

β -lactamase inhibitors

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اصحوا لأبوي

β -lactamase inhibitors

- Early attempts to combine β -lactamase inhibitors with penicillins failed

لا يقدرون ان يخلطوا مع بعض

- Also early attempts to combine penicillin β -lactamase resistant penicillins with wide-spectrum penicillinase sensitive penicillins failed to give synergistic activity

- Example Methicillin or Oxacillin with Ampicillin or Carbencillin.

- Reasons are:

not mixed together due to 8-

1. Failure of lipophilic penicillinase resistant agents to penetrate cell envelopes in Gram negative bacteria
2. Induction of β -lactamases by some penicillinase resistant penicillins.
3. The reversible binding of penicillinase-resistant penicillins to β -lactamase.. Higher concentration of this substance is needed to inhibit this enzyme.

لا تشكلوا covalent bond

كيف أقدر الامتحان ← Mechanism-Based β -lactamase inhibitors

Mechanism – Based β -lactamase inhibitors

- Examples:
- Clavulanic Acid (Natural): causes potent and progressive inactivation of β -lactamase
- Sulbactam (Synthetic)
- Tazobactam (Synthetic)
- Thienamycins: Natural, inhibit β -lactamases and bind to PBPs

(*) Clavulanic acid + Sulbactam + Tazobactam $\rightarrow$ class (I) β -lactamase inhibitors
(*) Thienamycins + Carbapenem $\rightarrow$ class (II) Beta lactamase inhibitors
(MOA + Structures) مختلفين.

Mechanism – Based β -lactamase inhibitors

- **Class I** inhibitors that have a heteroatom leaving group at position 1 (e.g., clavulanic acid and sulbactam) and **Class II** inhibitors that do not (e.g., the carbapenems).
*فلا تفسد الـ Penicillin
ممكن بدالـ Sulphur group
نفس (C) ←*
- Unlike competitive inhibitors, which bind reversibly to the enzyme they inhibit, mechanism-based inhibitors react with the enzyme in much the same way that the substrate does. With the β -lactamases, an acyl-enzyme intermediate is formed by reaction of the β -lactam with an active-site serine hydroxyl group of the enzyme.
*الـ Class II + (2)
cross linkage and enzyme becomes inactive*

MoA of class (1) + class (2)

in class (1)

مرحبة ال clavulanic acid يحتوي على good leaving group at C1 (S60)

enzyme β -lactamase يحتوي على Serine amine على السطح مع nucleophilic attack على C7 of clavulanic acid

على C7 يوجد acyl group وهذا الخطأ = تعتبر reversible transient inhibitors

يكون ممكن التراجع خلفه يرجع Free ولكن بسبب وجود good leaving group على C4

في C4 يوجد Not stable وايضا هناك على الحلقة (Ring strain)

مع تنكس الرابطة القوية فيها وهي كما ان تعتبر transient inhibitors وف تكون

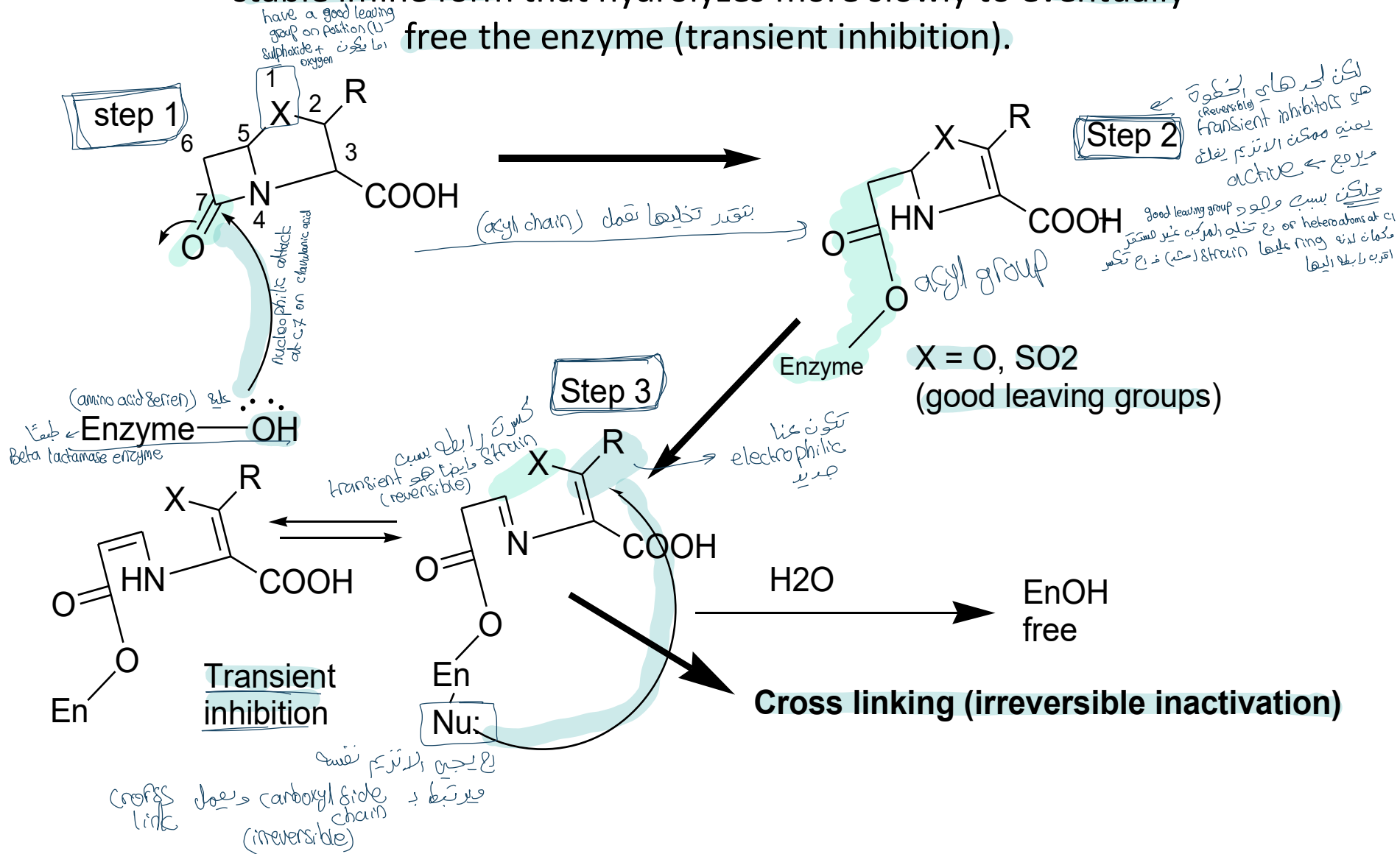
مع electrophilic على الحلقة التي انكسرت والآن نرى مع Nucleophilic attack cross linkage وهي (irreversible)

اما ال class (2) ← على C7 يوجد نفس الخطأ = ولكن بسبب عدم وجود good leaving group at C1

لذا هي (Carbon group) فارد يصير هنا cross link ويقتل بمرور الوقت transient inhibitory

وبالتالي يرجع β -lactamase يتكرر ← وهذا طرح السلايد (6+7)

Mechanism of inactivation **Class I** inhibitors the acyl-enzyme intermediate formed when a mechanism-based inhibitor is attacked by the enzyme is diverted by tautomerism to a more stable imine form that hydrolyzes more slowly to eventually free the enzyme (transient inhibition).



الافتلاف بين class (1) و class (2)

Differences in

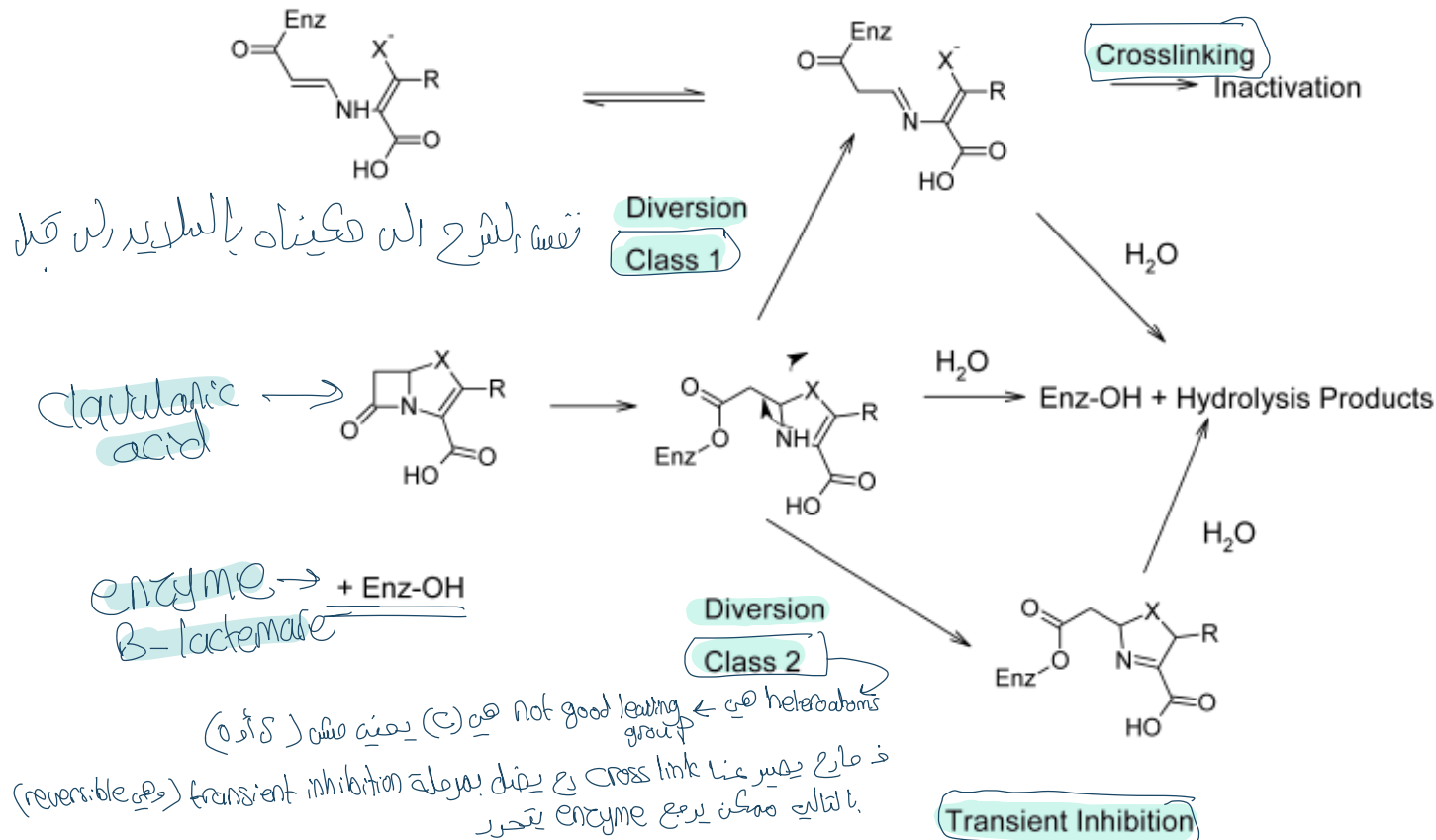


Figure 8.4 • Mechanism-based inhibition of β -lactamases.

Class I β -lactamase inhibitors

- Inhibitors include: Clavulanic acid and sulbactam, it contains a good leaving  heteroatom at the 5-membered rings.
- This type can lead to irreversible inhibition. *and cross linked*
- Used with Ampicillin and Amoxicillin.

Products

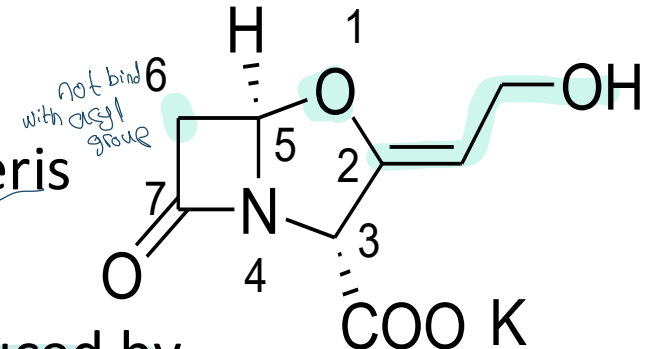
Clavulanate potassium

is class C1 ← ~~⊗~~ clavulanic acid structures 8

- not have acyl group in C6
- on C1 have an oxygen
- side chain is ethyl alcohol
- not have pharmacophore for activity
- not active for antibiotics

- Clavulanate potassium
- Antibiotic from Streptomyces clavuligeris
- Very weak antibacterial activity
- Potent inhibitor of β -lactamases produced by
- Staph. Aureus ⁽¹⁾ and Gram negative bacteria ⁽²⁾
- Combined with Amoxicillin for oral administration to treat
skin, respiratory, ear, UTI infections Augmentin®
- Oral bioavailability is similar to Amoxicillin
- Clavulanic acid is acid stable

only 10%
pharmacophore
penicillins



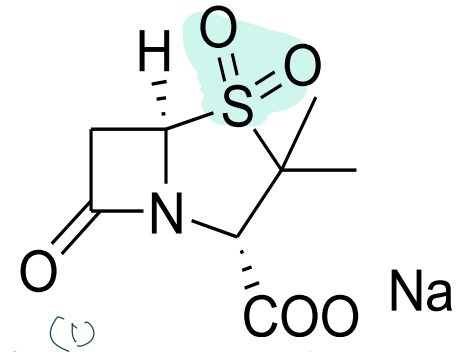
MoA → Form a covalent bond with Beta lactamase, when give with amoxicillin (Binding protein needed) ^{البروتين}

Products

cool
Sulfoxide

Sulbactam

class (1)
not give orally + synthetics



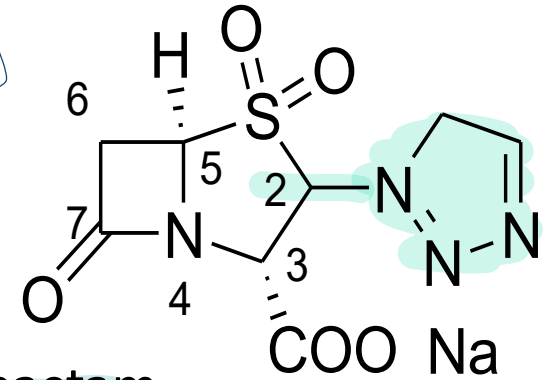
- Synthetic sulfone
- Potent inhibitor of β -lactamases produced by Staph aureus and many Gram negative bacteria
- Weak antibacterial activity → pharmacophore is similar to Penicillins
- Improves and potentiates the activity of Ampicillin and Carbencillins against Staph aureus (β -lactamases producing) and members of Enterobacteracia family, Gram (-) *P. aeruginosa* etc. *S. P. Carbapenem* is also *P. aeruginosa* is not Beta lactamase class (1) *clavulanic acid* *Amoxicillin*
- It does not, however, synergize with either carbenicillin or ticarcillin against *P. aeruginosa* strains resistant to these agents. Failure of sulbactam to penetrate the cell envelope is a possible explanation for the lack of synergy.
- Failure to penetrate cell envelope may be the reason
- Used parentally

Products

سولبكتامس نفس structure تبع
tri-thiazole ولكن له 2 عن tri-thiazole

Tazobactam

not give orally



- Synthetic sulfone
- More Potent inhibitor of β -lactamases than Sulbactam
- Weak antibacterial activity

Tazobactam is available in fixed-dose, injectable combinations with piperacillin, a broad-spectrum penicillin consisting of an 8:1 ratio of piperacillin sodium to tazobactam sodium by weight and marketed under the trade name Zosyn®.

- Used in combination with piperacillin for the treatment of appendicitis, postpartum endometritis, pelvic inflammatory disease, skin infections, and pneumonia.

- *Note: don't forget that if we have MSRA, we can't use these inhibitors, why? Because MRSA is resistant to penicillins because it modifies transpeptidase itself.*

MRSA مقاومتها ليست نتيجة ال Beta lactamase وانما نتيجة ال transpeptidase ال enzyme ال inhibition فهو يحور Binding pocket بفعلة Penicillin يعلق يرتبط فيها بسهولة بتغير تركيبها

ممكن ان يكون
الكثير من ال
قبل بآخر علاج

الاسم
التجاري
موجود في المستشفيات

3

2

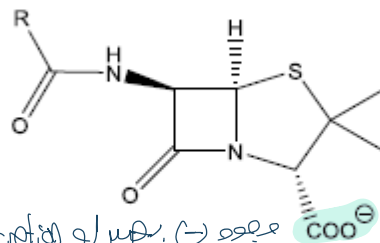
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⊗ زمان بالحرق الحامضية للمعدة حركتها صار في قلة في الامعاء
 فكل ما في الدم يدخل الامعاء في جسم المريض لفترة طويلة ليتجنبها
 يعطوه مريض متكررة كان يعطوا ال Probenecid هو منافس ال
 antibiotics (penicillin) على ال excretion

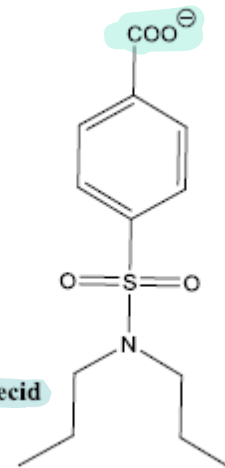
not Beta lactamase inhibitors probenecid

- Note: penicillins has carboxylic acid, so it becomes carboxylate in plasma, so tightly bound to plasma proteins (because albumin's charge is positive)
- also penicillins are candidate for renal secretion by anionic pump (which pumps carboxylic acids in the
- urine), that's why sometimes they add **probenecid** (adjuvant contains carboxylic acid) to compete with penicillin on the pump, so probenecid reduces elimination rate of penicillin and penicillin remains longer in the body.

they compete to plasma proteins and have short half life.



Penicillin



Probenecid

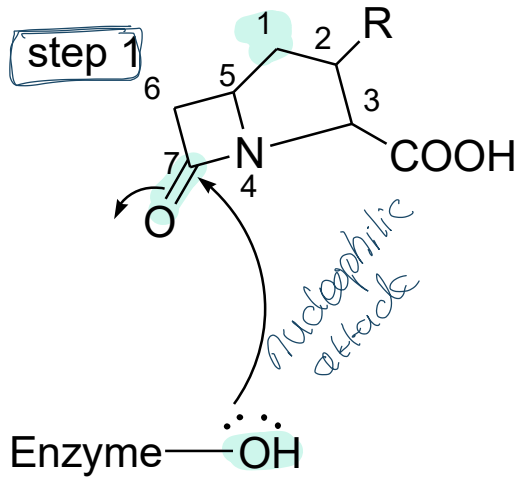
معدة (-) مفرط excretion في nephrons جبر ال
 anionic pump تخرج مع urine

لـ اذا سالت لهاي Pump تنبيه (inhibition)
 او يعطي مركب منافس ال Penicillin عليها بالتالي رجع
 يضل في جسم لفترة طويلة فلا ال Probenecid

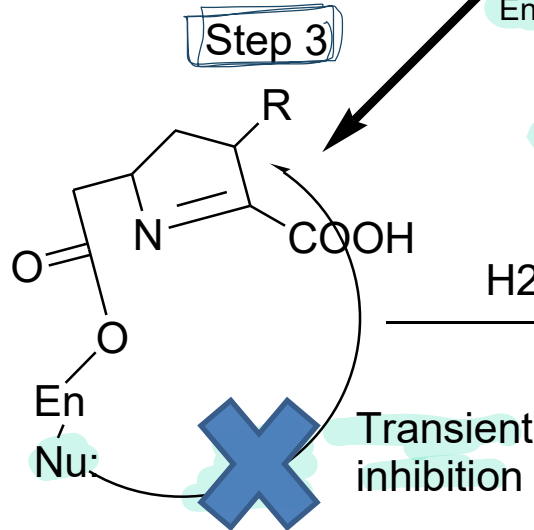
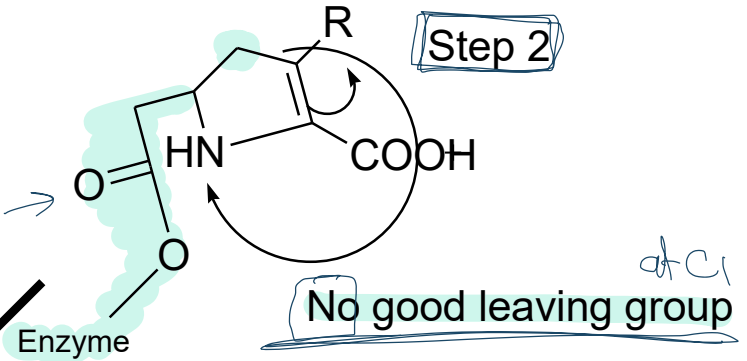
not contain leaving group → No cross link occurs → enzyme will still free after will

acetyl carbapenem

Class II inhibitors



acyl est group



H₂O

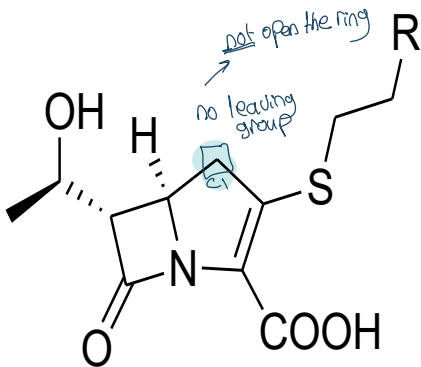
EnOH free + Hydrolysis products

Transient inhibition

no cross linking occurs

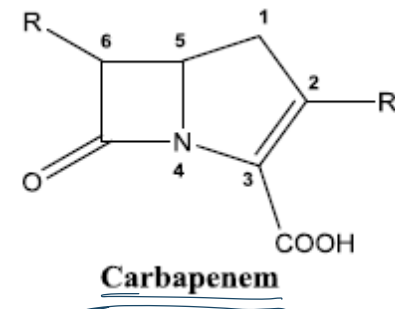
Class II inhibitors

- This class include Carbapenams, Imipenem
- Both have β -lactamase inhibition activity and antibacterial activity.



R = NH₂ Thienamycin

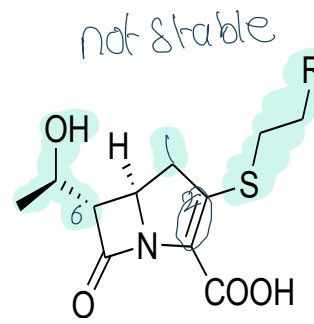
R = NHCH=NH imipenem



Carbapenems

Thienamycin

natural



R = NH₂ Thienamycin

R = NHCH=NH imipenem

- Thienamycin isolated from “Streptomyces cattleya”
- No (S atom) at position 1
- Double bond at C2-C3 (The double bond in the bicyclic structure creates considerable ring strain and increases the reactivity of the - lactam to ring opening reactions.)
- ^{8 at C2} Cystamine side chain at C2
- Simple alcohol side chain at position 6
- Stereochemistry is 5R:6S:8S
- They have two strained rings which decrease the chemical stability as well as acid stability. An unfortunate property of thienamycin is its chemical instability in solution. It is more susceptible to hydrolysis in both acidic and alkaline solutions
- Optimum stability pH 6-7 (7)
- Have broad spectrum activity.

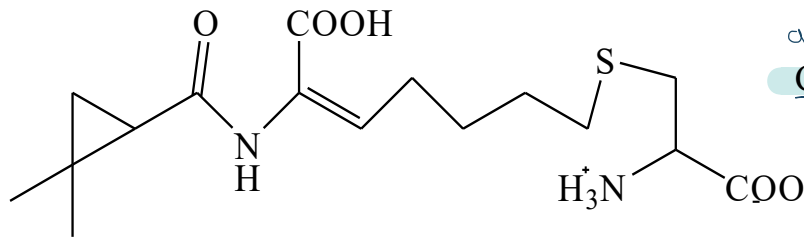
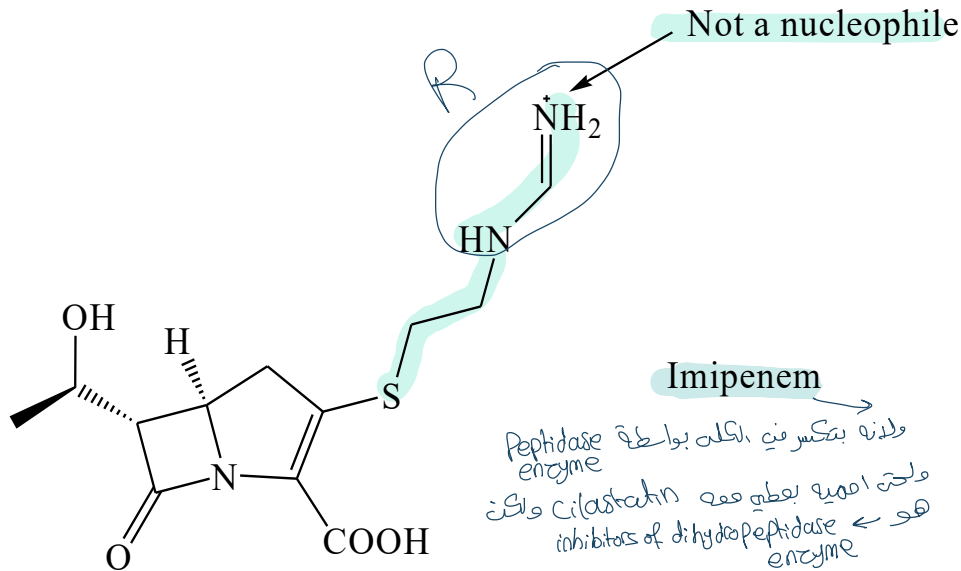
لديه بيفيدوس بوس ٩٥

Thienamycin properties:

- 1 • - Very potent β -lactamase inhibitor
- 2 • - Outstanding spectrum of activity that covers
gram +ve, gram -ve bacteria (including
pseudomonas aeruginosa), aerobic and
anaerobic bacteria.
- 3 • *Note: antibiotics that work on anaerobic*
bacteria is limited due to its high resistance.
- 4 • - Orally inactive.

Carbapenems

Imipenem – Cilastatin



Imipenem is indicated for the treatment of a wide variety of bacterial infections of the skin and tissues, lower respiratory tract, bones and joints, and genitourinary tract, as well as of septicemia and endocarditis caused by lactamase-producing strains of susceptible bacteria

Carbapenems

Imipenem – Cilastatin

- Chemically more stable than Thienamycin
- Cilastatin is inhibitor of (dehydropeptidase-1) ^{التي يحول breakdown of imipenem}
- Cilastatin give enzymatic stability for Imipenem ^{تنتج not chemical stability}
- $T_{1/2} = 1$ hour (short) due to renal secretion of penicillin
- Imipenem is extraordinary wide-spectrum antibacterial agent. It is an inhibitor of β -lactamases from certain Gram-negative bacteria resistant to other β -lactam antibiotics (e.g., *P. aeruginosa*, *S. marcescens*, and *Enterobacter* spp.).
- Some *Pseudomonas* spp. are resistant, such as *P. maltophilia* and *P. cepacia*, as are some methicillin-resistant staphylococci. Imipenem is effective against non-lactamase-producing strains of these and additional bacterial species, but other less expensive and equally effective antibiotics are preferred for the treatment of infections caused by these organisms.

ليس
imipenem is
expensive

Carbapenems

Imipenem – Cilastatin

- β -lactamases resistant and class II inhibitor
- Imipenem – Cilastatin are available as sterile powder for injection, in solution its stable for 4 hours at 25 C
- Synergistic action with Aminoglycosides, but chemically incompatible

Positive و negative charged لا يتفاعلان

Penicillins (negative charged)

Aminoglycoside (positive charged)

chemical incompatible لا يتفاعلان

Imipenem is:

قوة فاعلة منخفضة لاس = ضامة

- A **reserved** antibiotic, outstanding in spectrum, parenterally administered and stable under neutral conditions.
- Imipenem has very short half-life (1 hour) , because it has carboxylic acid which makes it candidate for active secretion and because it's unstable.
- So imipenem is coadministered with **cilastatin** (that contains COOH in its structure) , so increases the duration of imipenem remaining in the body and protects the kidneys from toxic metabolites of imipenem.
- Note: **reserved antibiotic** means that these antibiotics which are broad spectrum can
- be only available at hospitals due to resistance issues.

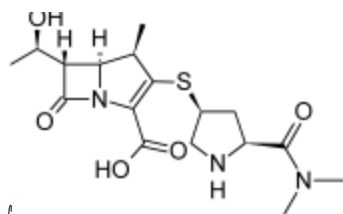
منحامل بفعال مركبات أكثر فعالية وأكثر *stability* فكانت مشكلة *thienomycin* في *stability*

NEWER CARBAPENEMS

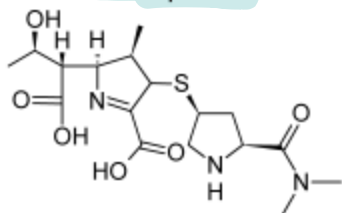
- The extended spectrum of antibacterial activity associated with the carbapenems together with their resistance to inactivation by most β -lactamases make this class of β -lactams an attractive target for drug development.
 - In the design of new carbapenems, structural variations are being investigated
 - with the objective of developing analogs with advantages
 - over imipenem.
1. Improvements that are particularly desired include stability to hydrolysis catalyzed by DHP-I,
نظير الـ imipenem يكون نفسه active against hydrolysis in DHP-I بعون ما أعطيه مع *clastin*
 2. stability to bacterial metallo--lactamases ("carbapenemases") that hydrolyze imipenem, activity against MRSA.
انزع بـ *carbapenem*
 3. increased potency against *P. aeruginosa*, especially imipenem-resistant strains.
ردنه في أنواع منها يكون Resistance
 4. Enhanced pharmacokinetic properties, such as oral bioavailability and a longer duration of action, have heretofore received little emphasis in carbapenem analog design.
ردنه هو ما يقطعه *orally*

NEWER CARBAPENEMS

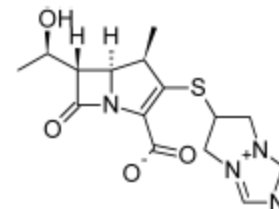
Refer to your book to read more about these •
compounds



Meropenem



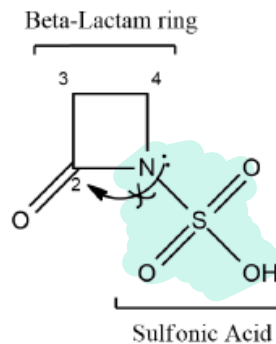
Meropenem metabolite



Biapenem

Single Ring *Monobactams*

- Monobactams –by their name- contain β -Lactam ring only without any other cycle, and as we said previously that β -Lactam ring is too stable here because the pair of electrons on nitrogen are in resonance with the carbonyl, so to decrease the stability, monobactam have a sulfonic acid at the nitrogen.
- Which means that the β -Lactam ring has been destabilized by the sulfonic acid which
- takes away the electrons, and this will prevent the resonance to occur –sulfonic acid destabilize the β -Lactam ring-



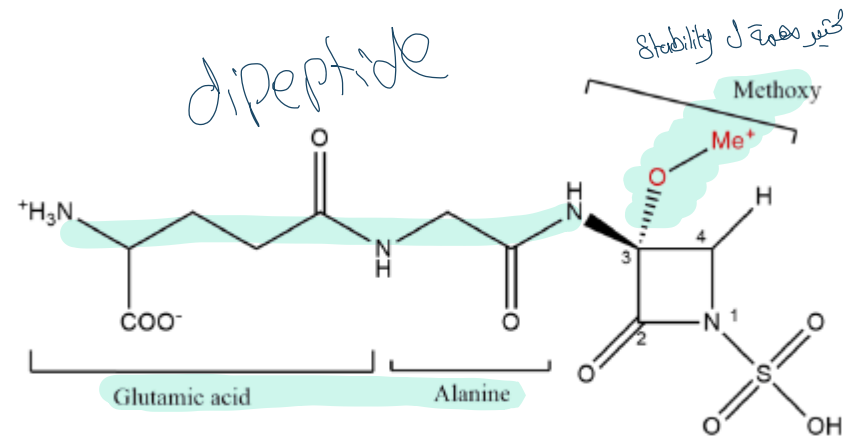
monobactam (without bind with sulfone group)
 المقرون هو stable بس ال resonance
 ولكن وجود S على N مع يحلله اكترونا = N
 ويمنع ال resonance وبالتالي يكون not stable

- monobactam (stable)
 - monobactam with (S-OH group) $\rightarrow$ not stable

- Monobactams:
- **1- Sulfazecin**
- 2- Aztreonam
- 3- Tigemonam
- Sulfazecin is a natural monobactam, at carbon #3 there is dipeptide –Alanine with glutamic acid- and methoxy, attached to the monobactam

Sulfazecin is:

- ❓ Orally inactive
- ❓ Chemically unstable (because of the protonation at the methoxy so it becomes a good leaving



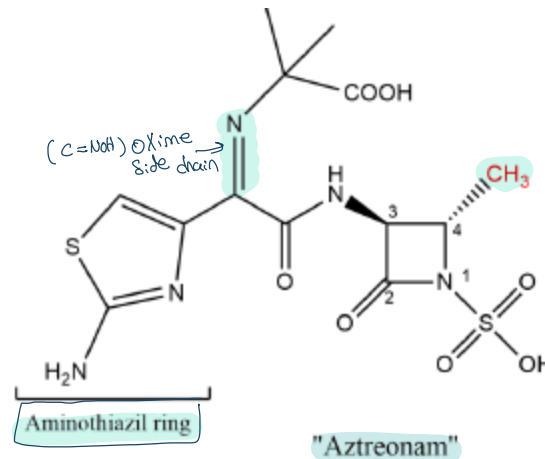
Sulfazecin not stable

2- Aztreonam

is mixed between
monobactam + cephalosporine

Aztreonam is an: ^{8 يتم امتصاصه لانه سلفوناميد} ^{sulfone group}

- Acid stable (but still not orally available)
- Used to treat local GI tract infections
- β -Lactamase resistant ^{مقاومة البكتيريا المتقدمة من cephalosporins}
- Gram negative antibacterial agent (only gram negative, not gram positive ⁺)
- because Aztreonam is too hydrophilic) – have an effect on *E. coli*, and no effect
- on *pseudomonas aeruginosa*, *streptococcus* and *staphylococcus*. not effect



3- Tigemonam

- Tigemonam is: *لديه بنية ال Cephalosporine*
- β -Lactamase resistant
- its spectrum of activity is the same as Aztreonam (against Gram -ve only except Pseudomonas)
- Orally stable (due to the hydrophilicity which is balanced here) *يختلف عن المركب، له قبل*

