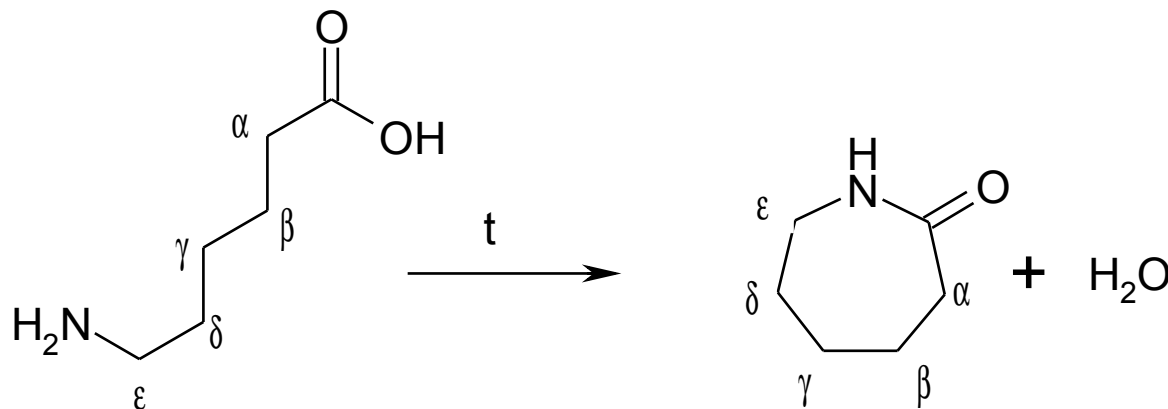
2. (Poly)peptide antibiotics

3. Macrolide antibiotics

4. Aminoglycoside antibiotics

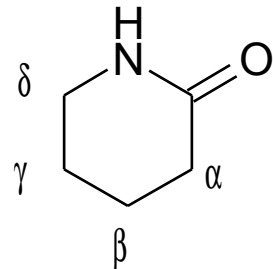
β -lactame antibiotics

Lactams = internal amides of amino acids

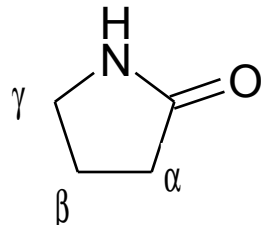


6-aminohexanoic acid
 ϵ -aminocaproic acid

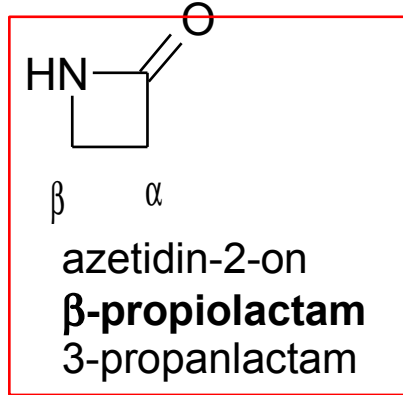
azepan-2-one
 ϵ -caprolactam
6-hexanlactam



piperidin-2-one
 δ -valerolactam
5-pentanlactam

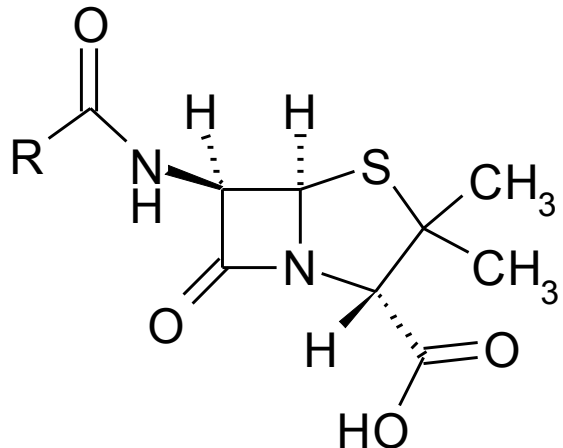


pyrrolidin-2-one
 γ -butyrolactam
4-butanlactam



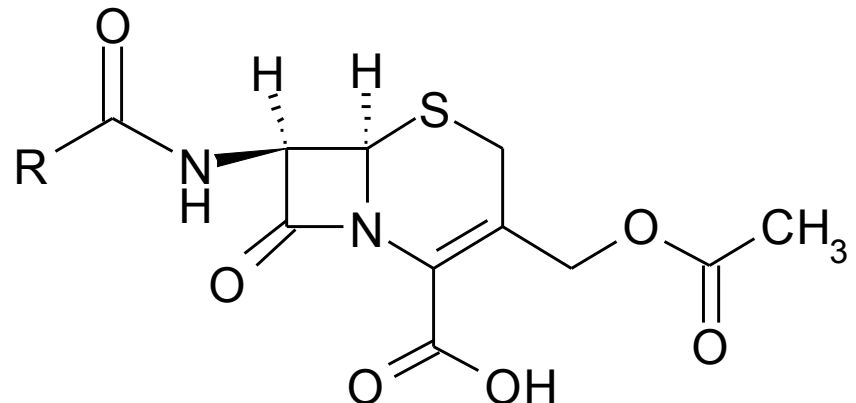
azetidin-2-one
 β -propiolactam
3-propanlactam

β -lactame antibiotics



Penicillins

N-acyl-6-aminopenicillanic acids



Cephalosporins

N-acyl-7-aminocephalosporanic acids

Mechanism of action

- inhibition of cell wall synthesis by binding to specific proteins

Penicilins History

World

1928(9) – **Alexander Fleming** – isolated a liquid concentrate inhibiting growth of bacteria from a mould of *Penicillium* species

1939 - 1943 Fleming, Florey, Chain & Johnson – isolation and constitution of penicillins

1945-Nobel prize for Fleming, Florey and Chain

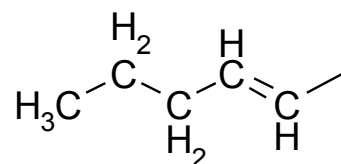
Czech territory

1943 – Málek, Fragner, Herold, Hais etc. – Mykoin BF 510

Penicillins

The initial „amorphous penicillin“ was a mixture of several compounds:

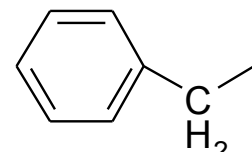
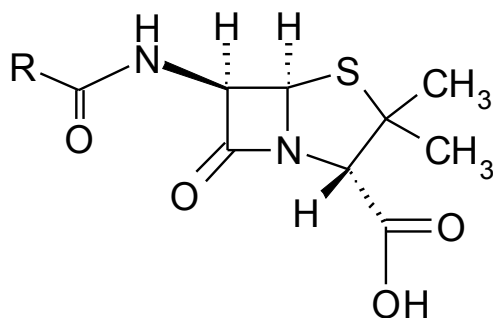
R



Penicillin

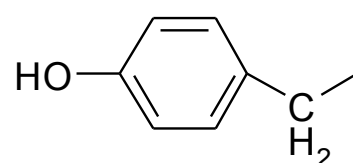
F

pentenyl-



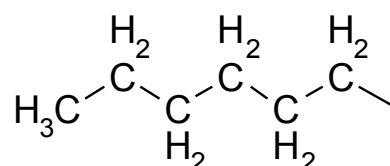
benzyl-

G



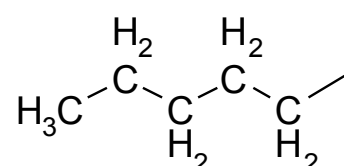
p-hydroxybenzyl-

X



heptyl-

K

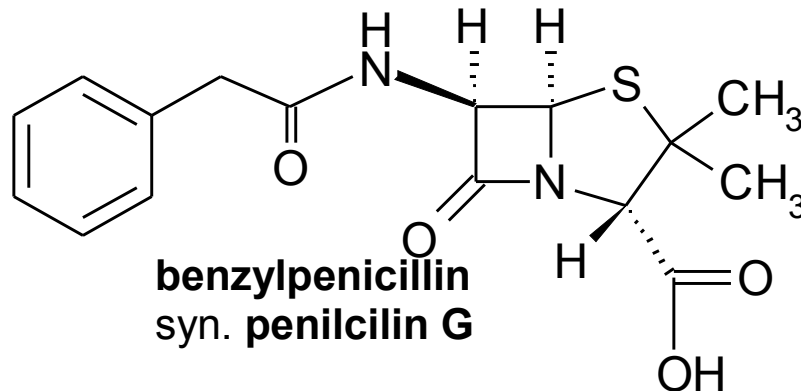


amyl-

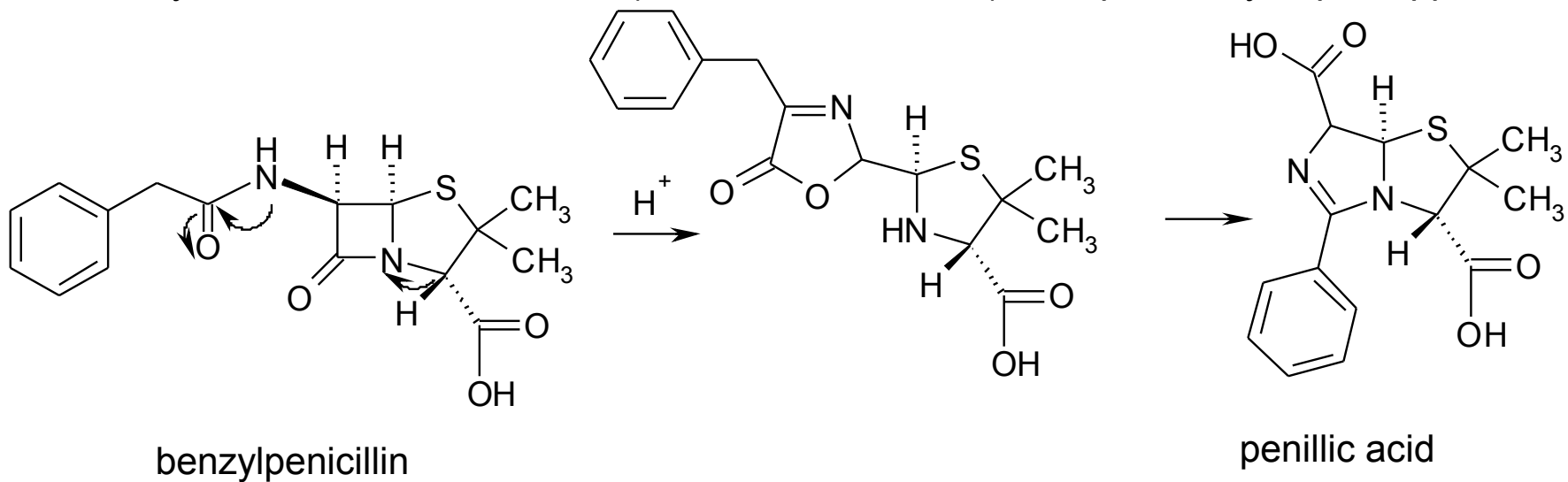
dihydro-F

Penicillins

Benzylpenicillin and its problems

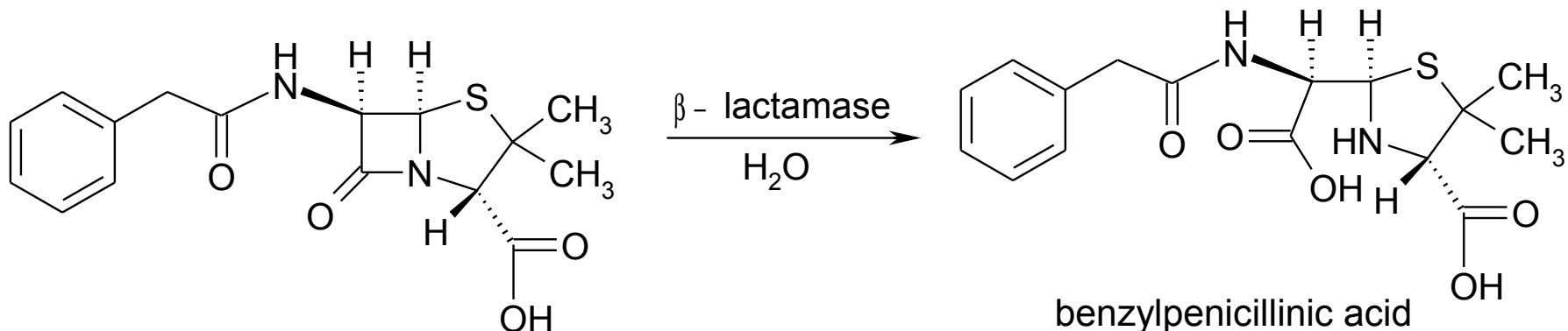


- production of benzylpenicillin by the mould ↑ by addition of phenylacetic acid into its broth
- Problems:
- weak binding to plasmatic proteins ⇒ fast excretion ⇒ frequent administration is necessary
 - instability in acid media of stomach (see reaction scheme) ⇒ impossibility of p.o. application



Penicillins **Benzylpenicillin** and its problems

3. Sensitivity to penicillinases (β -lactamases – enzymes catalysing hydrolytic cleavage of the β -lactame ring) – see the scheme

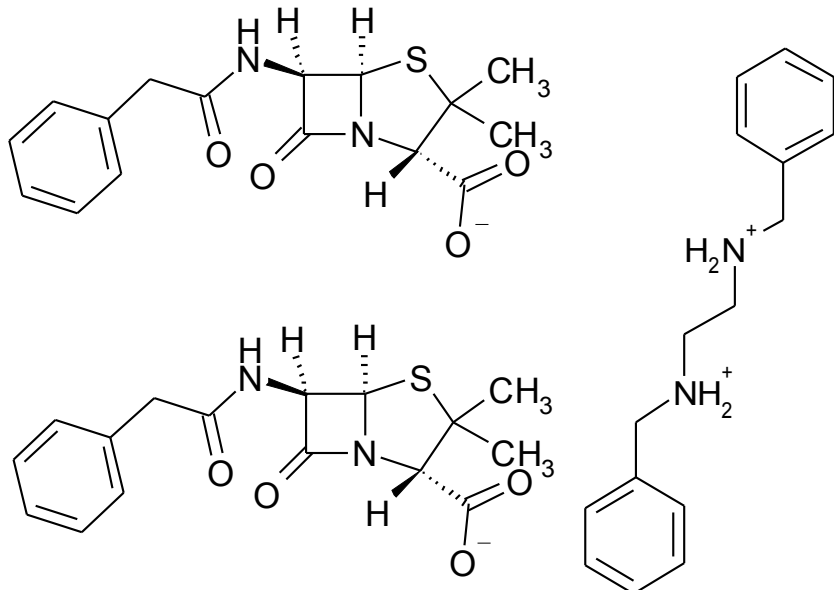


4. Rel. narrow spectrum – only G⁺ strains (*Streptococcus*, *Staphylococcus*, *Clostridium*, *Neisseria*, *Corynebacterium*, *Bacillus anthracis* ...)
5. Inducing allergies – anaphylactic shock – caused by 6-aminopenicillanic acid as the impurity – resolved by better purification (chromatography)

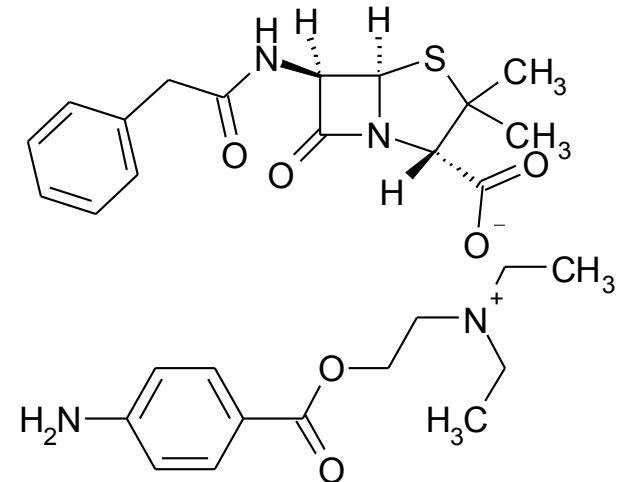
Penicillins

Resolving of benzylpenicillin problems

Ad 1. (necessity of frequent application) – poorly soluble salts with organic bases



benzathine benzylpenicillin
Pendepon® inj. sic.



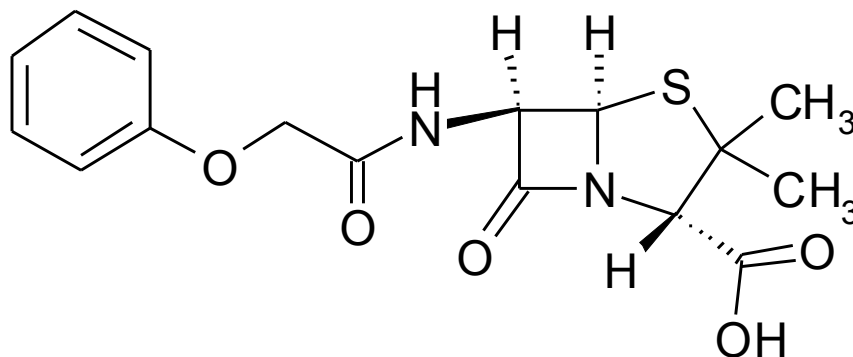
procaine benzylpenicillin
Prokain Penicilin G® Biotika inj. sic.

•depot (= long acting) forms for i.m. injections

Penicillins

Resolving of benzylpenicillin problems

Ad 2. – ↑ of stability in acid media



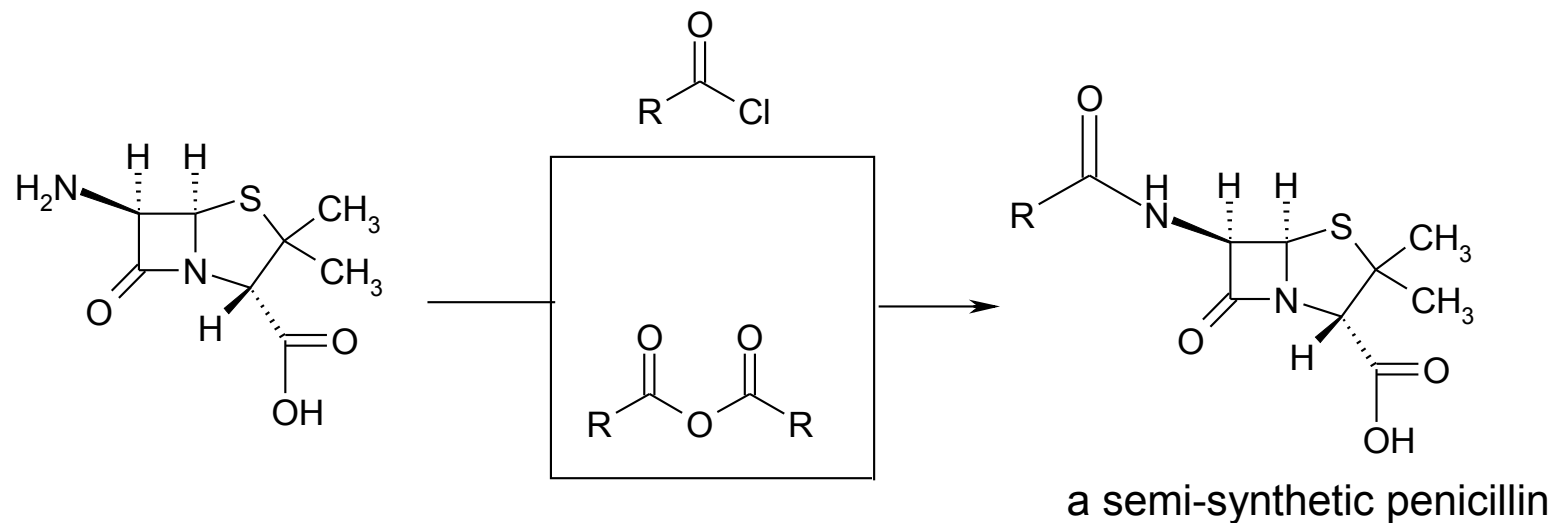
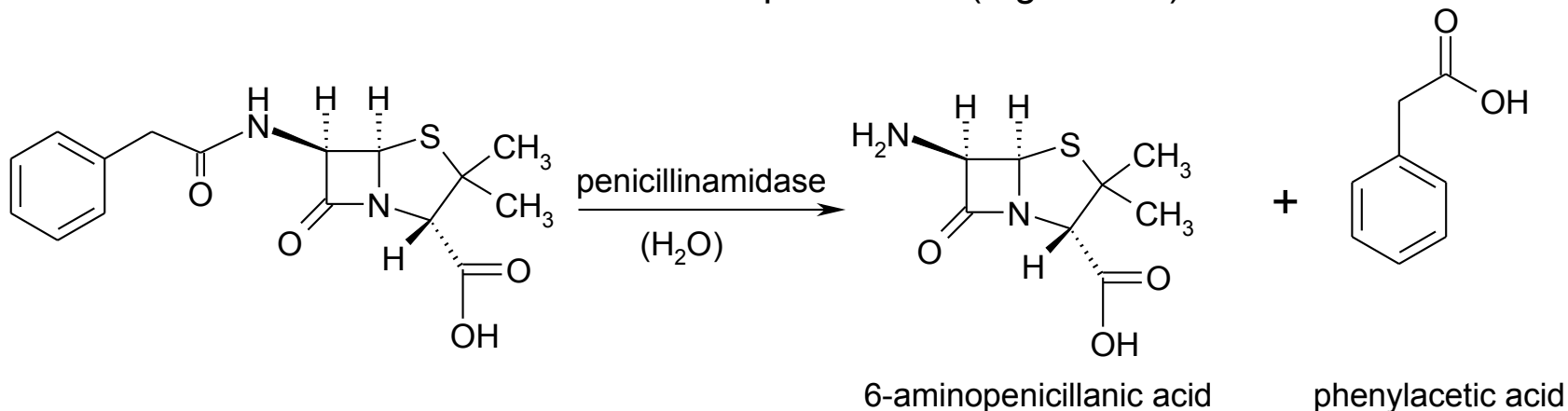
phenoxyethylpenicillin
syn. penicillin V

- acquired by addition of phenoxyacetic acid into the broth of the production strain
 - suitable for p.o. administration
V-Penicillin®, Ospen®

Penicillins

Overall resolving of benzylpenicillin problems – **semi-synthetic penicillins**

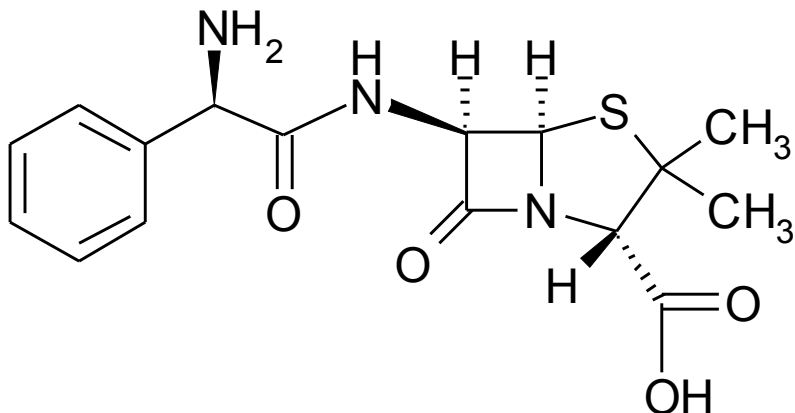
- **penicillinamidase (penicillinacylase)** – hydrolyzes acyclic amide bond, not β -lactame ring
 - used a microbe which produces it (e.g. *E. coli*)



Penicillins

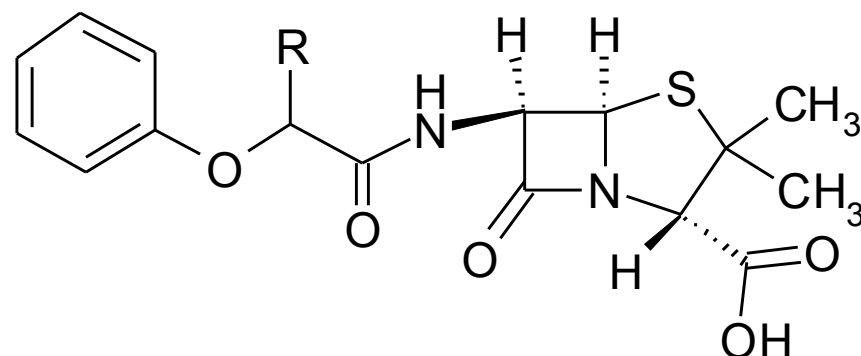
Mostly semi-synthetic penicillins stable in acid media

- stability against acids is increased by electron-donor substituents in N-acyl side chain (I+ or M+ effect)



ampicillin

Ampicilin® cps., inj sic.



R = -H

V-Penicillin® tbl., Ospen tbl. obd.

R = -CH₃

R = -CH₂CH₃

phenoxyethylpenicillin

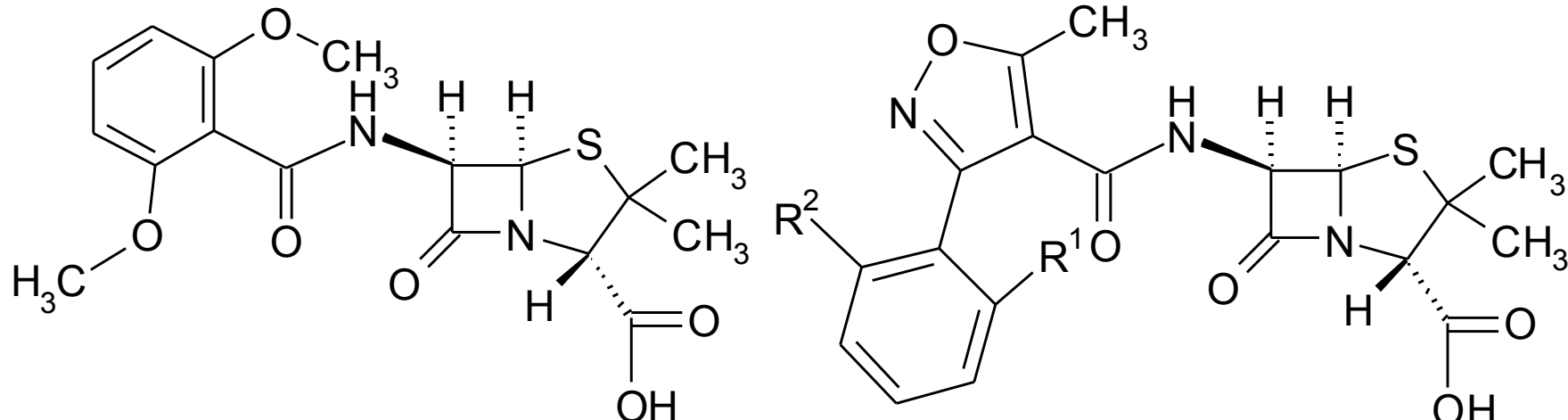
phenethicillin

propicillin

Penicillins

Semi-synthetic penicillins resistant to β -lactamases

- formed by acylation of amino group of 6-aminopenicillanic acid with bulky acyl rest; the lactame ring is then sterically hindered (\Rightarrow protected)



meticillin

$R^1 = R^2 = -H$ **oxacillin**

Prostaphlin® cps., inj. sic.

$R^1 = -Cl, R^2 = -H$ **cloxacillin**

$R^1 = R^2 = -Cl$ **dicloxacillin**

$R^1 = -Cl, R^2 = -F$ **flucloxacillin**

syn. floxacillin [USAN]

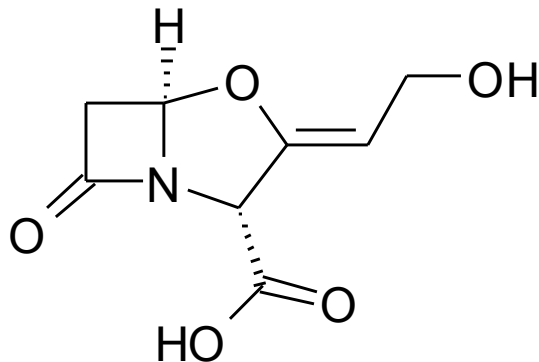
- resistant also to acid media; the resistance increases
oxacillin < cloxacillin < dicloxacillin = flucloxacillin

Penicillins

An alternative approach to ↑ of resistance to β -lactamases:

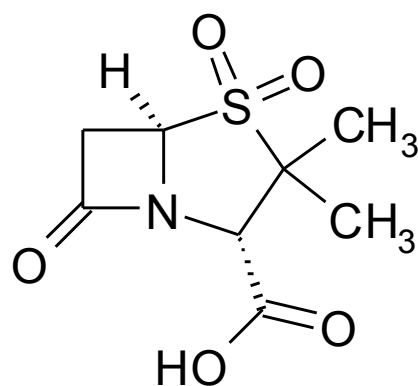
β -lactamases inhibitors

- compounds with β -lactam ring which binds to the enzyme active site with greater affinity and block this site
 - used in combination with penicillins



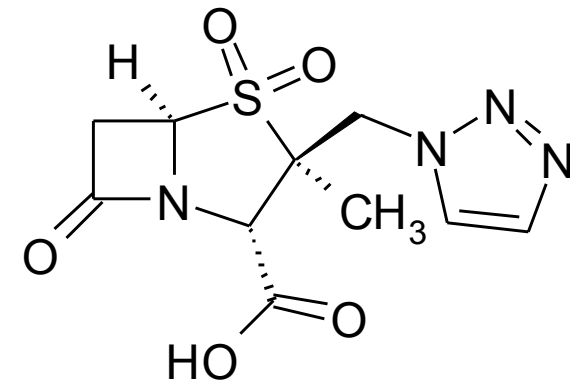
clavulanic acid

- isolated from *Streptomyces clavuligerus*
- + amoxicillin (= Amoxiklav®, Augmentin®)
- + ticarcillin (= Timentin® inj. sic.)



4,4-dioxopenicillanic acid

- sulbactam**
Betrion®
- + ampicillin (= Ampisucillin® inj. plv. sol.)

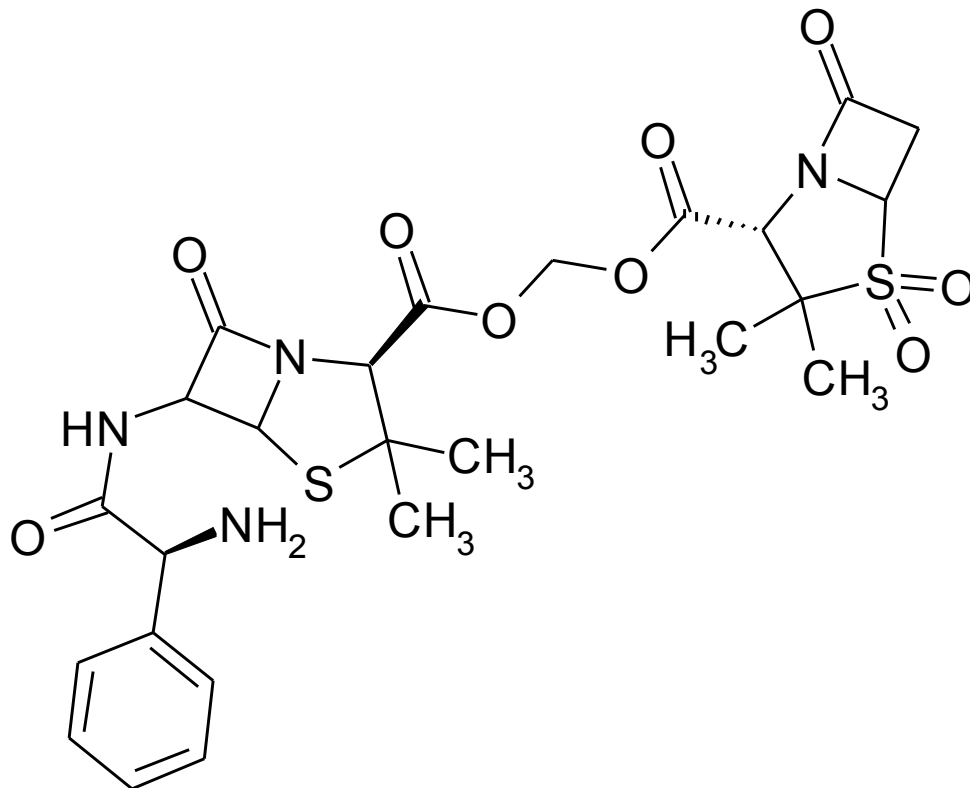


tazobactam

- + piperacillin (= Tazocin® inj. sic.)

Penicillins

A combination of a penicilline with a β -lactamase inhibitor in one molecule



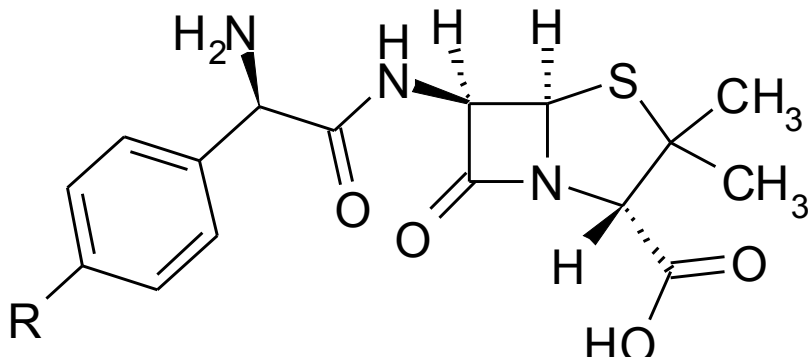
- a mixed ester of ampicillin and sulbactam with methanediol
 - a prodrug of both components
- sultamicillin**
- Bitamon® inj. sic., Unasyn® tbl. obd.

Penicillins

Penicillins with broadened spectrum

Ad 4. – introduction of a hydrophilic substituent to α -position of the acyl attached to amino group of 6-aminopenicillanic acid \Rightarrow **broadening of the antibacterial spectrum of penicillins also to G- strains**

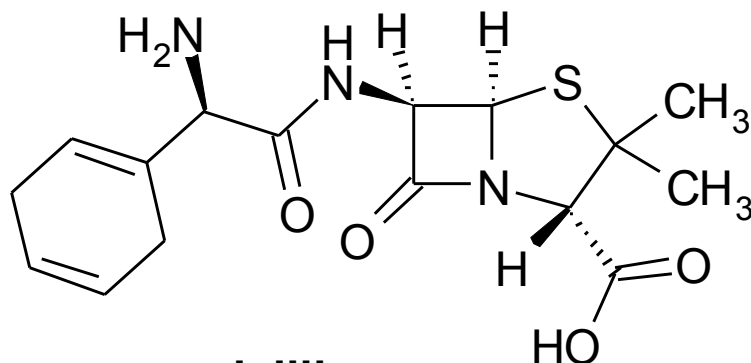
Compounds with free primary amino group



R = -H **ampicillin**

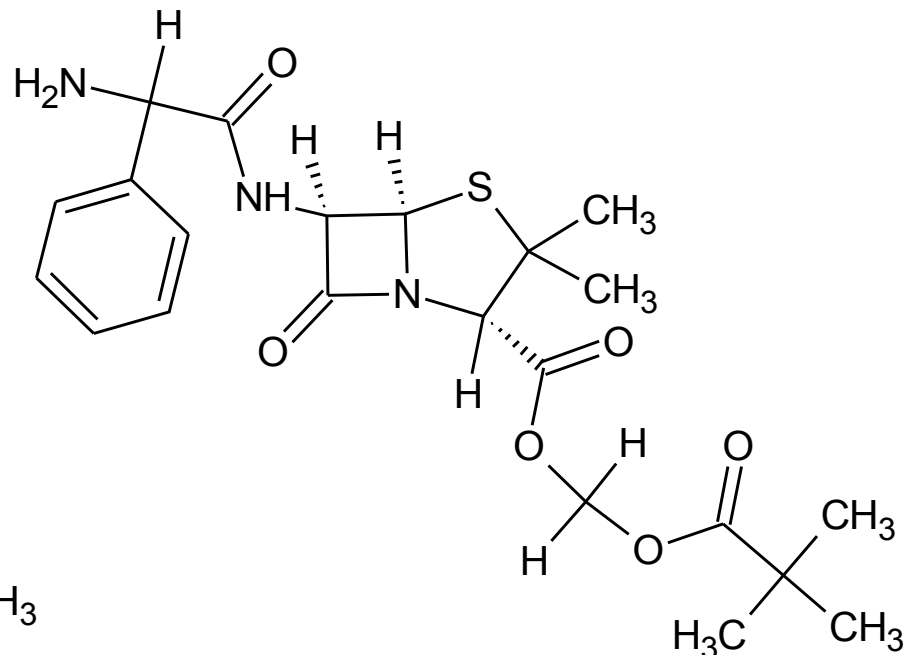
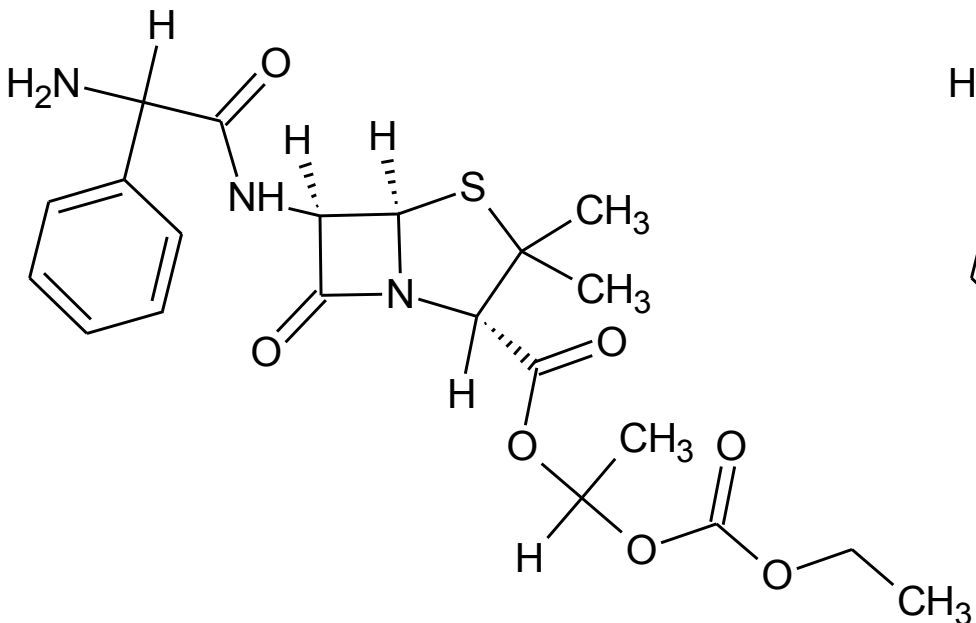
R= -OH **amoxycillin**

Amoclen[®], Amopen[®]



epicillin

Penicillins with broadened spectrum Ampicillin prodrugs



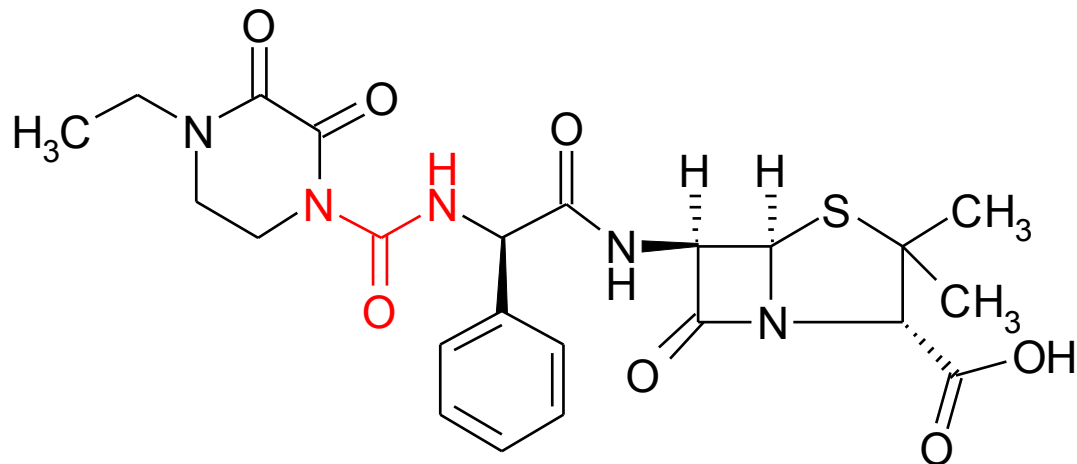
- hydrolyzed *in vivo* to ampicillin
- achieve significantly higher blood and tissue levels and attains peak blood levels more rapidly than equimolar doses of oral ampicillin
- more frequently used in veterinary (horses) than in human medicine
- models for design of prodrugs of cephalosporins

bacampicillin
ampicillin 1-
(ethoxycarbonyloxy)ethylester

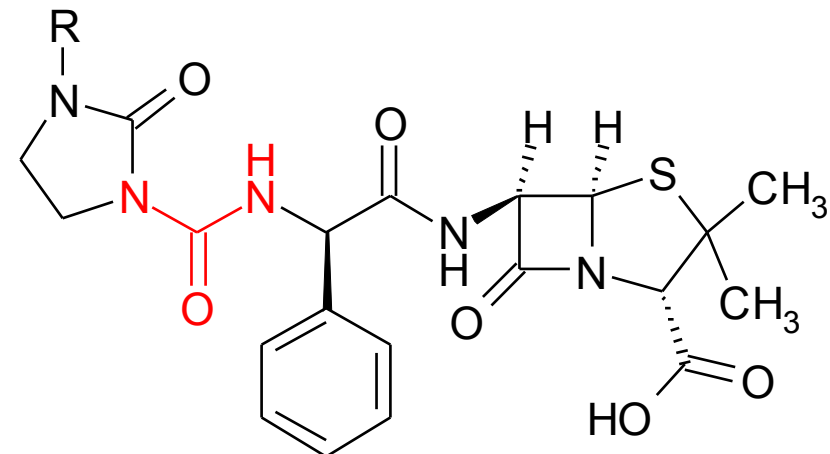
pivampicillin
ampicillin pivaloyloxymethylester
• successful in acute
exacerbations of chronic
bronchitis

Penicillins with broadened spectrum: ureidopenicillins

Compounds in which the amino group in α -position of the acyl is a part of urea moiety = **ureidopenicillins** = „anti-pseudomonas“ penicillins
•their spectrum includes *Pseudomonas aeruginosa*



piperacillin
Tazocin® inj. plv. sol.(+ tazobactam)



R = H-

azlocillin

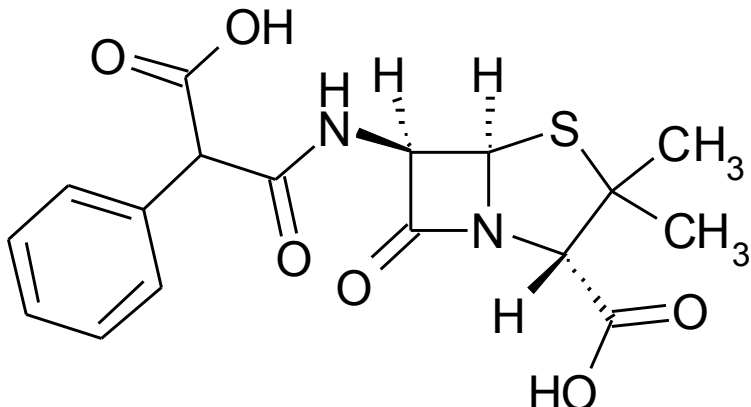
R = CH₃SO₂-

mezlocillin

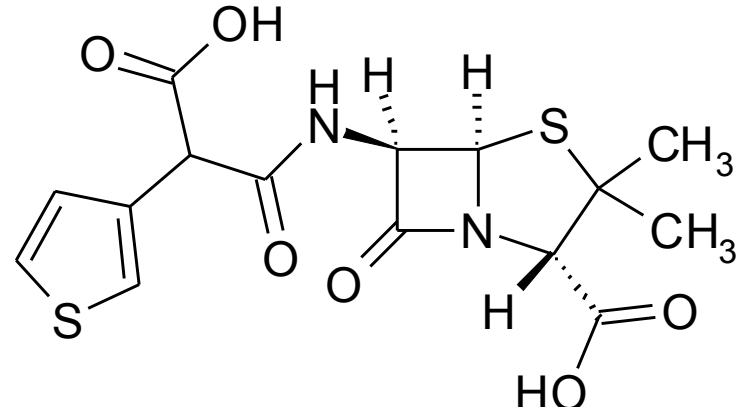
•serious infections including *otitis media*, CNS infections ...

Penicillins with broadened spectrum:

- compounds with the additional carboxyl in α -position of the acyl attached to amino group in position 6
 - in fact substituted malonic acids monoamides



carbenicillin



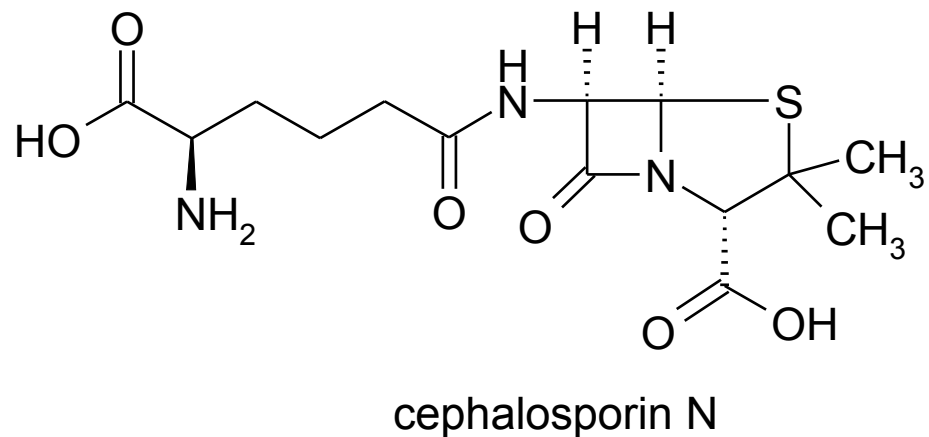
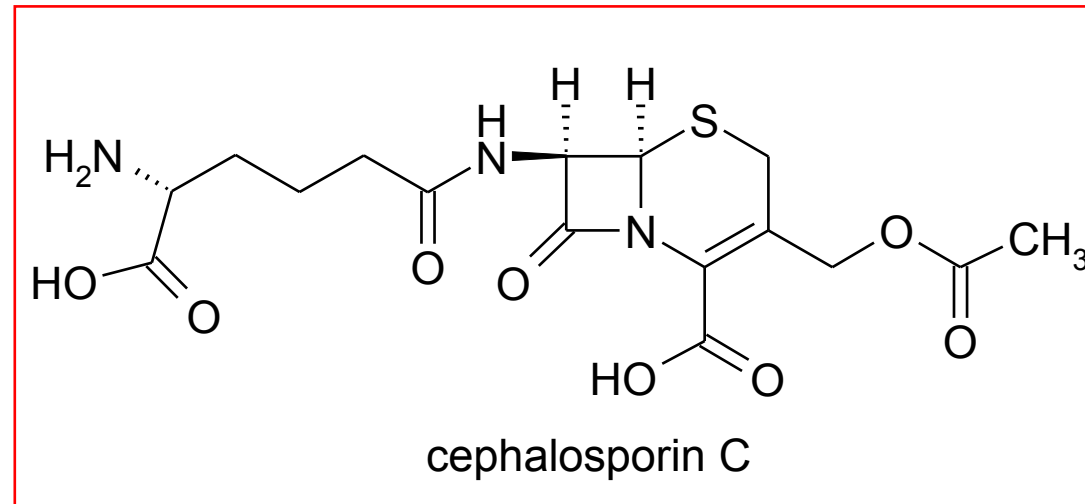
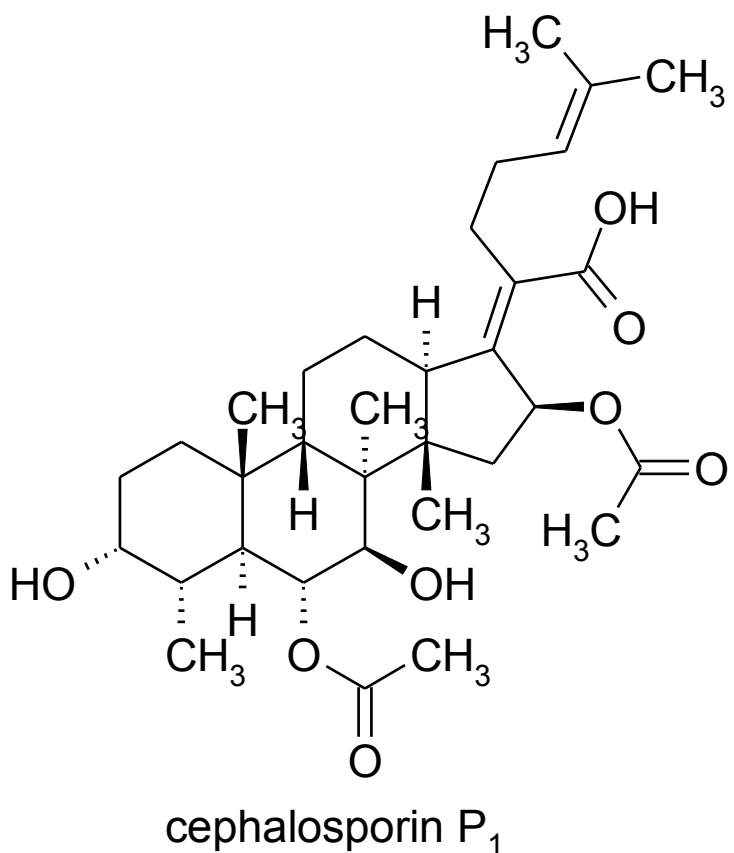
ticarcillin

Timentin® inj. sic. (+ clavulanic acid)
• kostní a kloubní infections of bones and
junctures (*Staphylococcus aureus*),
gynecological & abdominal infections ...

• ring analogy (benzene – thiophene)

Cephalosporins

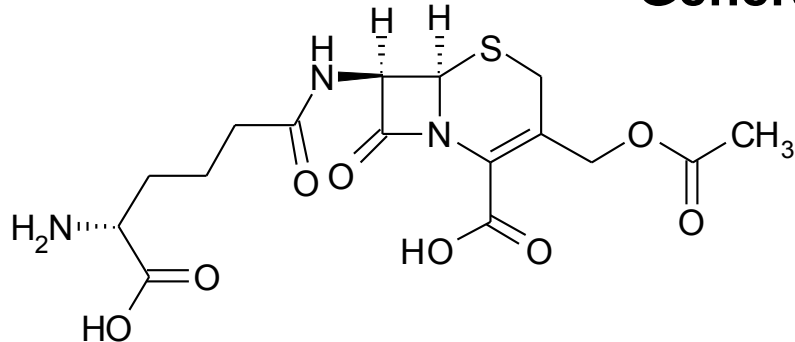
• fungi *Cephalosporium spp.* (1948)



... and other various structures

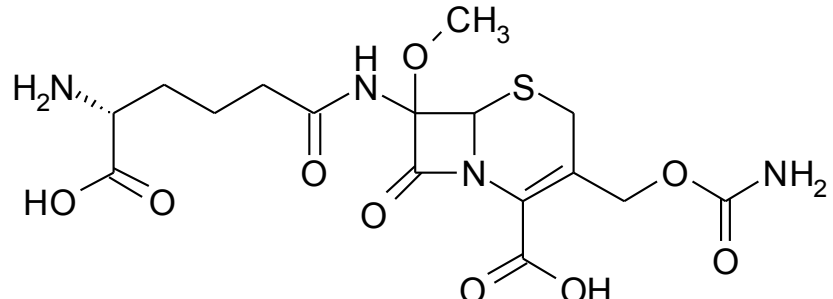
Cephalosporins

General structure



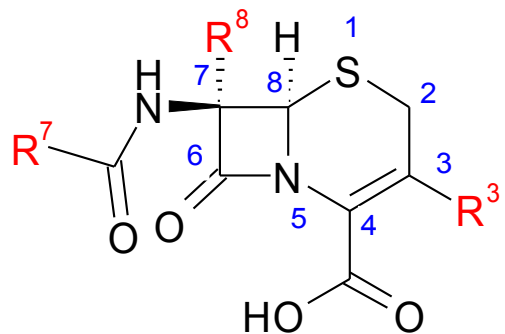
cephalosporin C

•isolated from *Cephalosporium spp.*



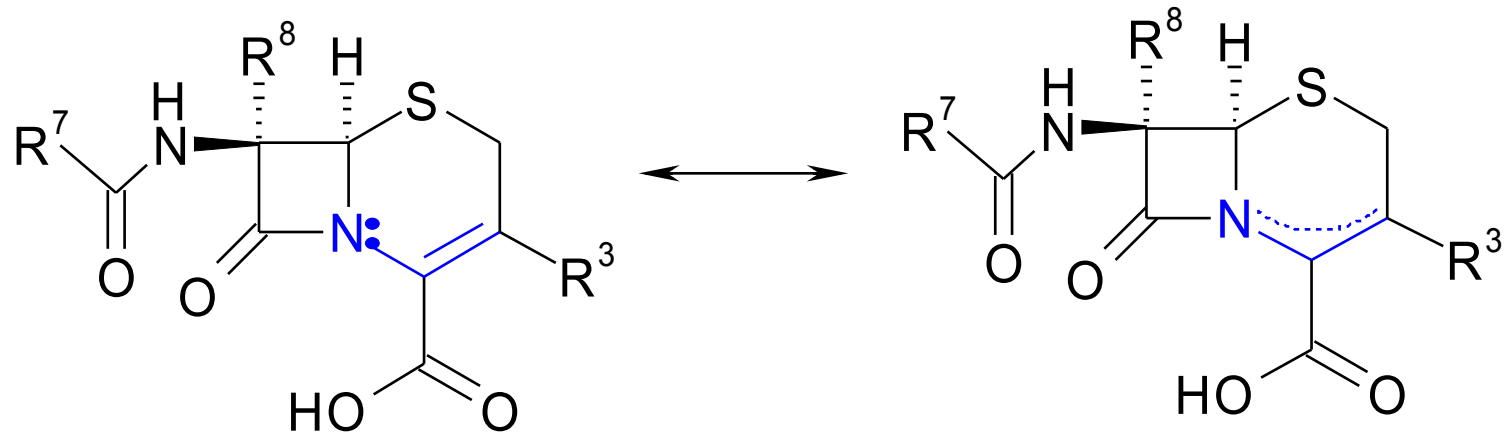
cephamycin C

•isolated from *Streptomyces lactadurans*



Cephalosporins

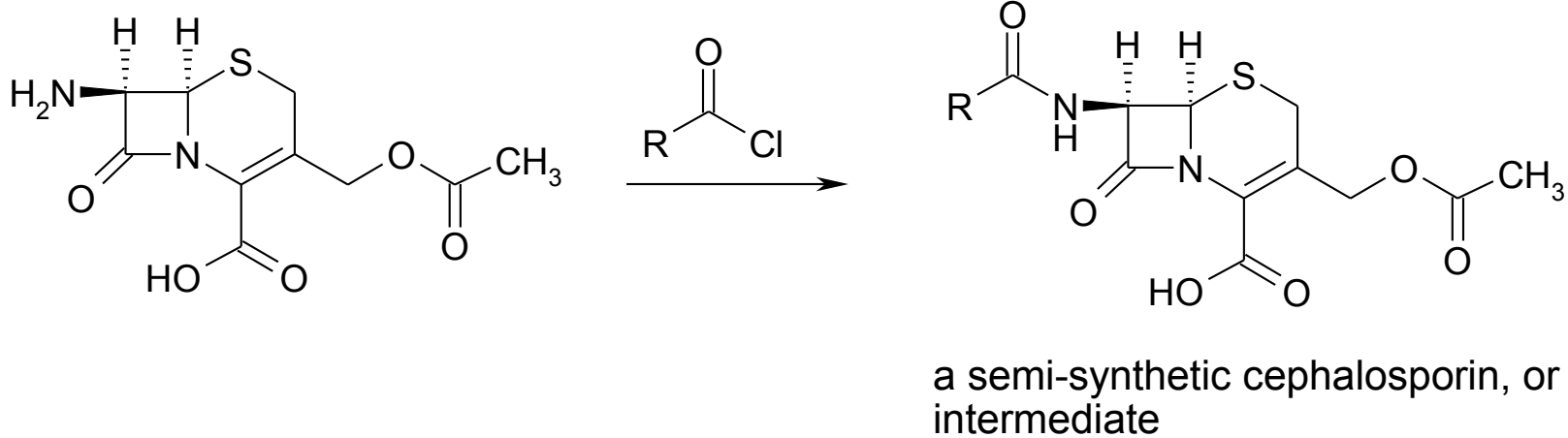
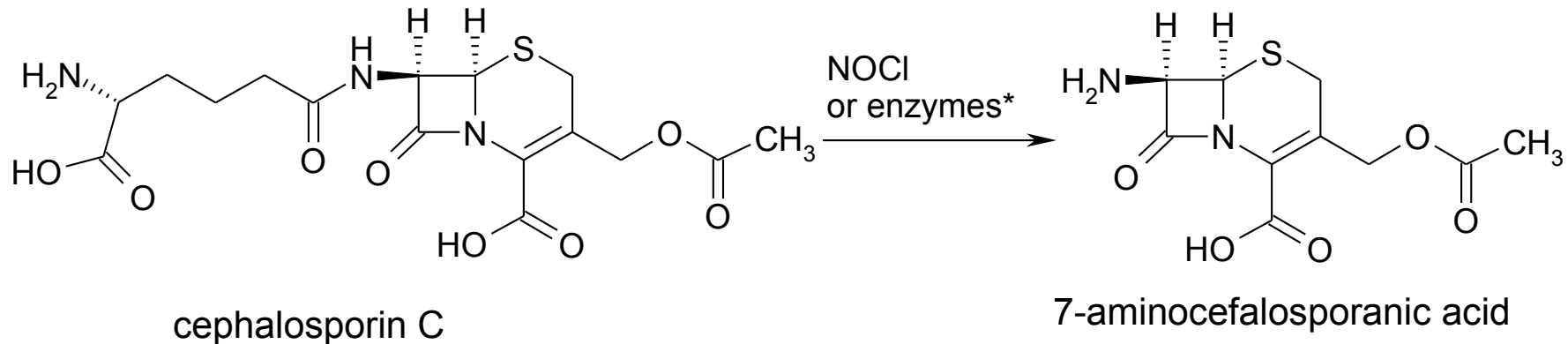
Properties



- electron pair on N5 is linked to conjugation with double bond $\Rightarrow \downarrow$ of electron density on N5 $\Rightarrow \downarrow$ of nucleophilicity of N5 \Rightarrow stability in acid media
- also \uparrow resistance to β -lactamases (cefalosporinases)

Cephalosporins

Compounds related to cephalosporin C, i.e. N-acylderivatives of 7-aminocephalosporanic acid.



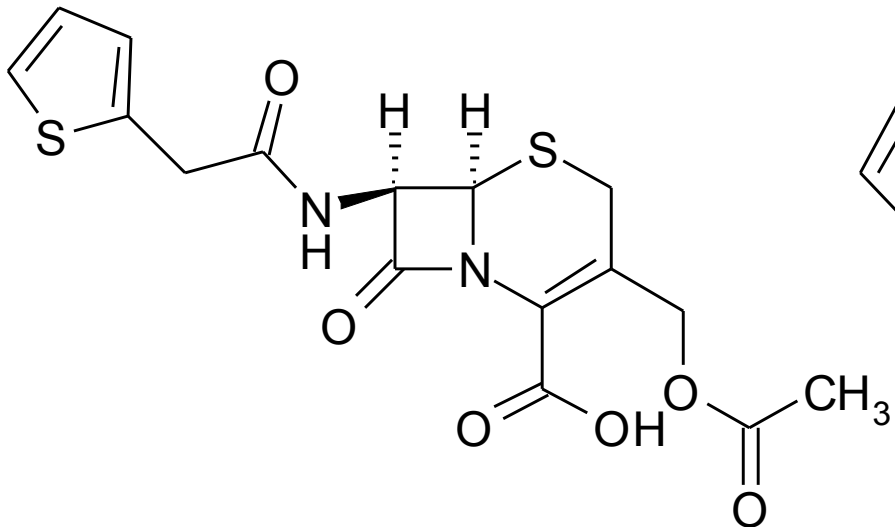
a semi-synthetic cephalosporin, or an intermediate

*glutarylacylase + D-amino acid oxidase

Cephalosporins

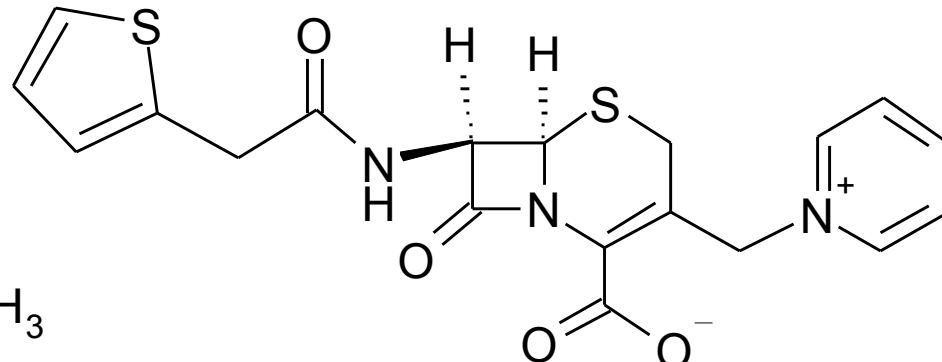
Compounds related to cephalosporin C, i.e. N-acylderivatives of 7-aminocephalosporanic acid

1st generation: for parenteral administration only (not absorbed from GIT)



cephalotin

Cefalotin® Biotika inj. sic.

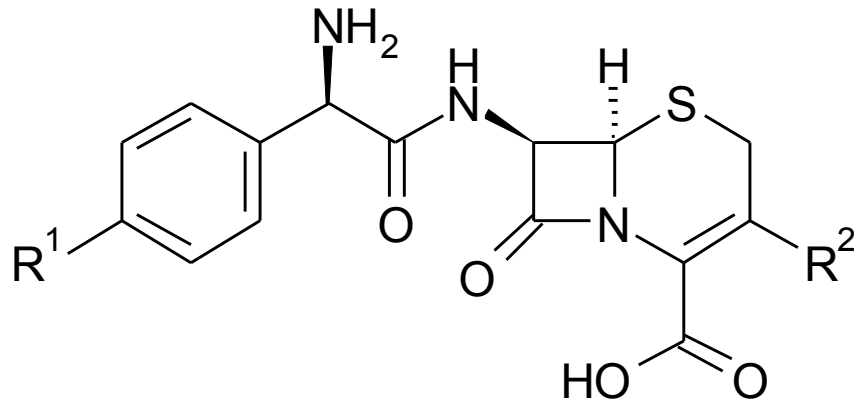


cefaloridin

Cephalosporins

Compounds related to cephalosporin C, i.e. N-acylderivatives of 7-aminocephalosporanic acid

2nd generation: for oral administration



R¹= -H, R²= -CH₃

cefalexin

Cefaclen® cps.

R¹= -OH, R²= -CH₃

cefadroxil

Biodroxil® tbl. obd.

R¹= -H, R²=Cl

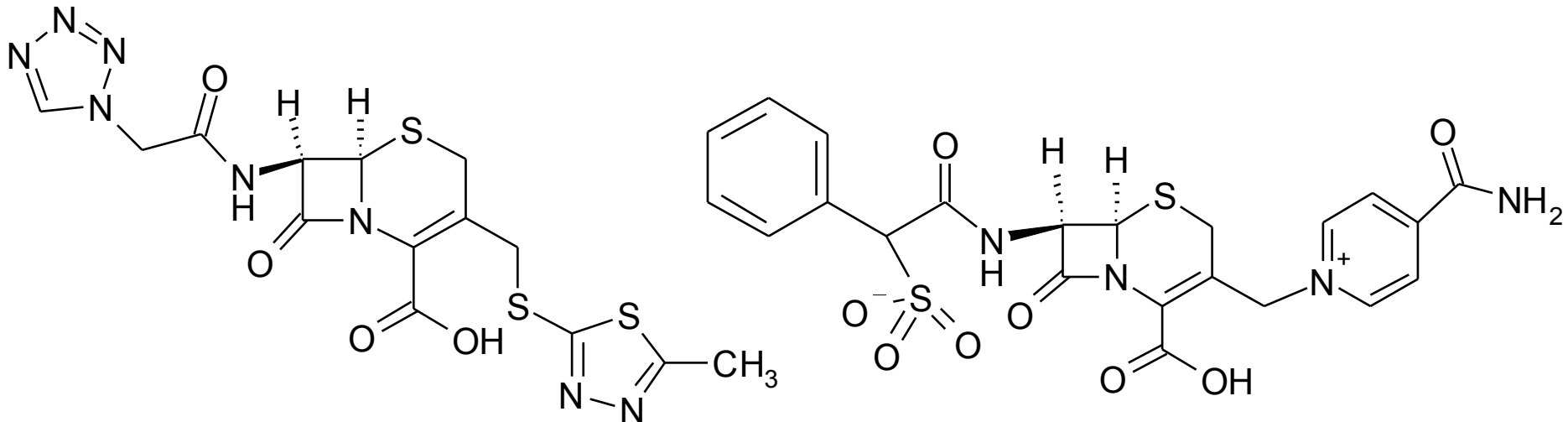
cefaklor

Ceclor® cps.

Cephalosporins

Compounds related to cephalosporin C, i.e. N-acylderivatives of 7-aminocephalosporanic acid

2nd generation: for parenteral use but with ↑ effect to G⁻, ↑ resistance to β-lactamases



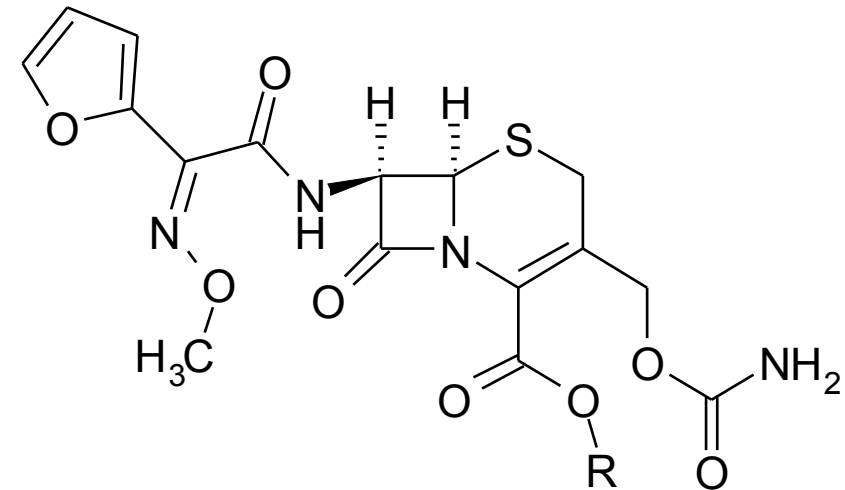
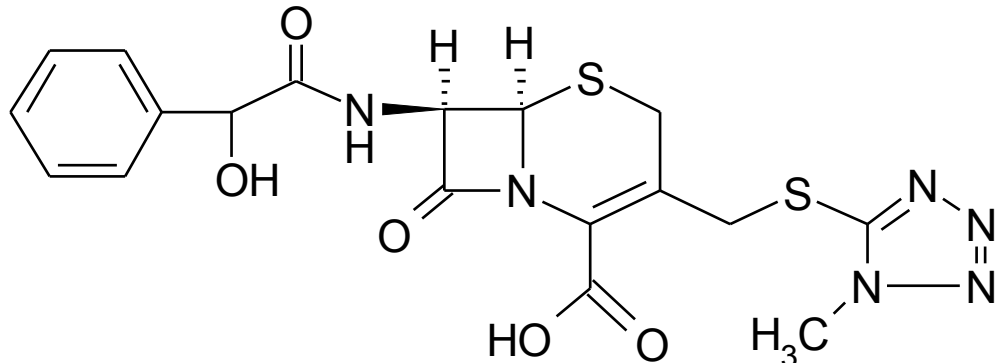
cefazolin
Kefzol® inj. sic.

cefsulodin
•*Pseudomonas*

Cephalosporins

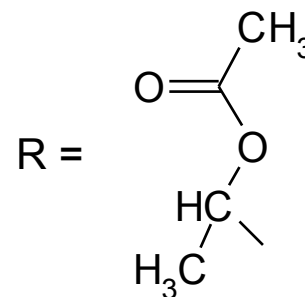
Compounds related to cephalosporin C, i.e. N-acylderivatives of 7-aminocephalosporanic acid

2nd generation: for both parenteral and p.o. administration, very resistant to β -lactamase



R = H-

Cefuroxime® tbl.

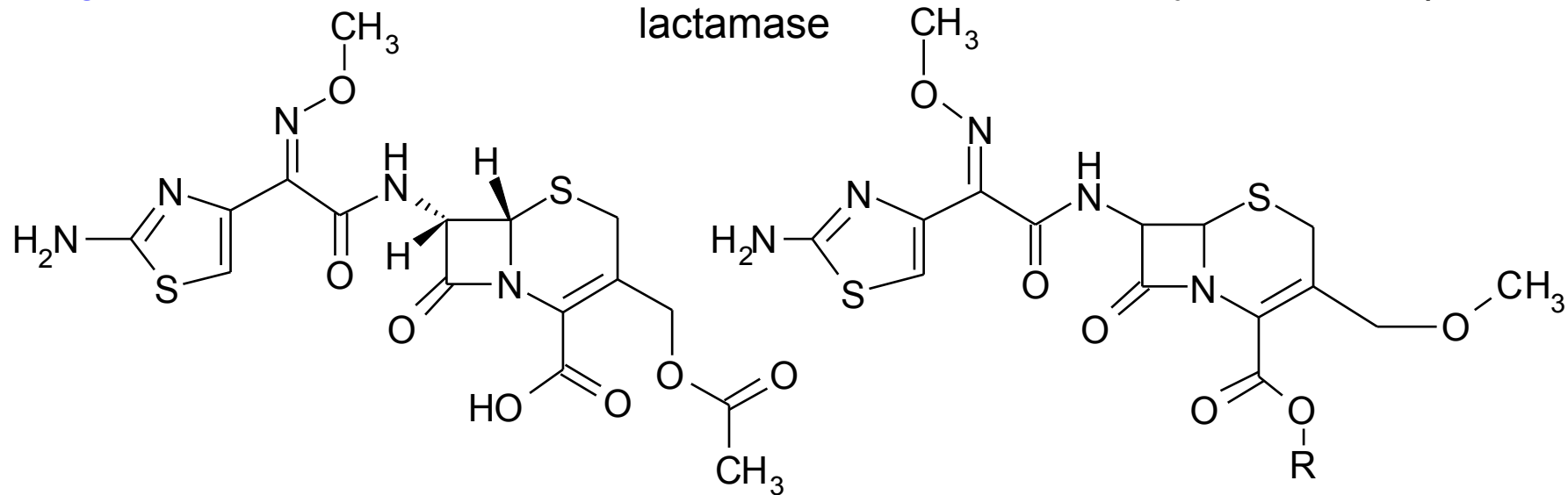


cefuroxime axetil
Zinnat® tbl. obd.

Cephalosporins

Compounds related to cephalosporin C, i.e. N-acylderivatives of 7-aminocephalosporanic acid

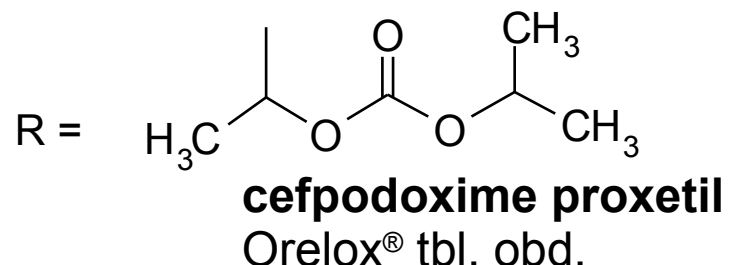
3rd generation: for both parenteral and p.o. administration, very resistant to β -lactamase



cefotaxime

Claforan® inj. sic.

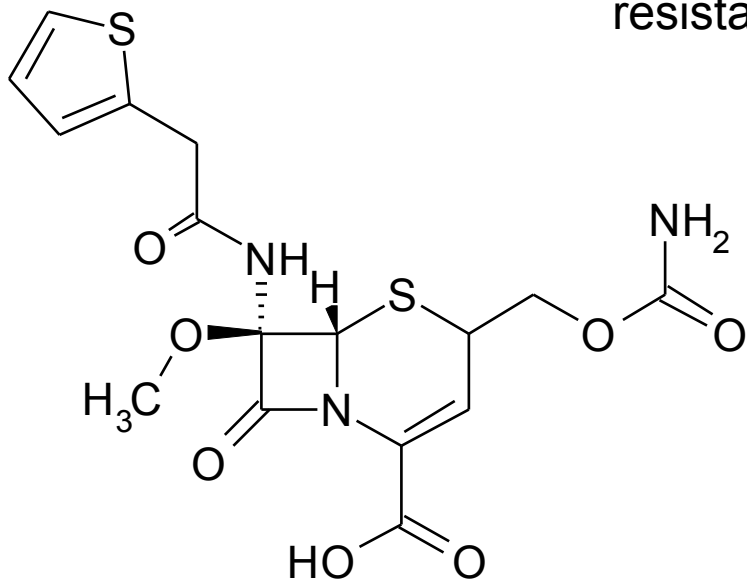
R = H- **cefpodoxime**



Cephalosporins

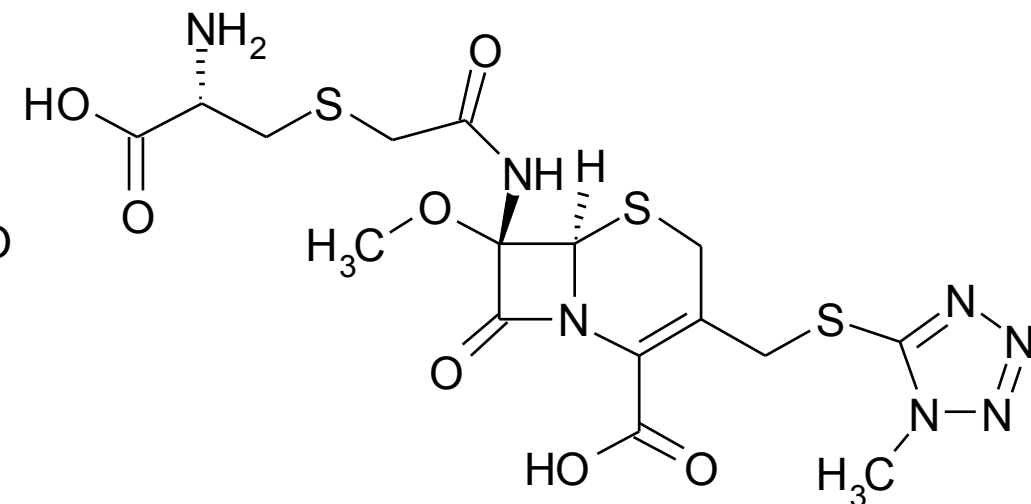
Compounds related to **cephamycin C**, i.e. N-acylderivatives of **7-methoxy-7-aminocephalosporanic acid**

„New class = 4th generation“ – for both parenteral and p.o administration –
resistant to β -lactamase



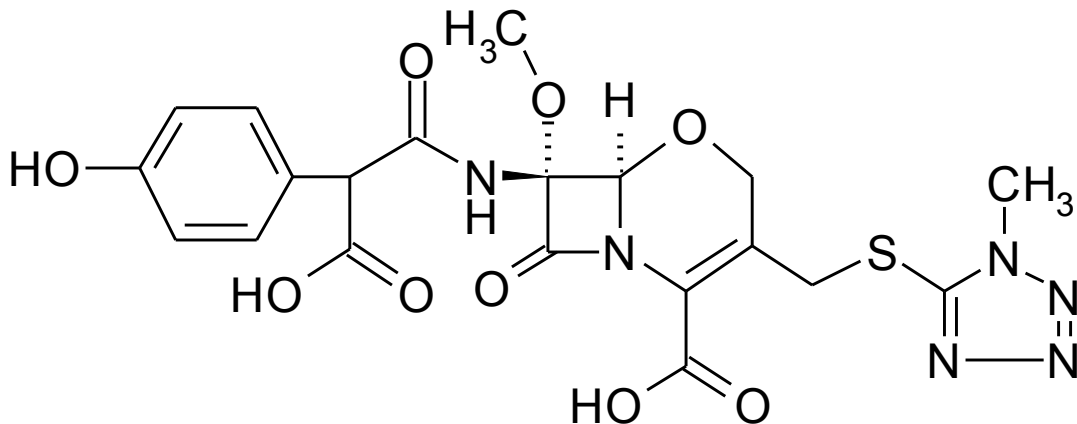
cefoxitin

Mefoxin® inj. sic.



cefminox

Cephalosporin analogues

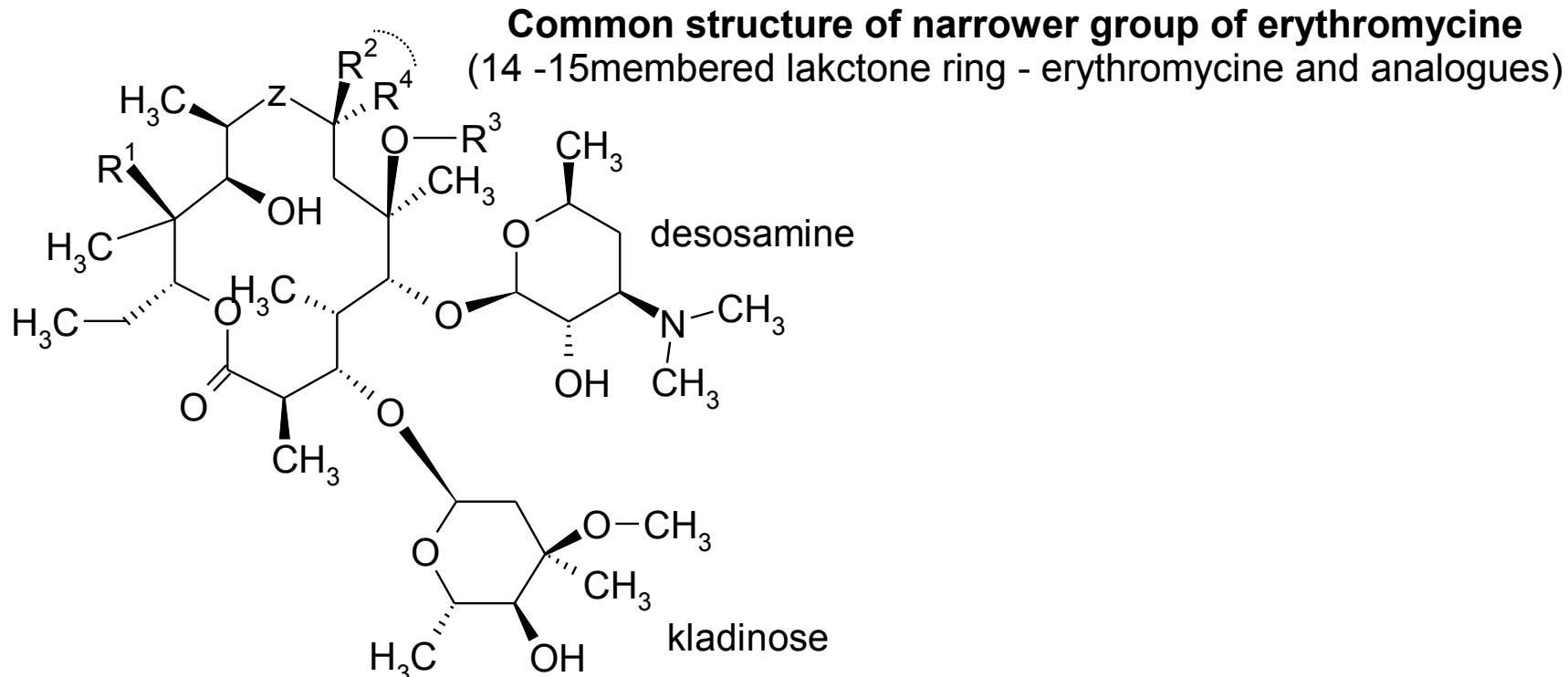


moxalactam

- dihydroooxazine derivative related to 4th generation of cephalosporins
- developed especially for treatment of meningitis (crosses the blood-brain barrier) and anaerobic infections

Macrolides

- makrocyclic lactones with 10 – 40membered ring with 1 aminomonosaccharide and 1 „neutral“ monosacharide which can have an additional aminosaccharide attached
- 1st group (with larger ring)- natamycine, nystatine, amphotericine B – see antimycotics
- 2nd group – **erythromycine group** (erythromycine and its analogues, spiramycin, tylosine)



R¹= -OH, -H

Z = >C=O, -CH₂N(CH₃)₂, >C=N-O-CH₃, >C=NOCH₂OCH₂CH₂OCH₃

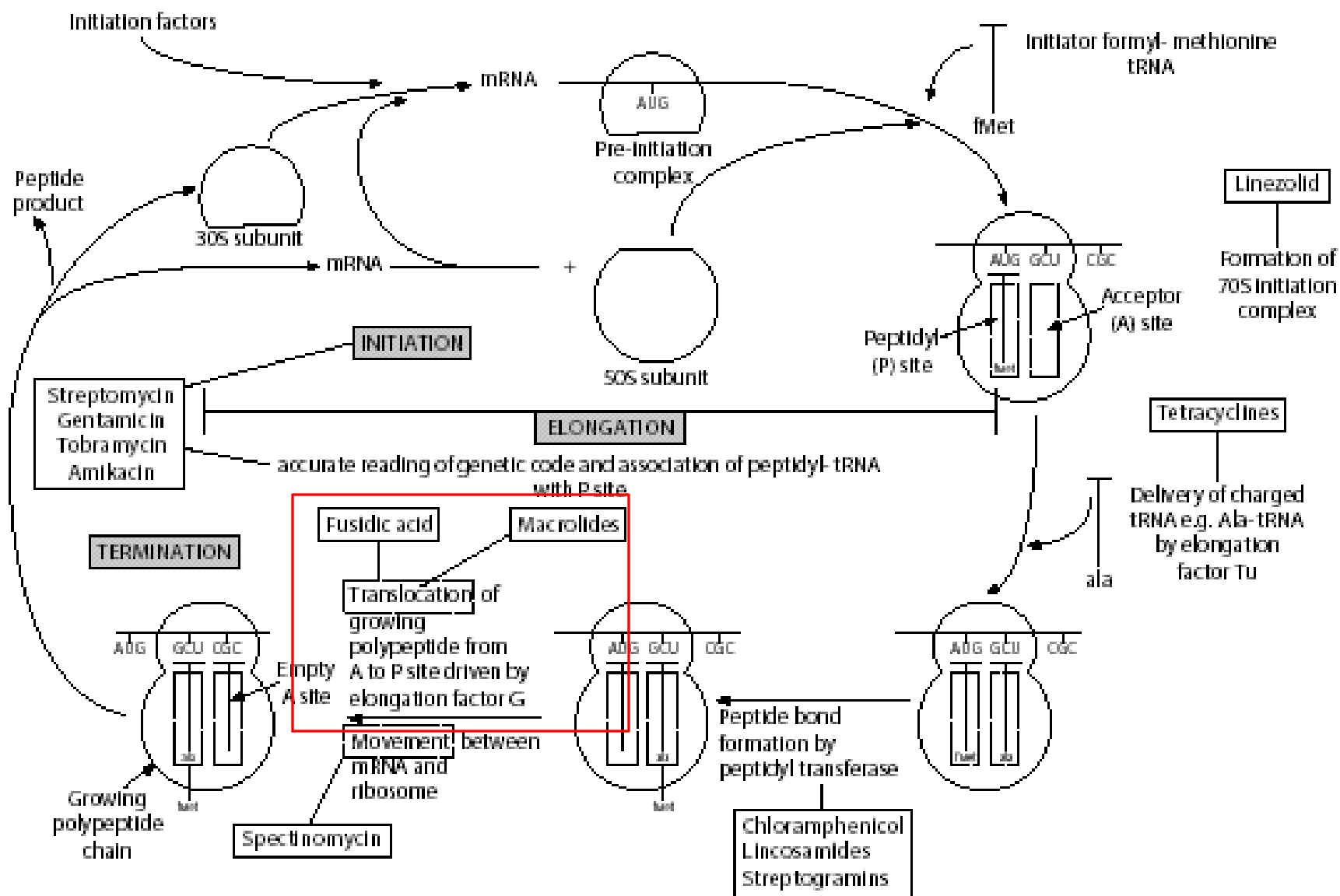
R²= -H, -F

R³= -H, -CH₃

R⁴= -CH₃ or R² + R⁴= oxirane

Macrolides

Site & mechanism of action



Macrolides

Site and mechanism of action

- **Proteosynthesis inhibition**
 - act at 50S ribosome subunit
- inhibit the translocation of growing peptide from acceptor (A) to peptide (P) site
 - **bacteriostatic effect**

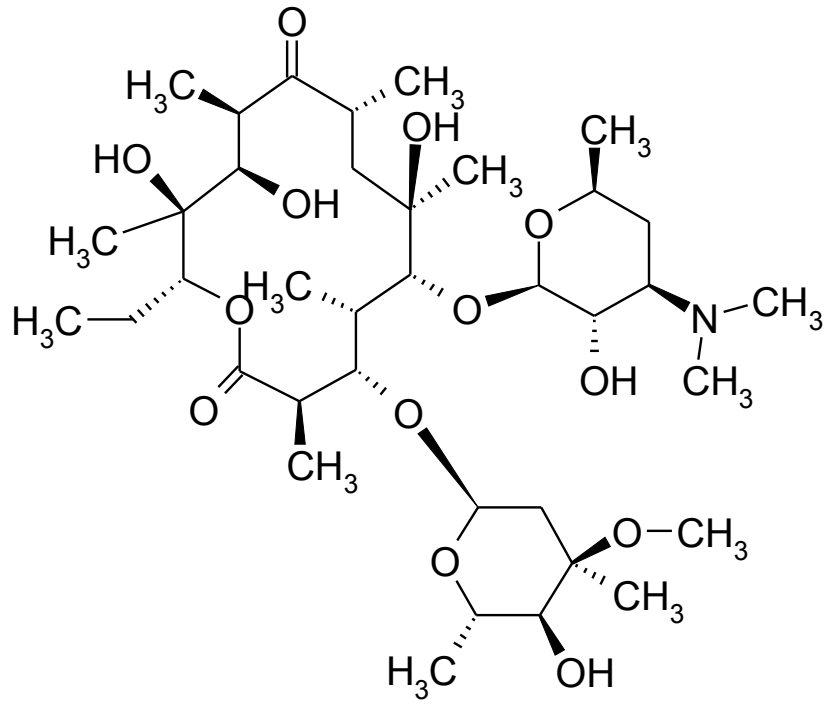
Spectrum:

- both G⁺ and G⁻

Neisseria, Haemophilus, Brahmanella, Legionella ...

Macrolides

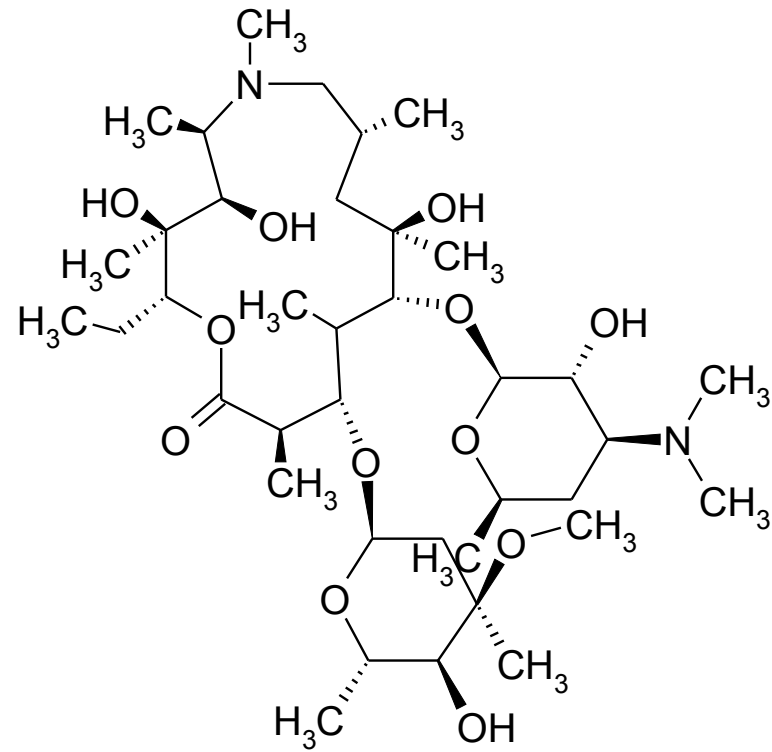
Erythromycine and its analogues



erythromycine

- isolated 1952 from *Streptomyces erythreus*

- poor biological availability \Rightarrow lipophilic salts (stearate, ethylsuccinate ...)
- external form (lotions ...) – treatment of *acne vulgaris*

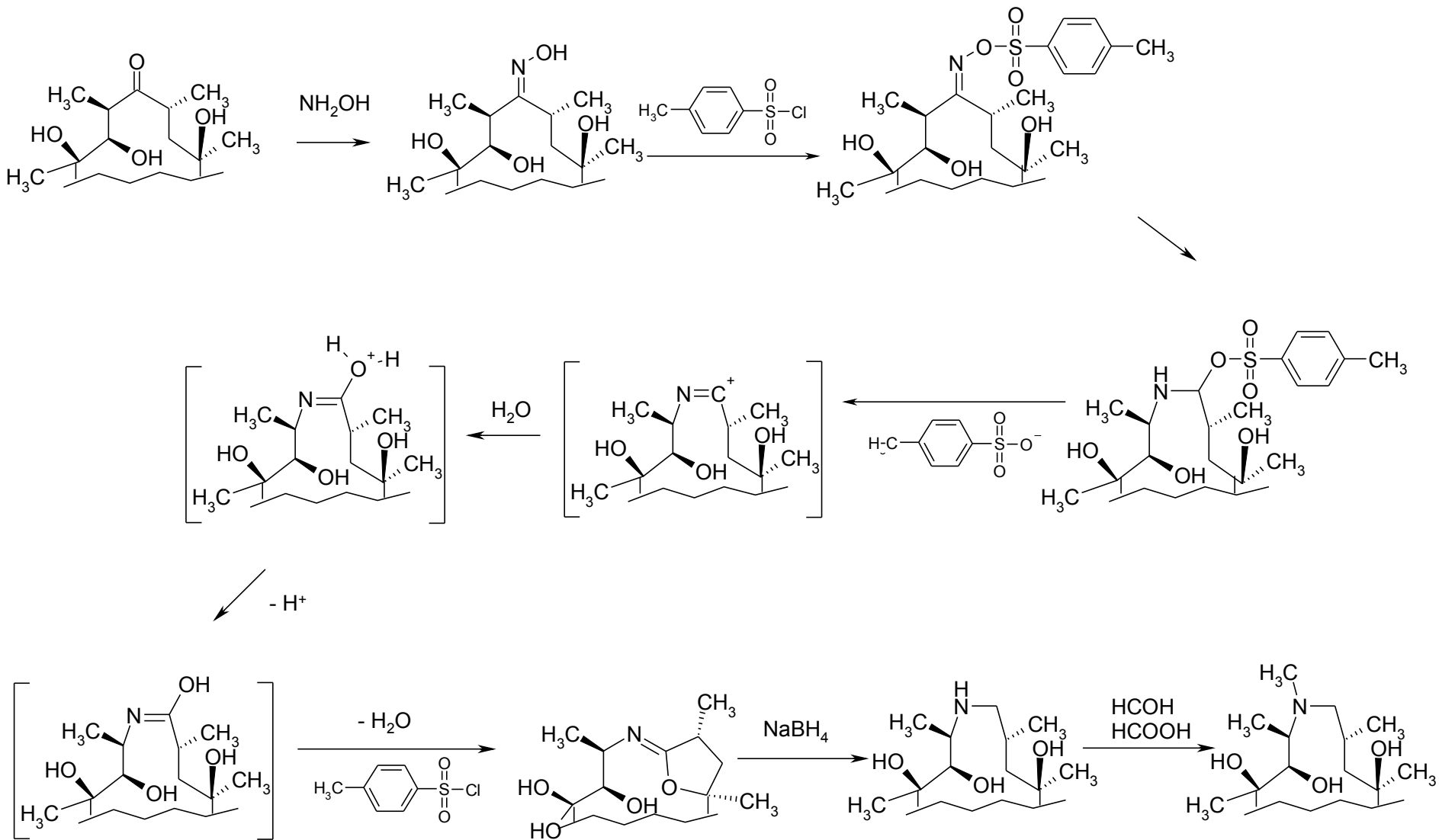


azithromycine

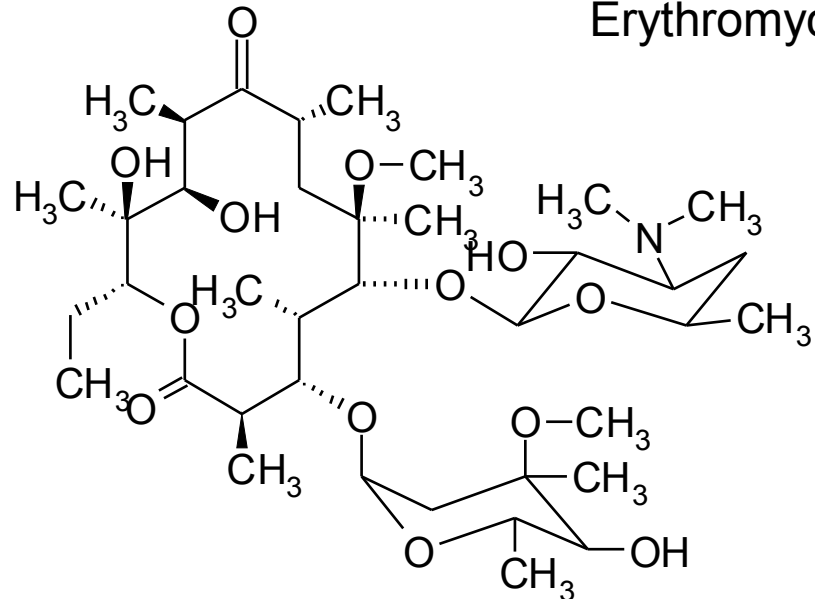
- semi-synthetic compound Sumamed® tbl. obd.

Macrolides

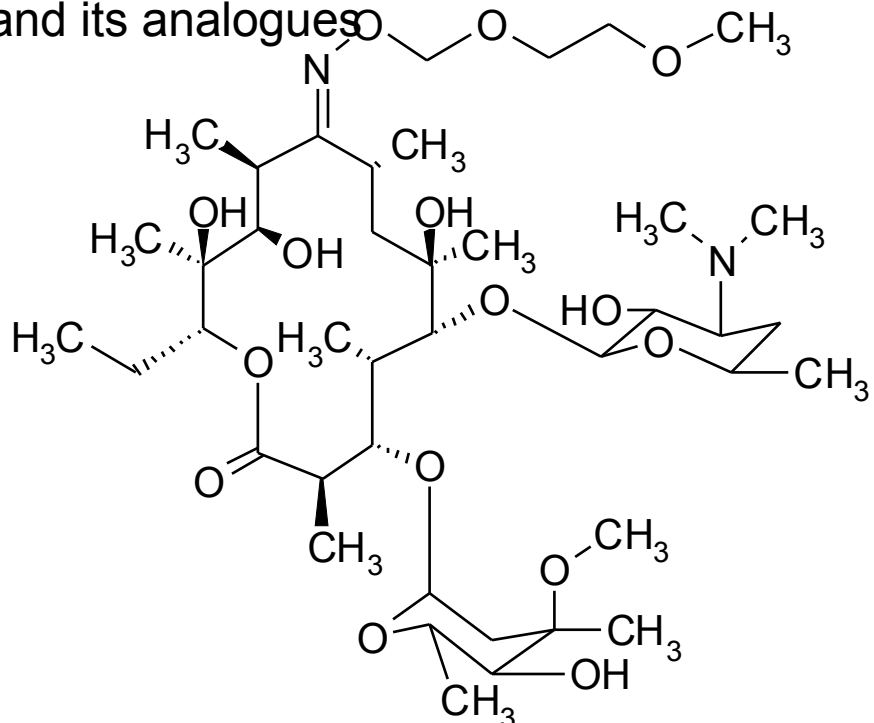
Synthesis of azithromycine from erythromycine



Macrolides



Erythromycine and its analogues



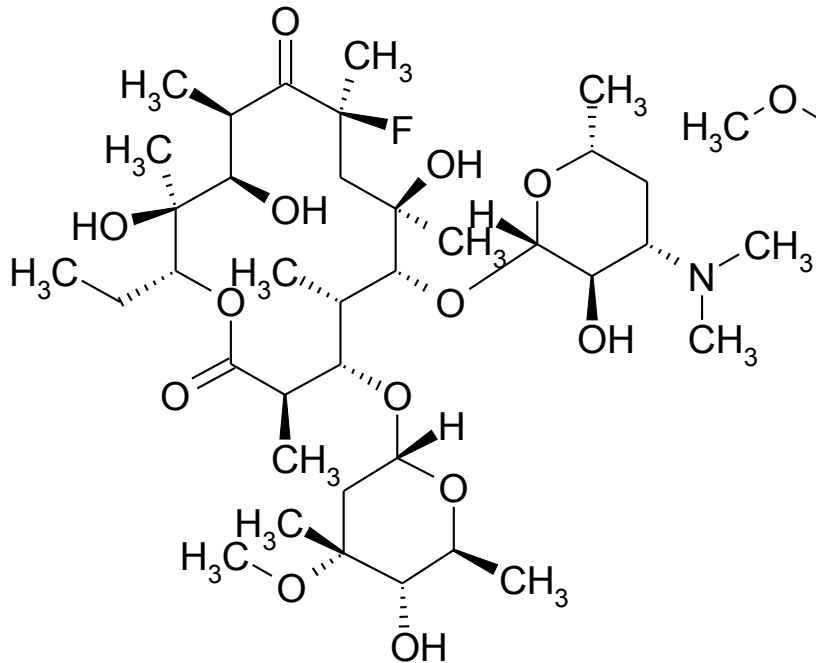
6-O-methylerythromycine
clarithromycine

•also some strains of *Mycobacterium avium*
Klacid® tbl. obd.

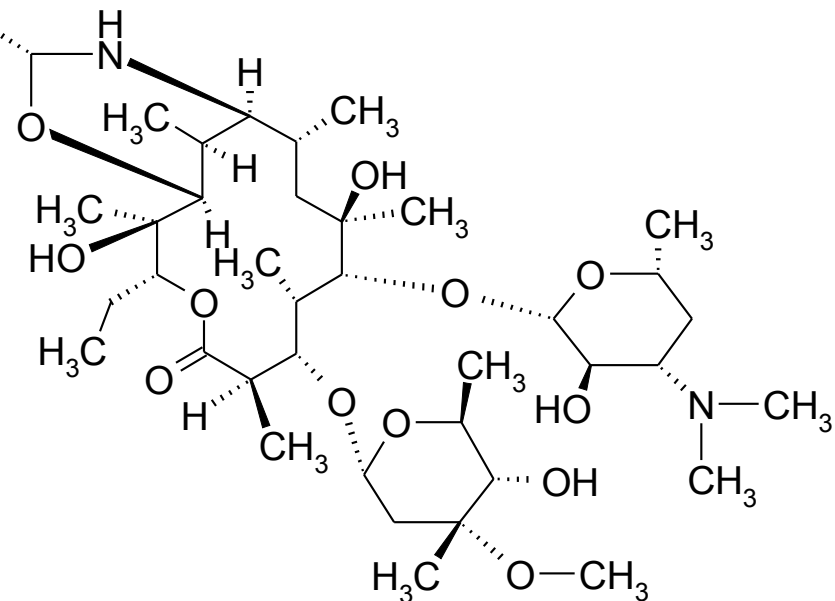
roxithromycine
Rulid® tbl.

Macrolides

Erythromycine and its analogues



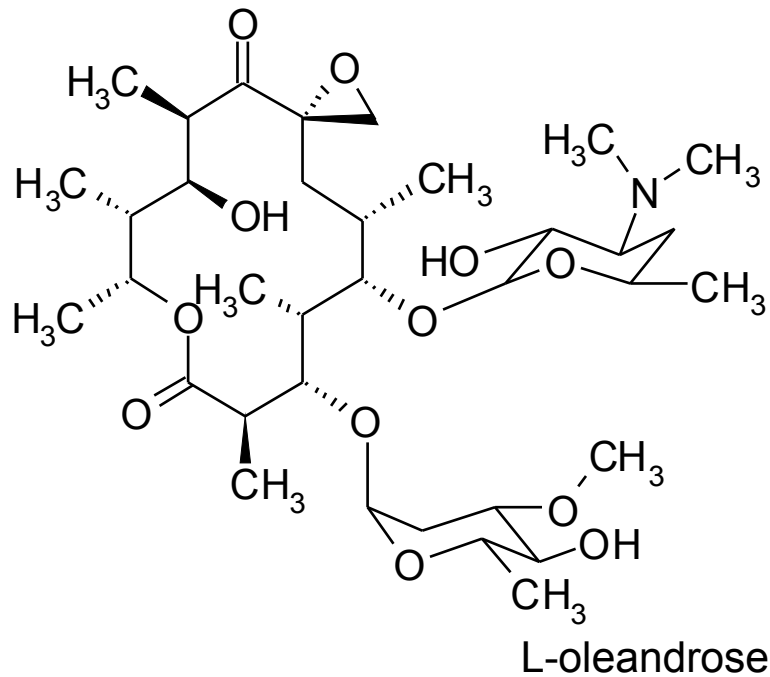
**8-fluoroerythromycin
flurithromycine**



dirithromycin

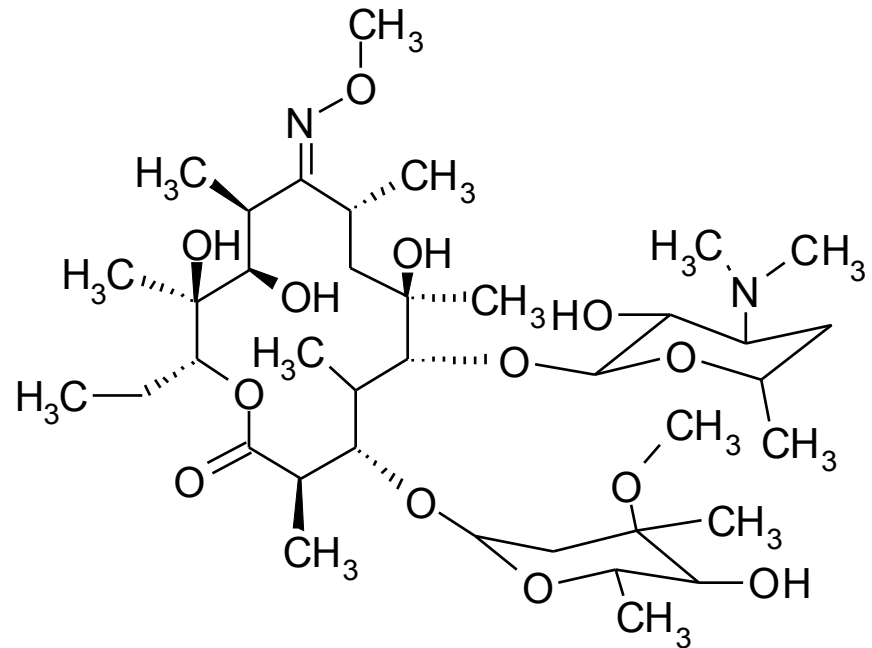
Macrolides

Erythromycine and its analogues

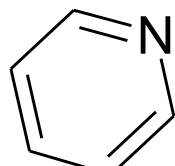


oleandomycin

•isolated 1954 from *Streptomyces antibioticus*



lexithromycin

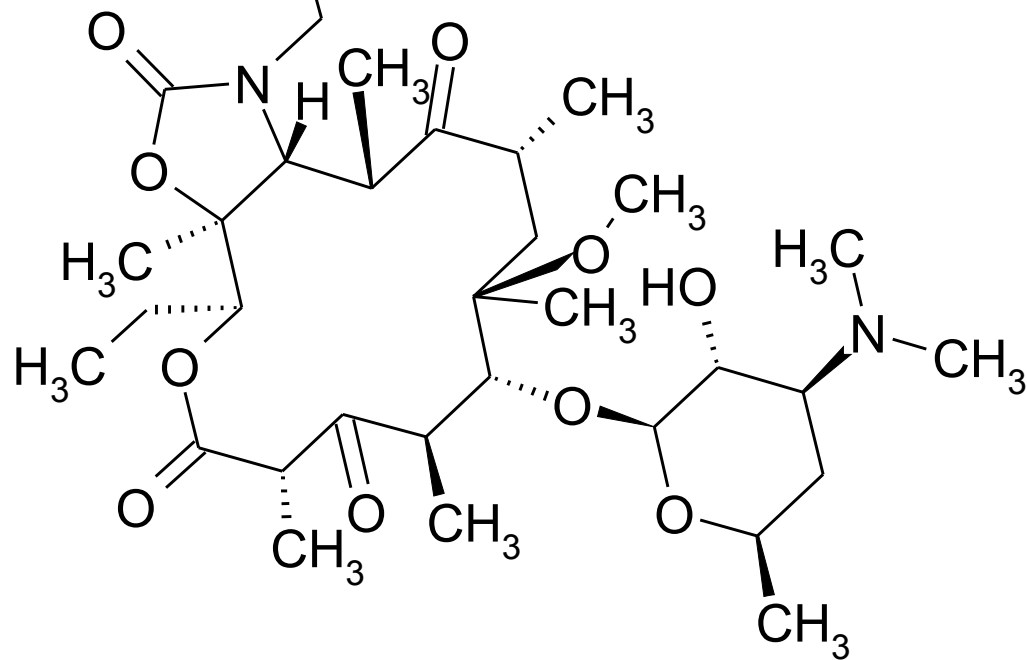


„More free“ erythromycine analogues: **Ketolides**

- keto-moieties in positions 2 and 9
 - good biological availability

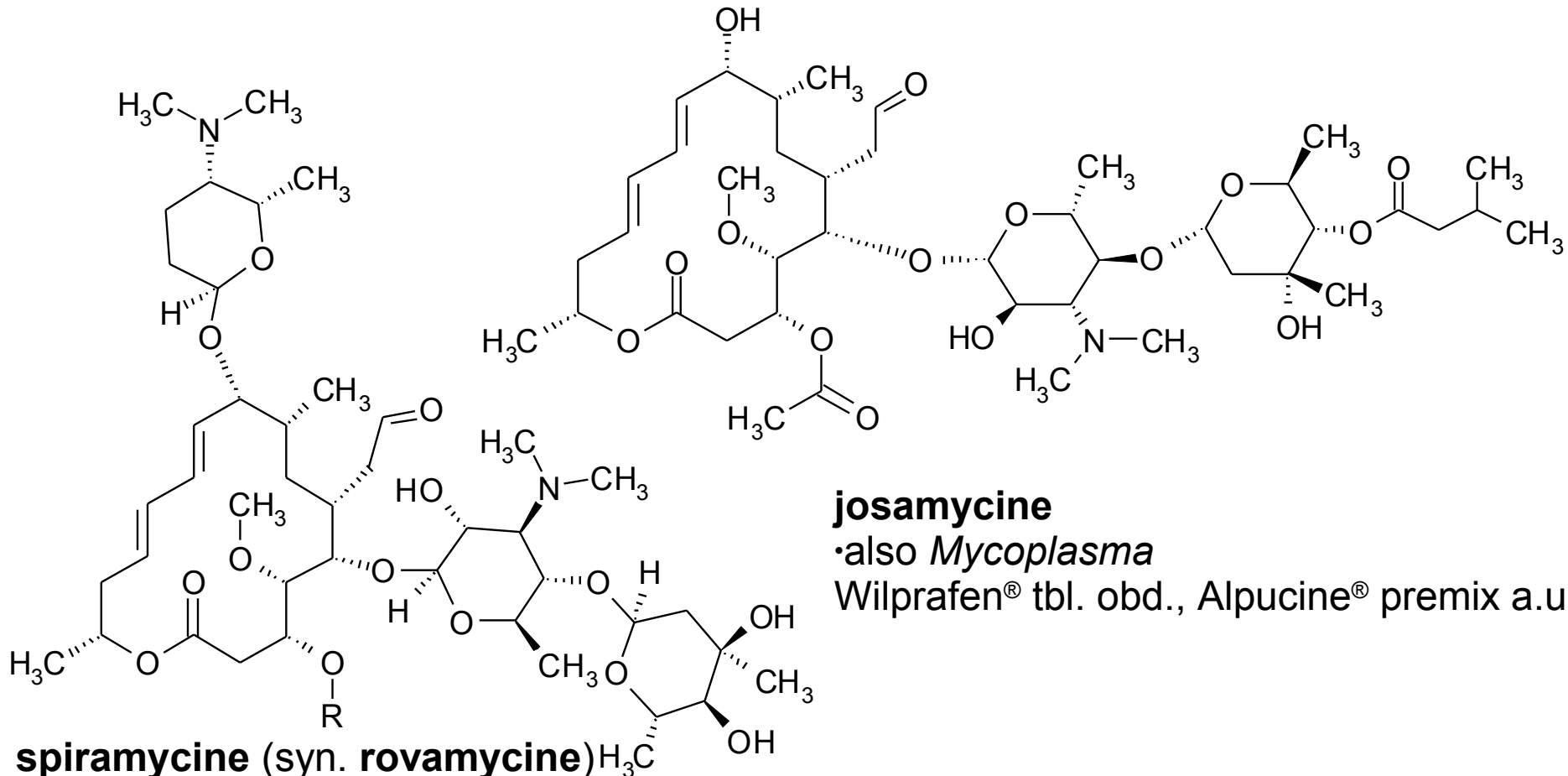
telithromycine

Ketek® tbl.



Macrolides

Compounds with 16membered lactone ring unsaturated in positions 10 and 12



spiramycin (syn. rovamycin)

•isolated 1954 from *Streptomyces ambofaciens*

R = HCO- spiramycin A = spiramycin I

CH₃CO- spiramycin B = spiramycin II

CH₃CH₂CO- spiramycin C = spiramycin III

Rovamycin® tbl. obd.

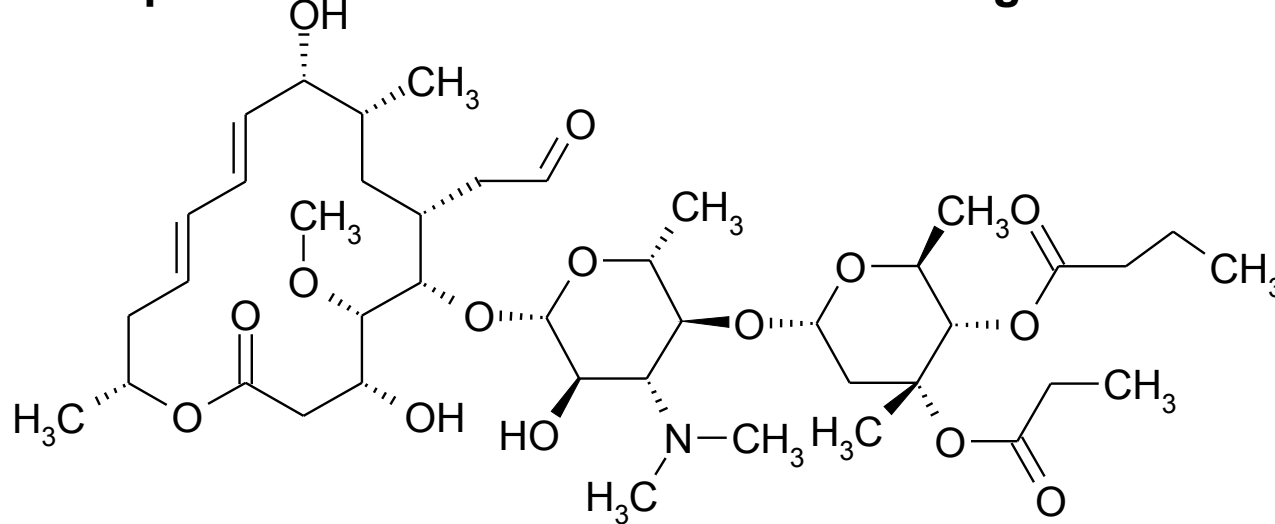
josamycin

•also *Mycoplasma*

Wilprafen® tbl. obd., Alpucine® premix a.u.

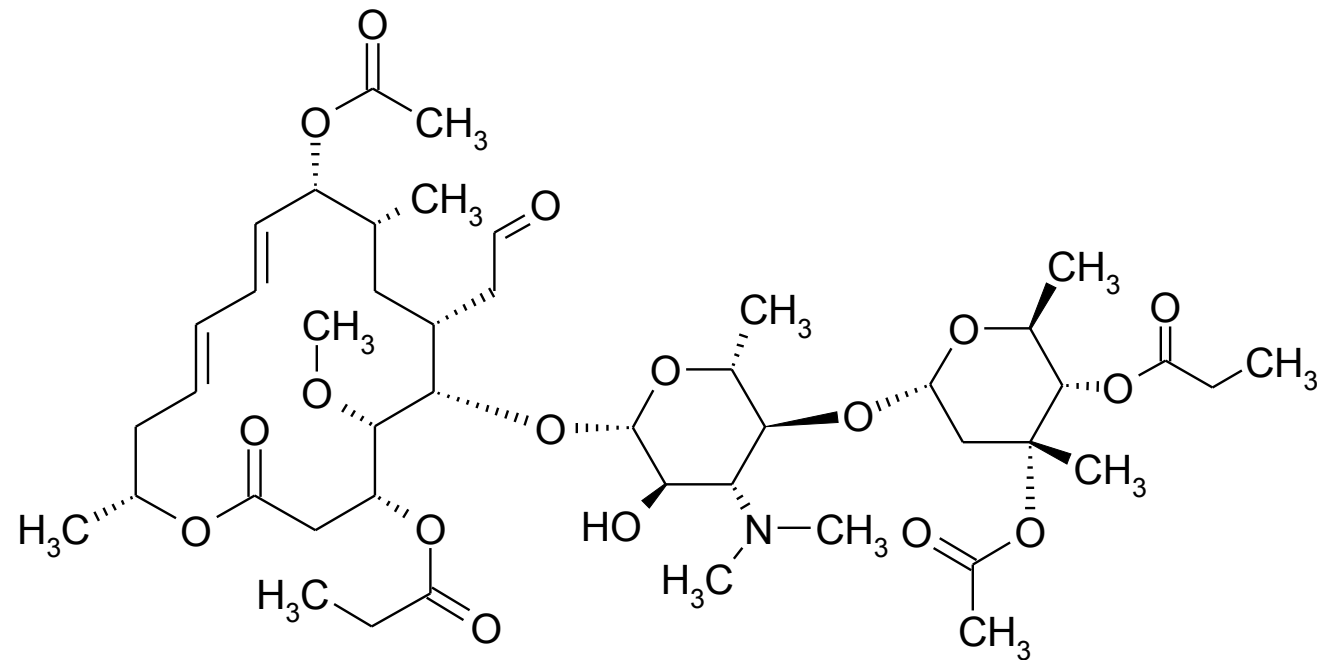
Macrolides

Compounds with 16membered lactone ring unsaturated in positions 10 and 12



rokitamycin

miokamycine



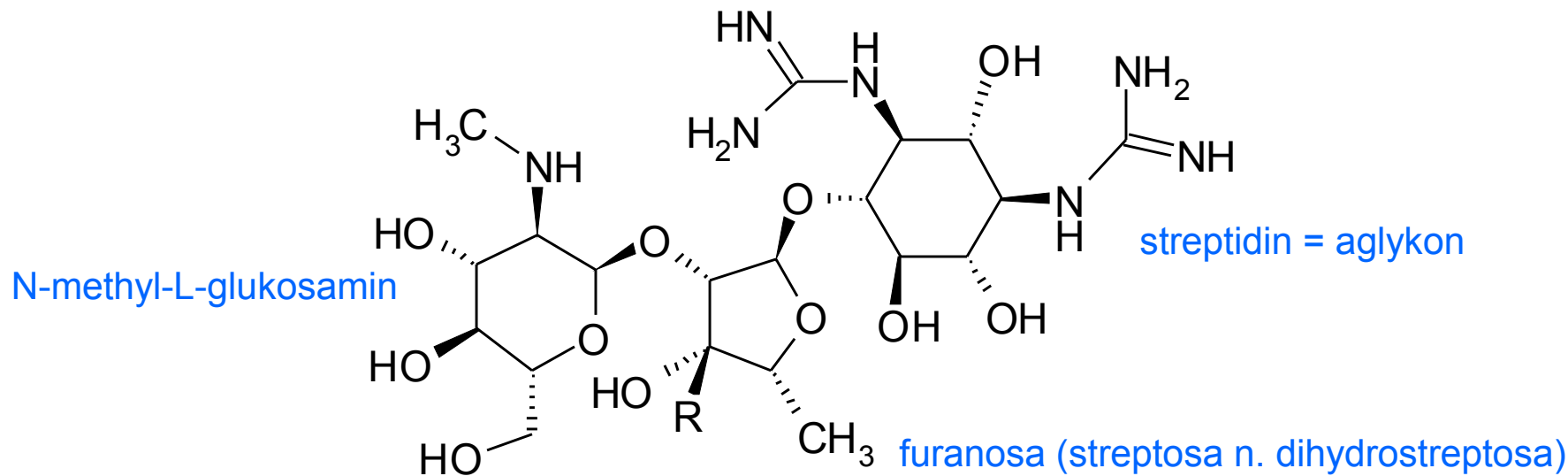
Aminoglycosids

- aminosaccharide glycosides produkované rodem *Streptomyces*
 - Skupina streptomycinu
 - Skupina neomycinu
 - Skupina kanamycinu a gentamycinu
- Mechanismus účinku
 - inhibice proteosynézy
- narušují přesné čtení genetického kódu a vazbu peptidyl-tRNA na peptidové vazebné místo
 - účinek bakteriostatický – baktericidní
- Spektrum
 - $G^+ < G^-$

Bacillus anthracis, Bordetella pertussis, Brucella, Corynebacterium diphtheriae, E. coli, Enterobacter, Haemophilus, Mycobacterium tuberculosis...

Aminoglykosidy

1. Skupina streptomycinu



R = -CHO

streptomycin

• izolován r. 1944 ze *Streptomyces fradiae*

• používán na *M. tuberculosis* v kombinaci s dalšími tuberkulostatiky

• baktericidní

Streptomycin „Grünenthal“® inj. sic., Streptowerfft® a.u.v

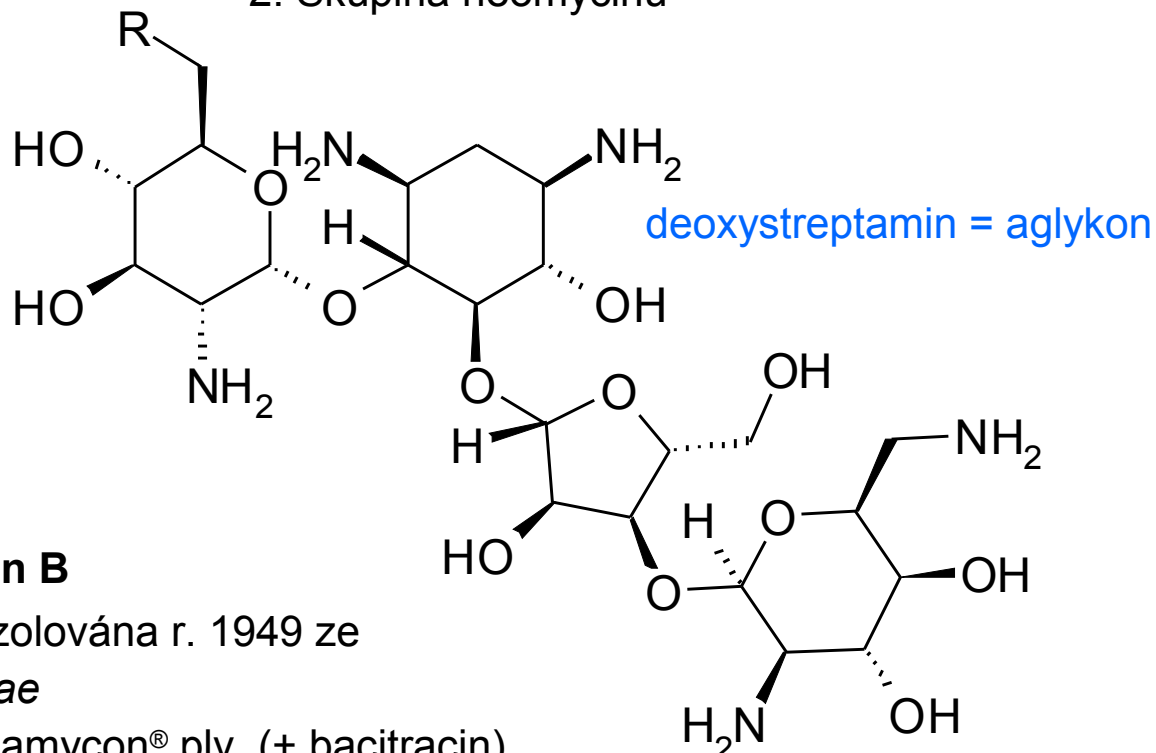
R = -CH₂OH

dihydrostreptomycin

Depomycine® a.u.v. inj. (+ benzylpenicilin)

Aminoglykosidy

2. Skupina neomycinu



R= -NH₂ **neomycin B**

směs neomycinů izolována r. 1949 ze

Streptomyces fradiae

Framykoin® ung., Pamycon® plv. (+ bacitracin)

R = -OH **paromomycin**

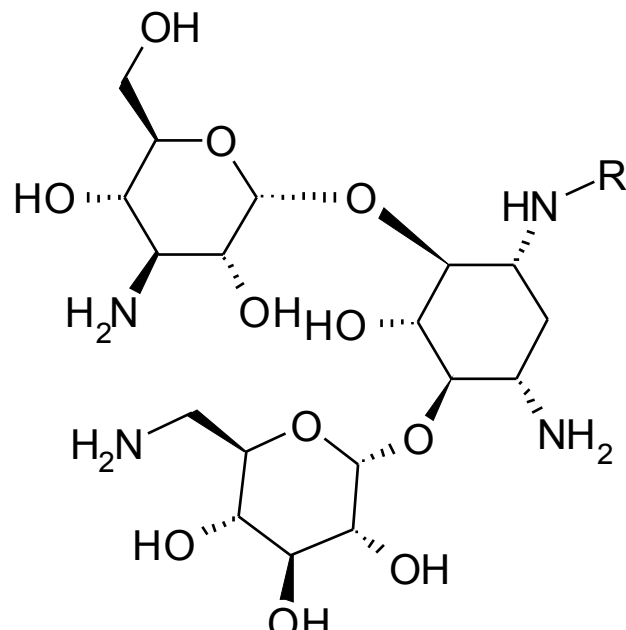
nevstřebává se z GIT

používán na *Entamoeba histolytica*

Humatin® cps.

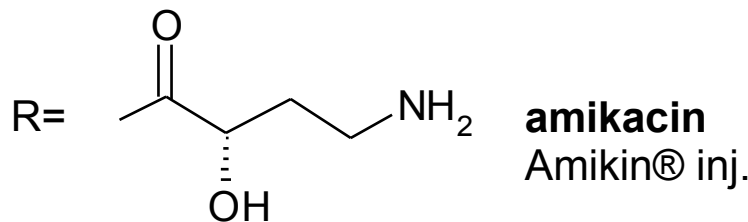
Aminoglykosidy

3. Skupina kanamycinu a gentamycinu

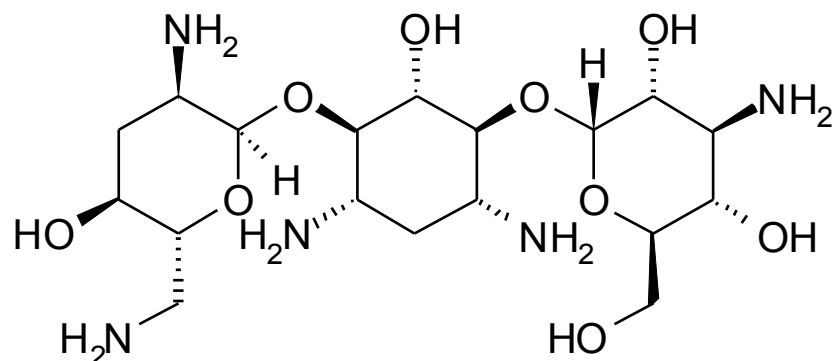


R = -H

Kanacol® a.u.v. inj.



Podskup.kanamycinu

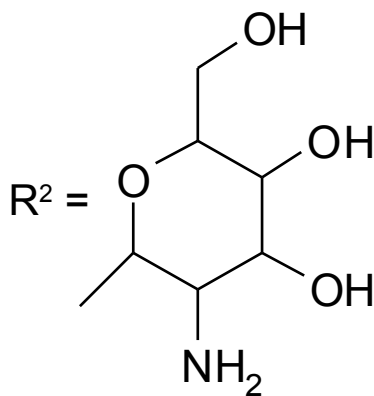
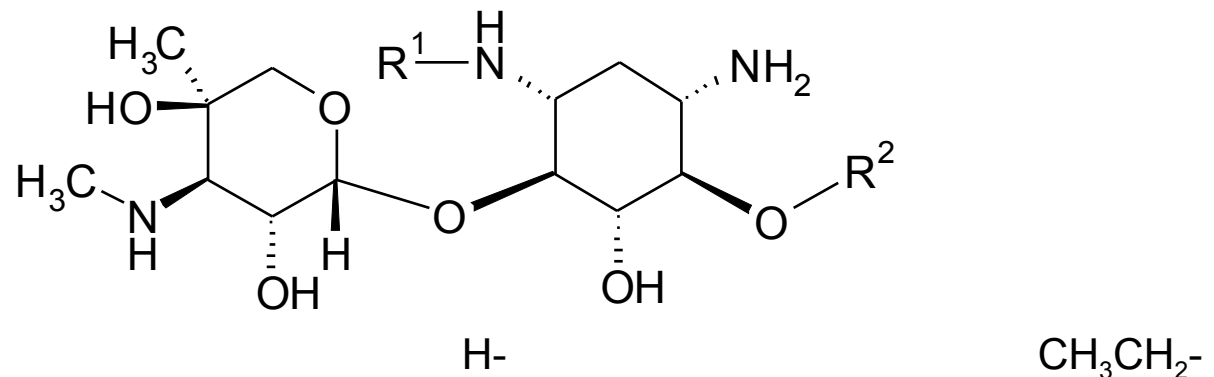


tobramycin

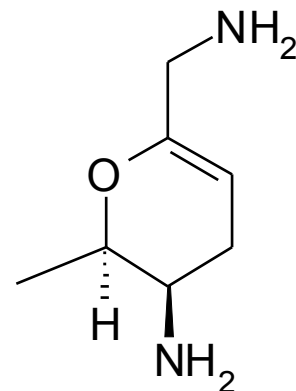
Tobi Nebuliser Solution® inh.
sol.

•léčba chronické
pseudomonádové pulmonální
infekce u pacientů s cystickou
fibrózou

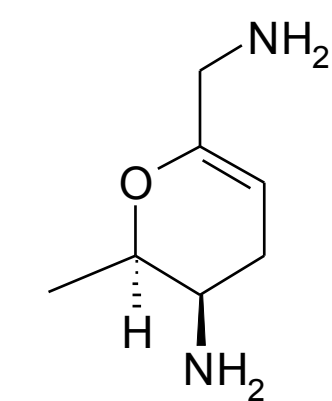
Aminoglykosidy
 3. Skupina kanamycinu a gentamycinu
 Podskup.gentamycinu



gentamycin
 Garasone® gtt. opht.
 (+betamethason)
 Diagen® a.u.v.



sisomycin



netilmycin
 Netromycine® inj.
 •těžké infekce, sepse ...