

Cell wall synthesis inhibitors

Part 2

Pharmacology 3
Dr. Rawan Abudalo

Department of Clinical Pharmacy and Pharmacy Practice
Faculty of Pharmaceutical Sciences
Hashemite University

By: Yana Hani
Dr: Heba

II. CEPHALOSPORINS

prefix : Cef or Ceph

- ① A wider spectrum than penicillins.
 - ② More resistance to B-lactmases enzyme.
 - ③ Eliminated by kidney.
 - ④ More expensive than penicillins.
- 

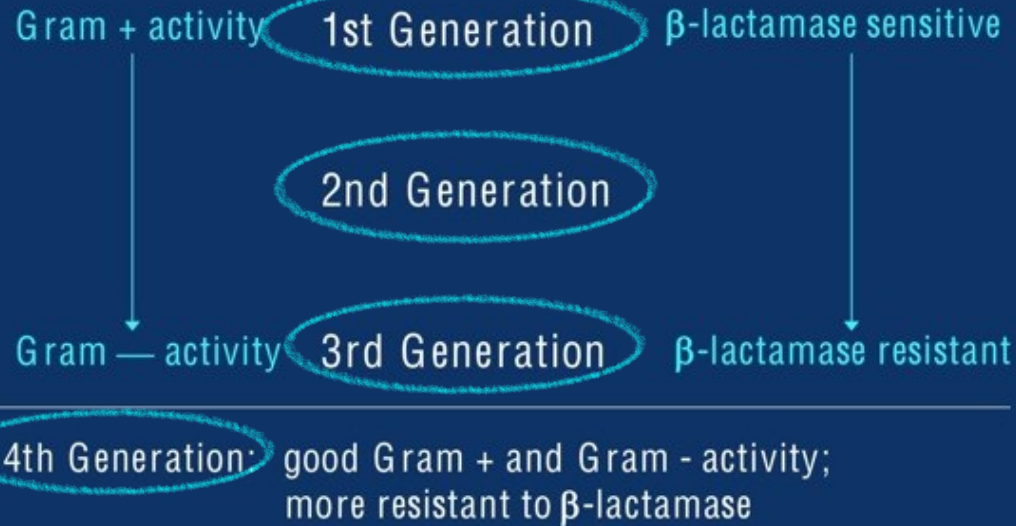
① يكون فعال ضد G^+ و G^- (وكل جيل نجون اقوى من الى قبل).

② منشآت هيكلية (Kinetic) لها بتشبه (penicillin)

③ يفرغهم افراج عن طريق الكلية.

④ هم أغلى سعراً من (penicillin).

Cephalosporins



بیشتر همد کل ما نیک جیل
جدید بکون (wide spectrum) اکتن

Summary of therapeutic applications of cephalosporins

First-generation cephalosporins	
Gram (+) cocci	<u>Staphylococcus aureus*</u> <u>Staphylococcus epidermidis</u> <u>Streptococcus pneumoniae</u> <u>Streptococcus pyogenes</u> <u>Anaerobic streptococci</u>
Gram (-) rods	<u>Escherichia coli</u> <u>Klebsiella pneumoniae</u> <u>Proteus mirabilis</u>
*Methicillin-resistant staphylococci are resistant	

Second-generation cephalosporins	
Gram (+) cocci	<u>Staphylococcus aureus</u> <u>Streptococcus pneumoniae</u> <u>Streptococcus pyogenes</u> <u>Anaerobic streptococci</u>
Gram (-) cocci	<u>Neisseria gonorrhoeae</u>
Gram (-) rods	<u>Enterobacter aerogenes</u> <u>Escherichia coli</u> <u>Haemophilus influenzae</u> <u>Klebsiella pneumoniae</u> <u>Proteus mirabilis</u>
Anaerobic organisms**	
***Cefoxitin and cefotetan have anaerobic coverage	

Third-generation cephalosporins	
Gram (+) cocci	<u>Streptococcus pneumoniae</u> <u>Streptococcus pyogenes</u> <u>Anaerobic streptococci</u>
Gram (-) cocci	<u>Neisseria gonorrhoeae</u>
Gram (-) rods	<u>Enterobacter aerogenes</u> <u>Escherichia coli</u> <u>Haemophilus influenzae</u> <u>Klebsiella pneumoniae</u> <u>Proteus mirabilis</u> <u>Pseudomonas aeruginosa</u> <u>Serratia marcescens</u>

هاد هو للحفاظ لكن حتى نقتنع المخت
أنه اول جيل كان فعال شوية منه
(G-ve) والجيل الي بعده المخت
حتى كان ضد (anaerobic) واي
بعده اكثر واكثر

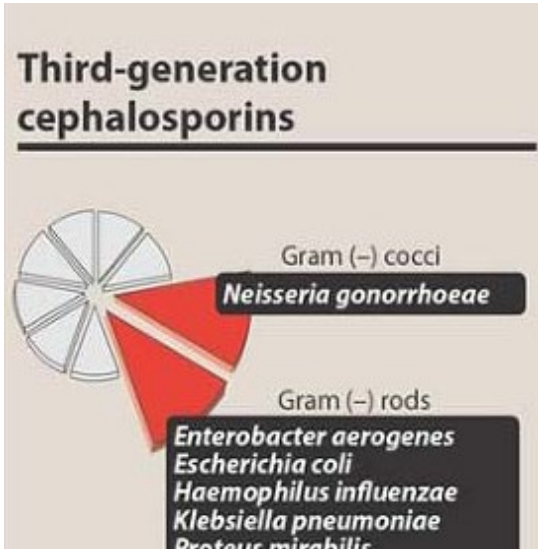
Cephalosporins

subgroups	indication
First generation (Cefazolin, cephalexin)	surgical prophylaxis
Second generation (Cefaclor, cefuroxime, and cefprozil)	active against <u>H</u> influenzae, <u>sin</u> usitis, <u>ot</u> itis, and lower <u>res</u> piratory tract infections
Third generation (ceftriaxone, cefotaxime, cefpodoxime, cefdinir, cefixime)	<u>M</u> eningitis, <u>end</u> ocarditis, <u>emp</u> irical therapy of sepsis in both the immunocompetent and the immunocompromised patient
Fourth generation (Cefepime)	<u>P</u> neumonia, <u>Emp</u> iric therapy in febrile <u>neutropenic</u> patients, <u>UTI</u>
Advanced generation-5 th (Ceftaroline)	complicated <u>skin</u> and <u>soft tissue</u> infections and community-acquired <u>pneumonia</u>

→ خصوصاً في عمليات جراحة العظام , غضاريف , رباط صليبي

→ في منهم شراب للأطفال وفي منهم صبوي

Third generation Cephalosporins



Adequate therapeutic levels in the CSF, regardless of inflammation, are achieved only with the **third-generation cephalosporins**.

Are effective in the treatment of neonatal and childhood **meningitis** caused by *H. influenzae*. meningococcal meningitis.

Third-generation cephalosporins must be used with caution, as they are associated with significant "**collateral damage**," including the induction of antimicrobial resistance and development of **Clostridium difficile infection**.

← بيوصل لـ (CNS) بشكل هينج وبستخدام لعلاج

(meningitis) و (spectrum) نوعاً ما واسع؛ فهو من
(antibiotic) الواردة إليها تعمل (superinfection) أو بمصطاع
تأني (Collateral Damage)

Cephalosporins Active against Methicillin-Resistant
Staphylococci (advanced generation; 5th generation)

- **Ceftaroline.**
- The unique structure allows ceftaroline to bind to PBPs found in MRSA and penicillin-resistant *Streptococcus pneumoniae*.
- Ceftaroline is currently approved for the treatment of complicated skin and soft tissue infections and community-acquired pneumonia.

Therapeutic advantages of some clinically useful cephalosporins

First Generation

Cefazolin

This first-generation parenteral cephalosporin has a longer duration of action and a similar spectrum of action, compared to other first-generation drugs. It penetrates well into bone.

Cefadroxil

Cephalexin

This is the prototype of first-generation, oral cephalosporins. Oral administration twice daily is effective against pharyngitis.

Second Generation

Cefuroxime sodium

This prototype second-generation, parenteral cephalosporin has a longer half-life than similar agents. It crosses the blood-brain barrier, and it can be used for community-acquired bronchitis or pneumonia in the elderly and for patients who are immunocompromised.

Cefuroxime axetil

Administered twice daily, this drug is well absorbed and is active against β -lactamase-producing organisms.

Third Generation

Cefdinir
Cefixime

These are administered orally once daily.

Cefotaxime

This penetrates well into the CSF.

Ceftazidime

This is active against *Pseudomonas aeruginosa*.

Ceftriaxone

This drug has the longest half-life of any cephalosporin (6 to 8 hours), which permits once-a-day dosing. High levels of the drug can be achieved in blood and CSF. It is effective against genital, anal, and pharyngeal penicillin-resistant *Neisseria gonorrhoeae*. The drug is excreted in bile and may be used in patients with renal insufficiency. It has good penetration into bone.

Fourth Generation

Cefepime

This is active against *Pseudomonas aeruginosa*.

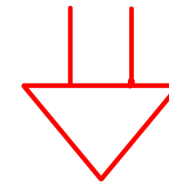
Advanced Generation

Ceftaroline

This is active against MRSA.

هاد تخيى لكل الي عيانه قبل ، الي بالاسف
رعت حكته الدكتورى بحال صرّة .

الي بالبوكس الاسود هون (orally)



الي تحت هاد تخيى ♥

First Generation

- Cefazolin :

هو (parenteral) ، الطول (longe duration) مقارنة
بالأنواع الثانية من الجيل الأول، لكن أنه نفس (spectrum of
action). ويوصل للعظم بشكل سريع.

- Cephalexin:

هو (prototype)، يستخدم مرتين باليوم (orally)
يستخدم لـ (pharyngitis).

- Cefadroxil: نفسه ↑

Second Generation

- Cefuroxime Sodium :

هو (prototype) يؤخذ (parenteral)، وعنده أطول (half-life)
في (2nd)، يعبر (BBB) وينتشر نسيجه في (community acquired
bronchitis) أو (Pneumonia) في كبار السن أو (immunocompromised).

- Cefuroxime axetil:

يؤخذ مرتين في اليوم (orally)، امتصاصه سريع وفعال ضد
العاثيات التي تنتج (β-lactamase).

Third Generation

- Cefdinir , Cefixime:

يؤخذ مرتين في اليوم (orally).

- Cefotaxime:

علاج بويصل (CSF).

- Ceftriaxime:

يستخدم لعلاج (Pseudomonas
aeruginosa).

- Ceftriaxone:

هو أطول (half-life) في كل (Cephalosporin)، يؤخذ مرة واحدة يومياً، يعبر امتصاص
امتصاص كبير في الدم وفي (CSF) وفعال ضد (penicillin resistant).

Neisseria gonorrhoeae
genital
anal
pharyngeal

يعبره إخراج عن طريق (bile) لذلك يمكن استخدامه في مرضى
(renal insufficiency) ويمنح بعبء امتصاص في العظام.

Fourth Generation

- Cefepime:

فعال ضد (Pseudomonas aeruginosa).

Advanced Generation

- Ceftaroline:

فعال ضد (MRSA).



Pharmacokinetics

1. Administration:

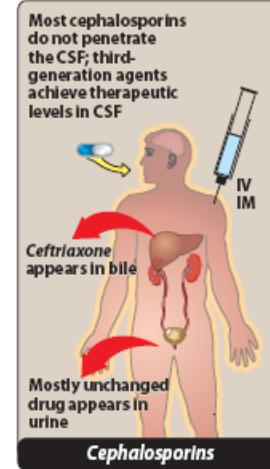
- Many of them must be administered **IV** or **IM** because of their poor oral absorption (however some can be given orally)

2. Distribution:

- CPNs distribute very well into body fluids but **not** to CSF.
- **Cefazolin** penetrates well into most tissues. It is a drug of choice for **surgical prophylaxis** including **orthopedic surgery** because of its ability to penetrate **bone**.
- Only **ceftriaxone** or **cefotaxime** achieve therapeutic levels in the CSF and have become agents of **choice for meningitis**.
- All CPNs cross the placenta.

3. Elimination:

- Tubular secretion and/or glomerular filtration
- Doses must be adjusted in cases of renal failure
- Exception: **Ceftriaxone**, excreted through the bile.....Employed in patients with renal insufficiency



← أغلب الادوية تكون **IM & IV** وشوية **orally**.

← بغيرهم امتصاص منه في الجسم لكن هو تكلم بيوصلها لـ (CSF).

(Cefazolin) ← **(surgical prophylaxis)** بها فيها **(orthopedic surgery)**.

(Ceftriaxone) / (Cefotaxime) ← يستخدم في **(meningitis)**

← تكلم بغيرهم **(excretion)** بواسطة الكلى إلا

Ceftriaxone

Adverse effects

✓ Hypersensitivity reaction

Current data suggest that the cross-reactivity between penicillin and cephalosporins is around 3% to 5% and is determined by the similarity in the side chain, not the β -lactam structure.

The highest rate of allergic cross-sensitivity is between penicillin and **first-generation cephalosporins**.

Cephalosporins should be avoided or used with caution in individuals with penicillin allergy.

Patients who have had an **anaphylactic response**, **Stevens-Johnson syndrome** or toxic epidermal necrolysis to penicillins **should not receive cephalosporins**.

Adverse effects

- ✓ pain after injection.
- ✓ Diarrhea.

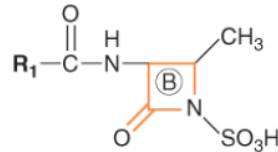
Some have anti-Vitamin K effect (bleeding).

Other B-Lactam Antibiotics- **Monobactams** | *Aztreonam*

Other B-Lactam Antibiotics- **Monobactams** | *Aztreonam*

Other B-Lactam Antibiotics- **Monobactams** | *Aztreonam*

- They are drugs with a monocyclic β -lactam ring.
- Their spectrum of activity is limited to aerobic Gram-negative organisms (including *P. aeruginosa*).
- **Aztreonam** is resistant to the action of β -lactamases.
- It is administered either IV or IM. Every 8 hrs
- this drug may offer a safe alternative for treating patients who are allergic to penic &/or cephalosporins.



Monobactams
↓



Aztreonam

✓ G is aerobic.

✓ *Pseudomonas aeruginosa*.

X anaerobic, G⁺.

IM or IV

كل مرضي كتي منيج لك الشخص الي عنهم حساسية من البنسلين و
السيفالوسبورين.

Other B-Lactam Antibiotics- Carbapenems

- broad-spectrum B-lactam antibiotics.
- Examples: **Doripenem, Imipenem, Meropenem, Etrapanem.**
- They resist hydrolysis by most B-lactamases.
- Carbapenems are active against *P aeruginosa* and *Acinetobacter* species (except ertapenem).
- These agents have a very broad spectrum of action and are usually restricted to use in hospitals for treatment of serious infections.

• broad عادةً يستخدمون فيهم (empirical Therapy) في

Other B-Lactam Antibiotics- Carbapenems

- All are cleared renally and the dose must be reduced in patients with renal insufficiency.
- Excessive levels of imipenem in patients with renal failure may lead to seizures.
- Imipenem undergoes cleavage by a dehydropeptidase found in the brush border of the proximal renal tubule. This enzyme **forms an inactive metabolite that is potentially nephrotoxic**.
 - Imipenem is formulated with cilastatin, which prevents hydrolysis of imipenem by renal dehydropeptidase.

(Imipenem): هاد الدواء بيحلل (hydrolysis) بواسطة إنزيم
(Dehydropeptidase) الموجودة في (Proximal tubules) ونتيجة لذلك
بيكون (inactive metabolite) وهاد ممكن يكون Nephrotoxic

← متشابه هيك بينه وبين (Cilastatin) هاد الي بيحلل على
نسيجه (Dehydropeptidase) وبيمنع تكون (metabolite)
الي بيحلل (nephrotoxicity).

Glycopeptide Antibiotics

- **Vancomycin**

Vancomycin inhibits synthesis of bacterial cell wall by binding to the D-Ala-D-Ala terminus preventing further crosslinking.

- IV for systemic infections

- Not absorbed after oral administration (the use of the oral formulation is limited to the treatment of severe antibiotic-associated C. difficile colitis.)

Narrow spectrum(G+ ve)

Bactericidal /not B-lactam.

Gram (+) cocci

*Staphylococcus aureus**
Staphylococcus epidermidis
Streptococcus groups A, B, C
Streptococcus pneumoniae
Enterococcus faecalis

*(Including methicillin-resistant strains)

Gram (+) bacilli

Listeria monocytogenes
Corynebacterium jeikeium

Gram (-) cocci

Gram (-) rods

Anaerobic organisms

*Clostridium species***

Spirochetes

Mycoplasma

Chlamydia

**Oral vancomycin only
for *C. difficile*

Other

Actinomyces

Glycopeptide Antibiotics

- **Vancomycin**
- **Orally:- every 6 hrs** for refractory pseudomembranous colitis due to C. difficile.
- **Slow IV infusion (1-2 hrs)** for treatment of systemic infections or prophylaxis.
 - is effective against MRSA. (DOC) *aminoglycosid*
 - Vancomycin in combination with A.G alternative regimen to treatment of enterococcal endocarditis.
- **Teicoplanin** is a glycopeptide antibiotic that is very similar to vancomycin in mechanism of action and antibacterial spectrum.

Gram (+) cocci
Staphylococcus aureus*
Staphylococcus epidermidis
Streptococcus groups A, B, C
Streptococcus pneumoniae
Enterococcus faecalis
* (including methicillin-resistant strains)
Gram (+) bacilli
Listeria monocytogenes
Corynebacterium jeikeium
Gram (-) cocci
Gram (-) rods
Anaerobic organisms
Clostridium species**
Spirochetes
Mycoplasma
Chlamydia
**Oral vancomycin only for C. difficile
Other
Actinomyces

Vancomycin

- S.E:-
- 1-Flushing (**red man syndrome**) with a **rapid** infusion.(**More common**)
- Prevented by prolonging the infusion period OR pretreatment with an antihistamine such as diphenhydramine.
- 2- **phlebitis**(inflammation of vein) at site of injection.
- 3- **ototoxicity & nephrotoxicity** (rare) but increased risk when administered with A.G.

↙
aminoglycosid



Other Cell wall synthesis inhibitors

Fosfomycin:

Inhibits the formation N -acetylmuramic acid precursor.

- **Therapeutic use** : It is indicated for urinary tract infections caused by E. coli or E. faecalis.
- Rapidly absorbed after oral administration & distributes well to the kidneys, bladder, and prostate

Bacitracin:

Inhibits the carrier that transfers peptidoglycan subunits to the growing cell wall.

- It is highly nephrotoxic when administered systemically and is only used topically

Cell membrane active agents

- **Daptomycin** – is a new lipopeptide antibacterial drug.
- Binds to cell membrane causing depolarization and rapid cell death (doesn't work on cell wall).
- **Therapeutic use:**
 - for treating infections caused by resistant gram-positive organisms, including MRSA and *vancomycin* resistant enterococci (VRE)
 - is indicated for the treatment of complicated skin and skin structure infections and bacteremia caused by *S. aureus*,
- **Adverse effect :**
Myopathy and creatine phosphokinase levels elevation.