

# Antibacterial antibiotics

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أدىها لأبوي الله يرحمه

# Antibacterial antibiotics

في عصر طبيعه

anticancer = antibiotics = chemotherapy agent

- A substance is classified as antibiotic if the following conditions are met:

تعريف ال antibiotics -

1. It is a product of metabolism even if it was later duplicated by synthesis
2. Synthetic analogues of naturally occurring antibiotics
3. It antagonizes growth or survival of one or more species of microorganisms (Either kills the microbe (microbiocidal) or prevent its growth (microbiostatic)).
4. It is effective at low concentrations

# Antibacterial agents:

Penicillinum natatum

- The accidental discovery of penicillin is by Fleming in 1928 is the main reason for the initiation of modern antibiotic era.

- Clinically useful antibiotics need to have the following criteria:

1. Combat infection or neoplastic disease
2. Selective toxicity → انما يقتل وفعال سائر  
بدون صدمه الانسان
3. Stable for a period of time inside the body.
4. Ease of administration by oral or parenteral route
5. Rates of biotransformation and urinary elimination are
  - 1 slow enough to allow convenient dosing schedule and
  - 2 rapid enough to remove the drug and its metabolites after discontinuation

لازم اذوار ال Penicillin G لانه 8-

1- narrow spectrum

2- Resistance

3- Break down by Beta lactemase

\*) مختصر مرق نقطة (2 و 5) انه ← لازم يكون عنا Balance Inside the body

# Potential targets for antibacterial agents

- ❑ Protein synthesis
- ❑ Nucleic acid synthesis
- ❑ Cell metabolism (e.g. folate synthesis)
- ❑ Cytoplasmic membrane
- ❑ Bacterial cell wall synthesis

# Potential targets for antibacterial agents

Sulfonamides  
(on metabolic enzymes)

Penicillins  
Cephalosporins

Aminoglycosides  
Tetracycline  
Chloramphenicol

Quinolones  
Rifampicin

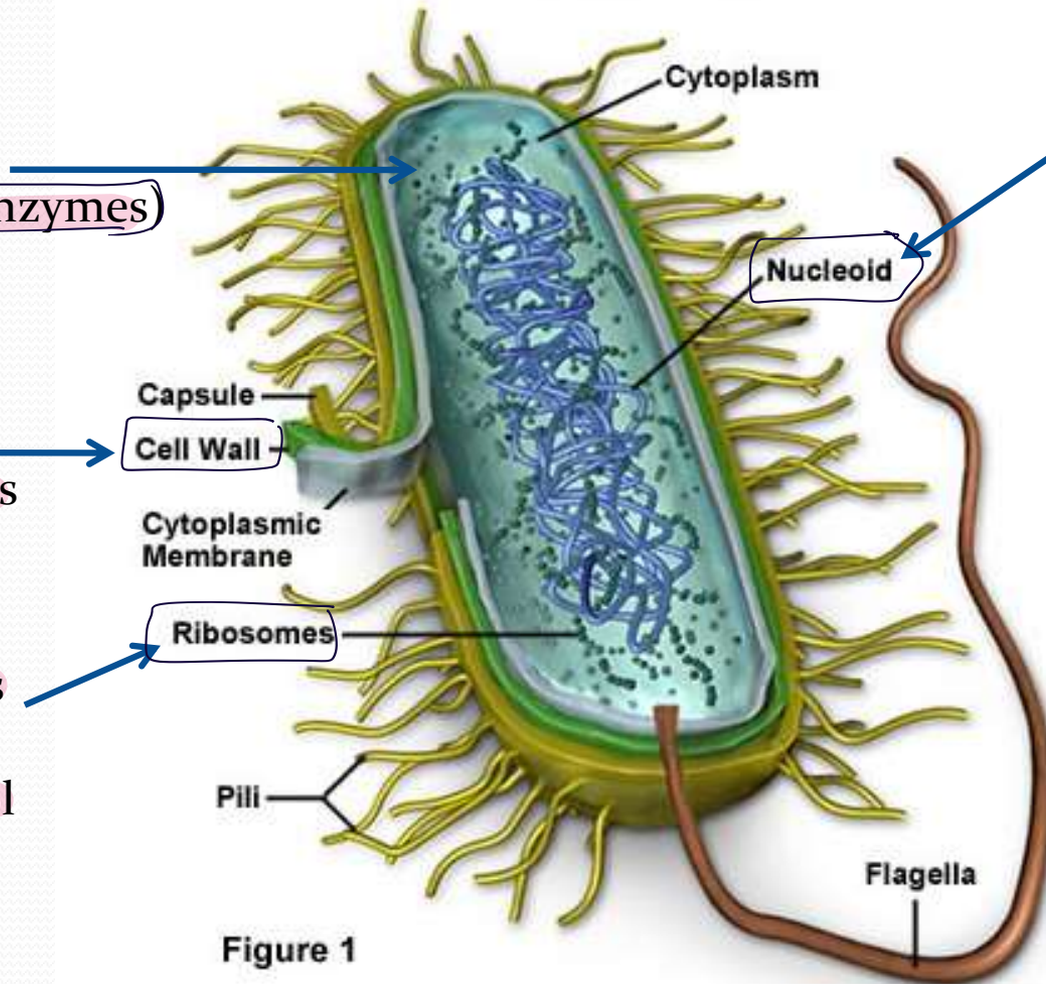
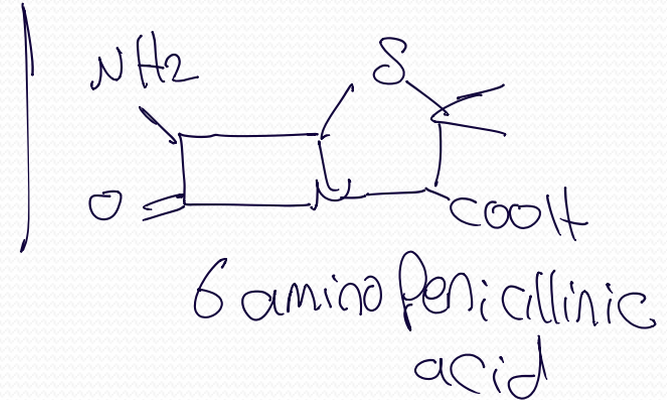
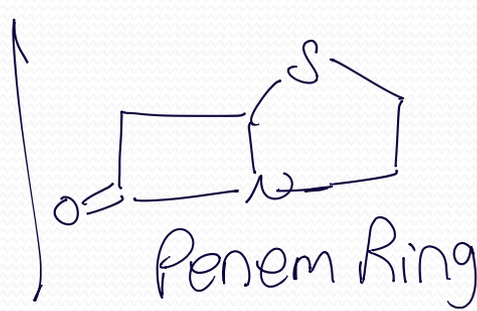
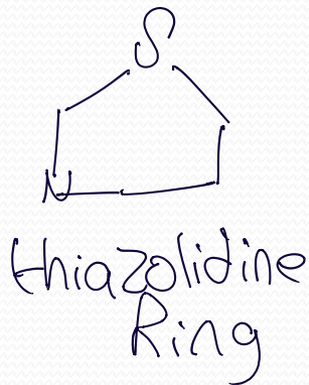


Figure 1

# Antibacterial agents acting on the cell wall biosynthesis

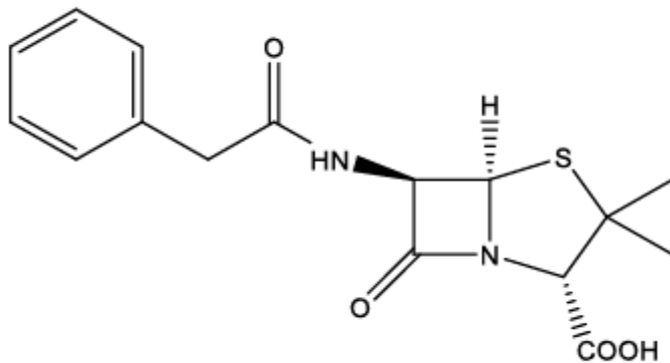
## Penicillins and Cephalosporins



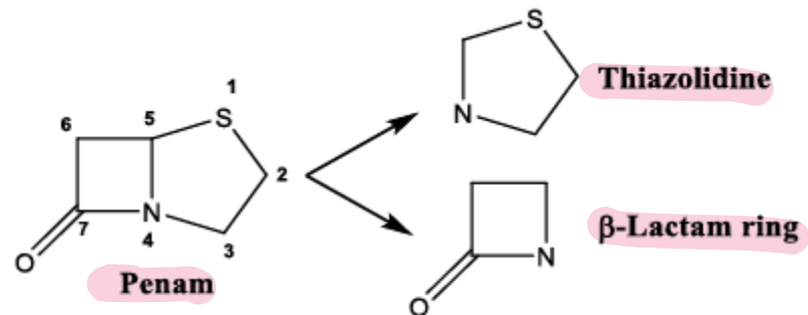
# $\beta$ -Lactam antibiotics

Lactame  $\rightarrow$  cyclic amide  
amide  $\rightarrow$   $\text{C}=\text{O}$  و  $\text{N}$  في الحلقة  
Lactone  $\rightarrow$  cyclic ester  $\rightarrow$  

- These **includes** both penicillins and cephalosporins.
- The name **“lactam”** is given to cyclic amides and is analog to the name **“lactone”**, which is given to cyclic esters.
- This ring ultimately proven to be the main component of pharmacophore. (Beta lactame Ring)
- This ring is more reactive and sensitive to nucleophilic attack when compared to normal planar amides.



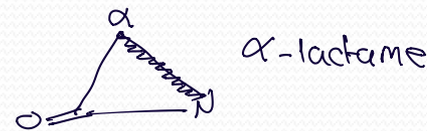
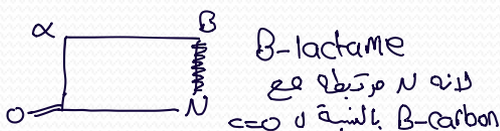
**Benzylpenicillin (Penicillin G)**



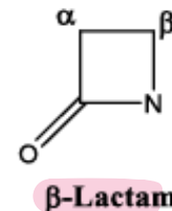
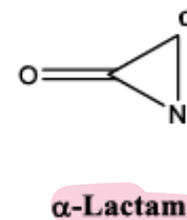
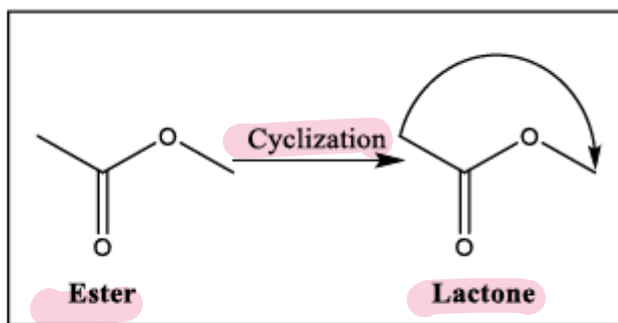
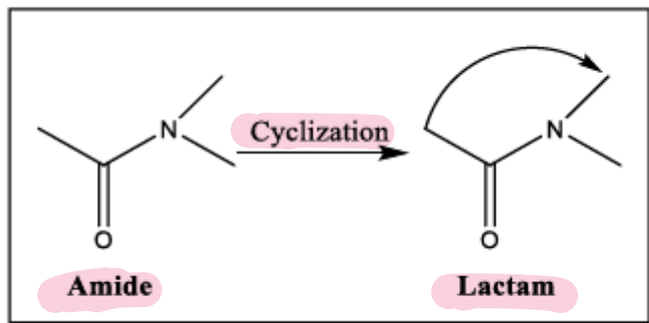
# $\beta$ -Lactam (the cyclic amide)

This ring is the result of cyclization of the amide group, as in case of Lactone which is the cyclic structure of ester.

Why it's called  $\beta$ -Lactam?



Because the first carbon beside the carbonyl group is  $\alpha$  carbon and beside this  $\alpha$  carbon is  $\beta$  carbon, so when the nitrogen atom substituting the  $\beta$  carbon we call it Lactam ring ( $\beta$ -Lactam), also there is  $\alpha$ -Lactam rings

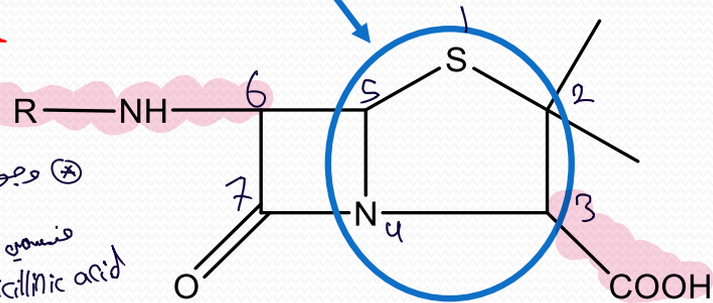


# $\beta$ -lactam antibiotics

**A**

5-membered thiazolidine ring

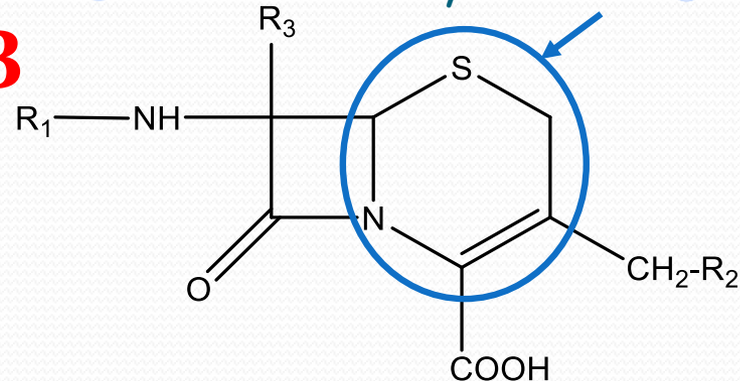
⊗ وجود ثغرابا على  
C3+C6  
فسيكون المركب  
6-aminoPenicillanic acid



Penicillin nucleus

**B**

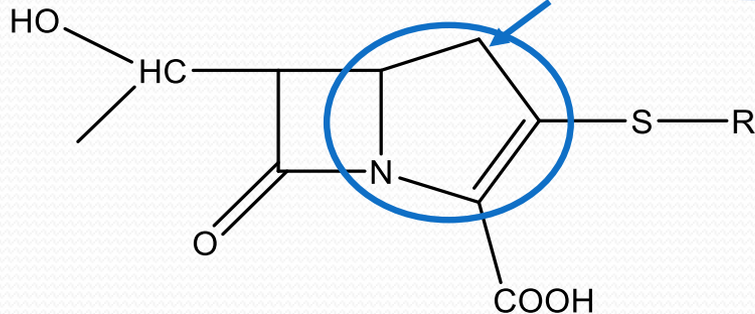
6-membered dihydrothiazine



Cephalosporin nucleus

**C**

Carbon atom

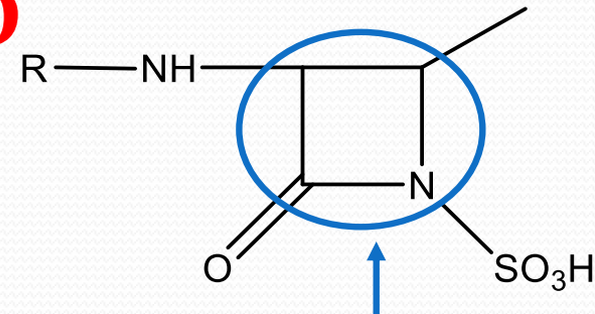


يستخدم في التهابات = Soft tissue

Carbapenem nucleus

**D**

monocyclic

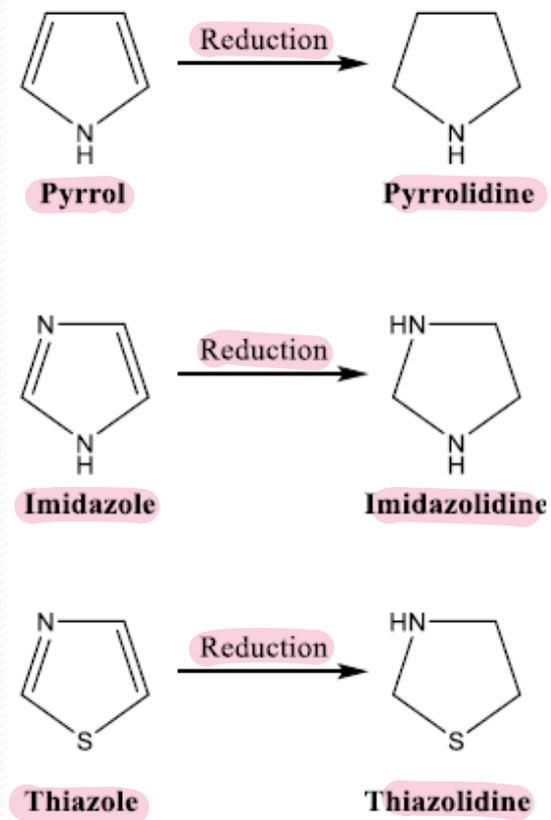


Monobactam nucleus

يتحول المركب مشبع

**Thiazolidine ring:** (S-containing N-containing reduced heterocyclic ring)

- Any **heterocyclic compound** with **5-membered ring** ends with "**ole**", so we have for examples Pyrrole, Imidazole, and thiazole.
- Upon reduction these compounds will be Pyrrolidine, Imidazolidine, and Thiazolidine (thia=sulphur, aza= nitrogen) respectively.

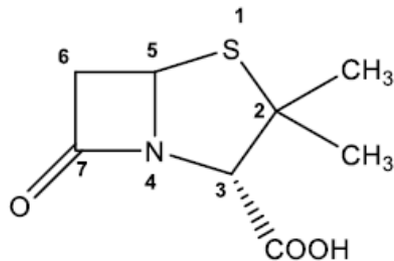


- To say that our compound is penicillin, firstly you have to look for the Penam system then the following

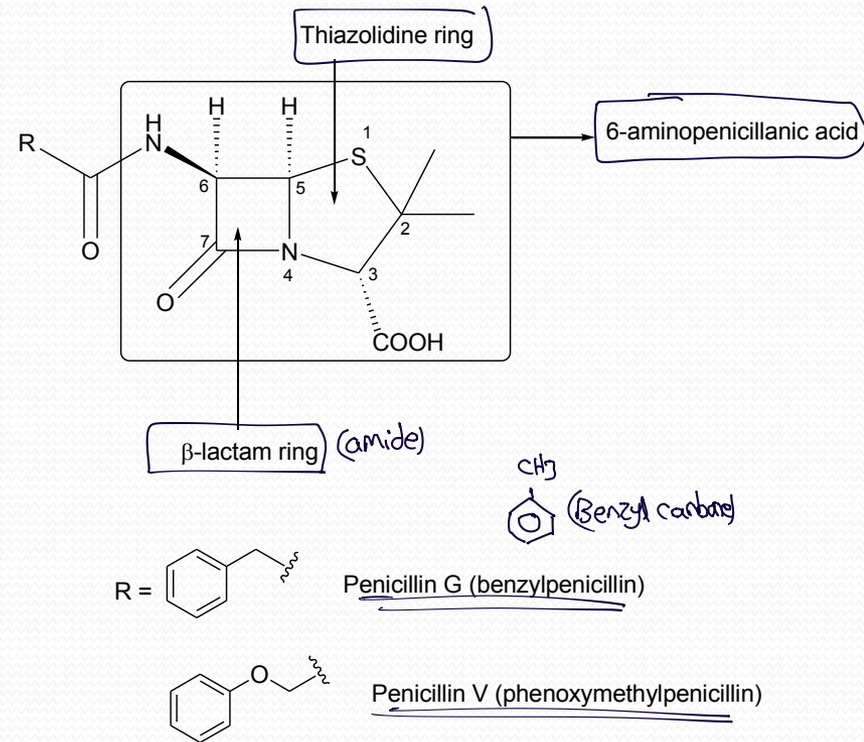
substitutions also must present in all penicillins:

- Di-methyl group at the position 2.
- A carboxylic acid at position 3, below the plan, (S) oriented.

Until now, this compound is called "Penicillanic acid", a 2, 2-dimethyl-3-carboxy Penam.

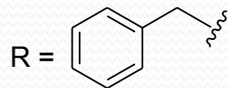
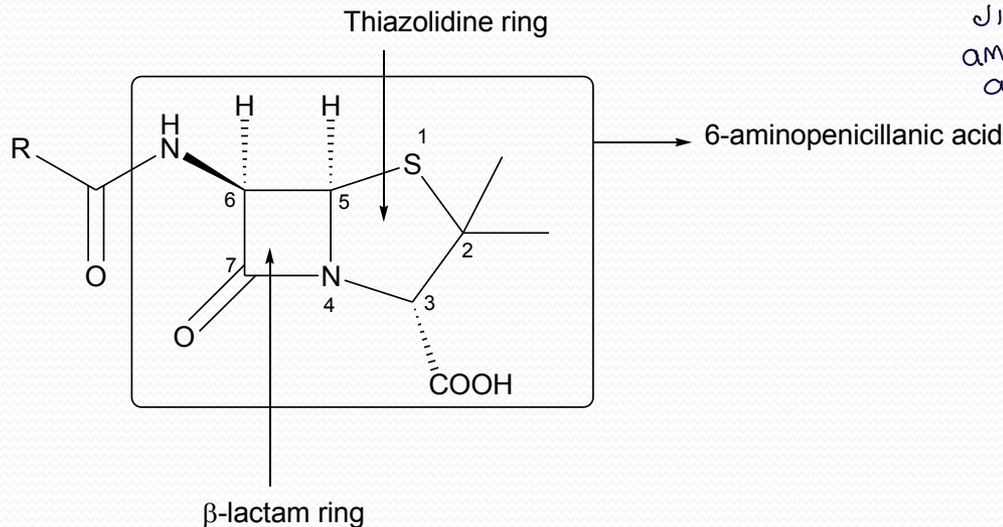


**Penicillanic acid**

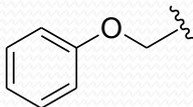


# Penicillin's naming is problematic

1<sup>st</sup> naming system related to the chemical abstracts



Penicillin G (benzylpenicillin)



Penicillin V (phenoxymethylpenicillin)

سؤال من اين يتصنع

فان ال  
amino  
acid

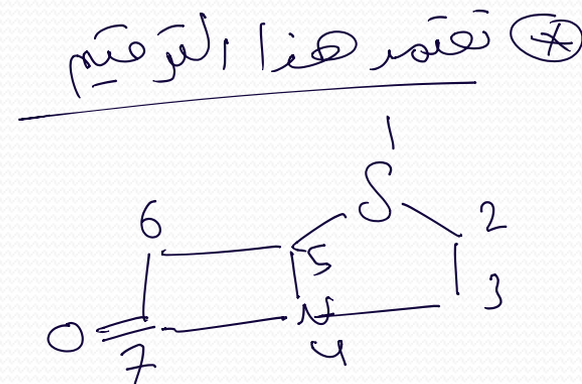
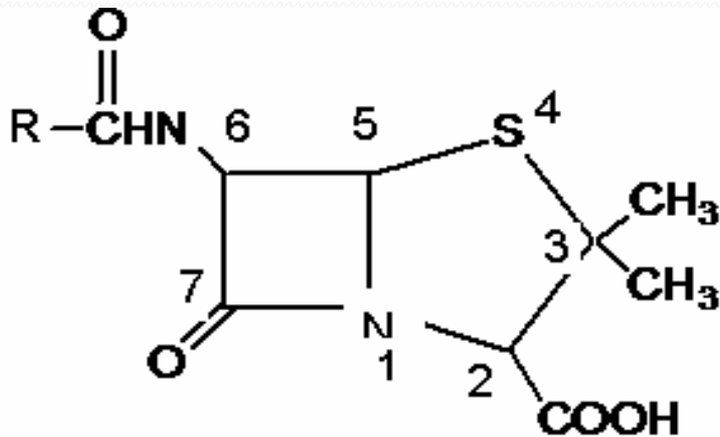
**Fungus synthesizes**  
penicillin using cysteine,  
valine and some of the  
fermentation products

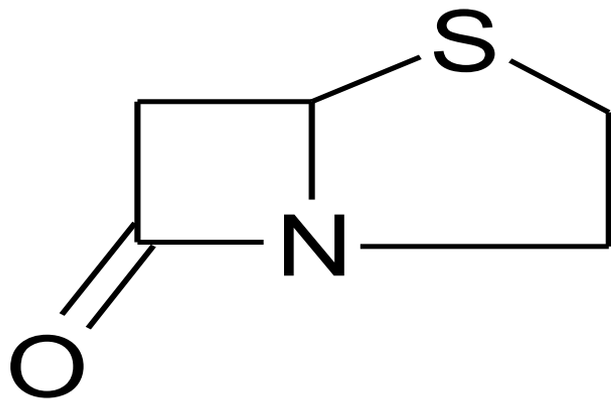
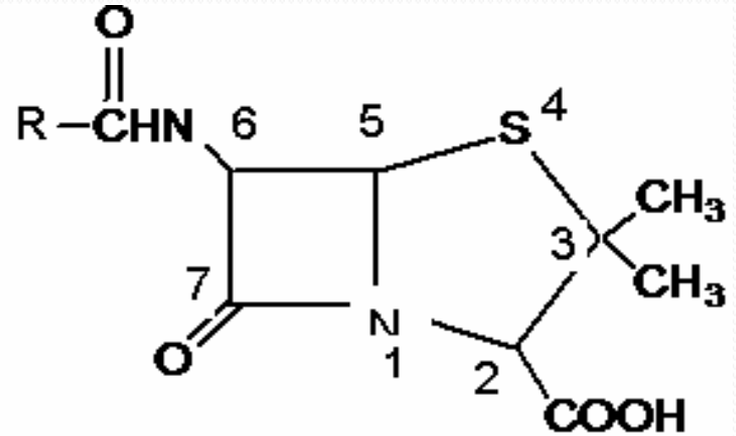
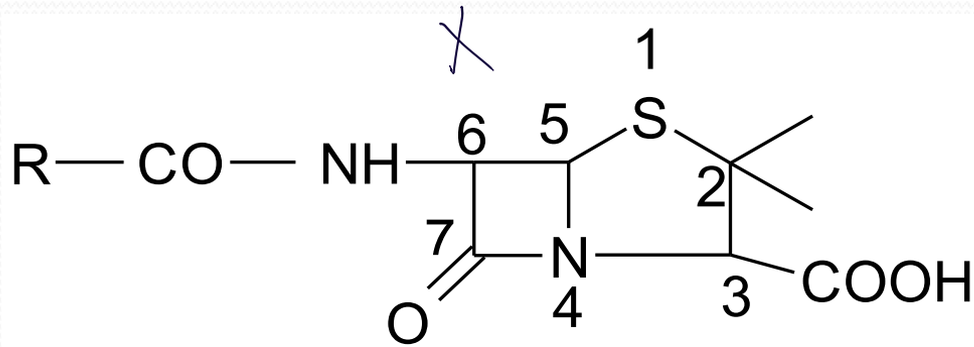
**Difficult to synthesize**  
**in the lab due to:**

- The unstable highly  
1. strained ring system.  
2. The three chiral  
center it has which  
should be with certain  
stereochemistry

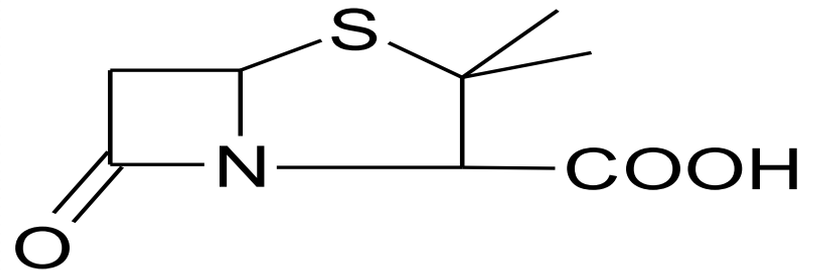
# Penicillin's naming is problematic

- 1st naming system related to the USP:
- The correct IUPAC name of penicillin is 4-Thia-1-azabicyclo[3.2.0] heptanes





Penam



Penicillanic Acid

# Stereochemistry

⊗ 3 chiral center &

C3 / C5 / C6

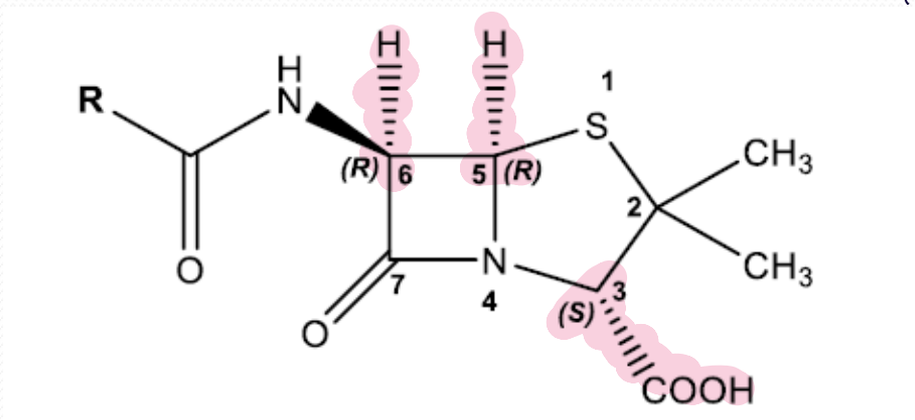
\* C3 → S

\* C5 → R

\* C6 → R

\* C3 + C5 → Sis of each other

\* C3 + C6 → trans of each other



\* ما بقدر اصفه في الاختير = لانه اله

change conformation

1- unstable strain Ring

2- three chiral center

It contains three chiral carbon atoms at C<sub>3</sub>, C<sub>5</sub> and C<sub>6</sub>.

C<sub>6</sub>-L configuration, C<sub>3</sub> and C<sub>6</sub> chiral centers are trans to each other.

All synthetic and semi synthetic penicillin have same absolute configuration that of natural

3S:5R: 6R

⊗ الإنسان فا كنده cell wall اما في البكتريا فيه ويختلف حسب Gram(+)/Gram(-)

1 Gram(+) ← كنده multi layer من peptidoglycan مع cross-linked

Peptidoglycan يتكون من (D-alanine + NAM + NAG) ← يكونوا قسايبين مع بعض يواظف  
انزيم trans peptidase ← يحص phospholipid تبعه اكلد

2 Gram(-) ← كنده Single layer of peptidoglycan وبالماء Purins (Water filled)

⊗ لازم الحماه الكيوي لحتى يعبر Purin لازم له حقا = hydrophilic فين  
(soluble) R-group is hydrophilic

# penicillins

بدي يرتبط معو لحتى يعطي التأثير هو (target)

transpeptidase

لما يرتبط Penicillin مع transpeptidase  
يعمل irreversible covalent bond

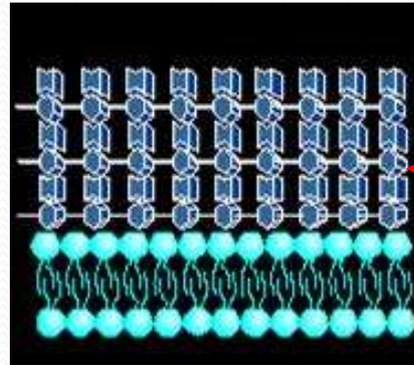
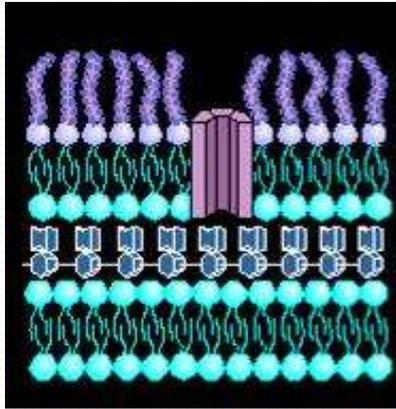
inhibition of cross link ← مع يرفد الماء لافك  
بسب ضغط الاسموزي ويسير his swelling ثم  
cell lysis (تمو = فلالا)

بدي افغ انه يرتبط معو

B-lactamase (penicillinase)

يعمل القليه مع Penicillins  
يعمل covalent bond

# The bacterial cell wall



Peptidoglycan

-  N-acetylglucosamine (NAG)
-  N-acetylmuramic acid (NAM)
-  D-alanine

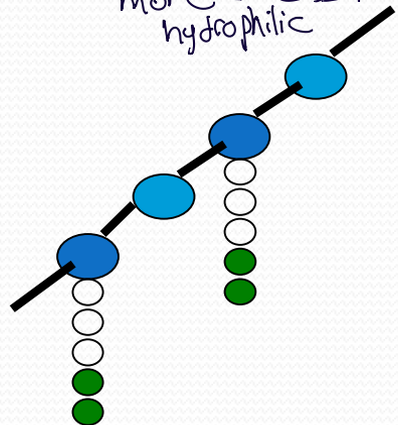
**Gram -**

Only two layers of peptidoglycan

ملحقة اقلية البعاء  
بأثر على  
Gram (-)  
لازم يكون  
more ← hydrophilic

**Gram +**

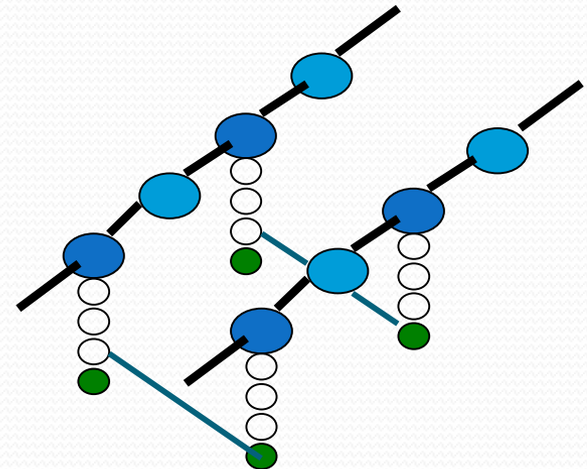
Consists of 50-200 peptidoglycan layers



Transpeptidase

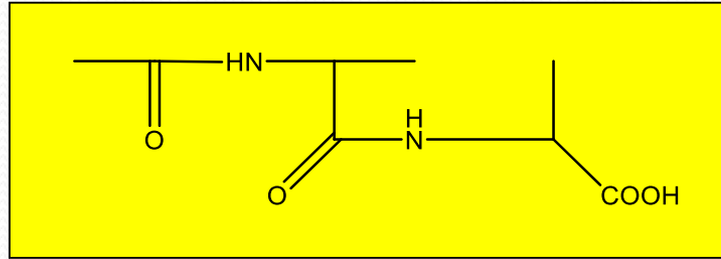


Involved in cross-linking



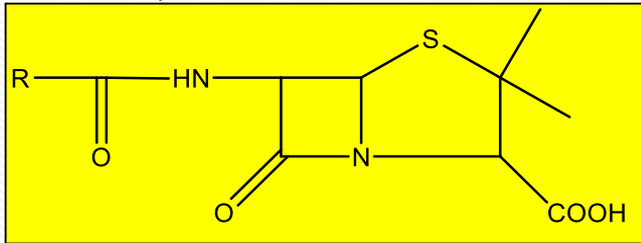
\* طريقة العمل ← Penicillin يمتص بسهولة في الخلية

(D-ala) يرفع الـ Penicillin يرتبط مع الإنزيم transpeptidase  
 ويرتبط بـ (D-ala) (irreversible covalent bond)

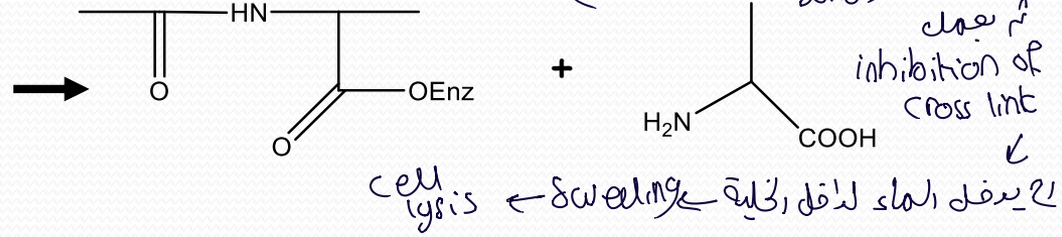


D-ala D-ala (natural substrate)

Same structures



Penicillins



Penicillin-enzyme complex

cross-linking inhibited

The wall become fragile and can no longer prevent the cell from swelling and bursting

Bacterial cell lysis

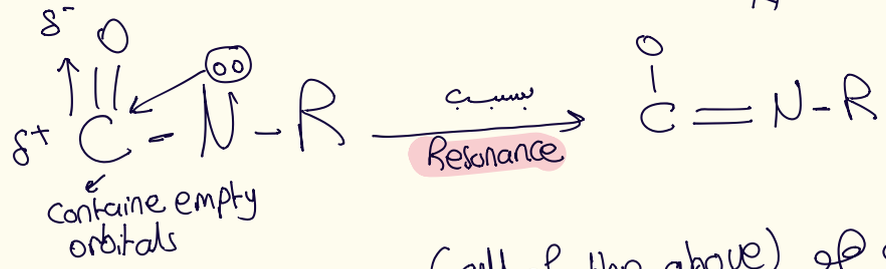
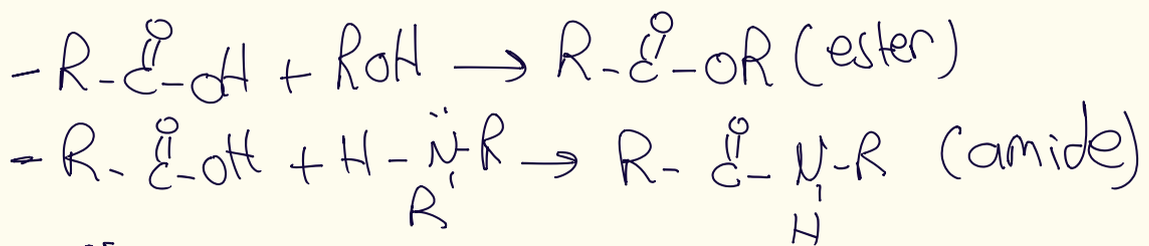
Excellent selective toxicity

لأنه (1) ينتقى الـ target يستهدف الـ Bacteria  
 (2) لأنه فيه الـ Dextro amino acid ← الـ Bacteria  
 أما فيه الـ L- amino acid ← الـ human

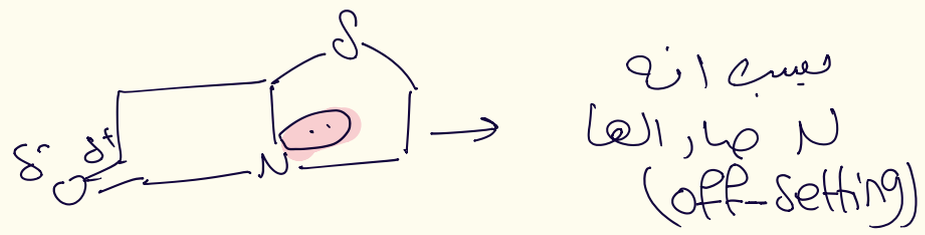
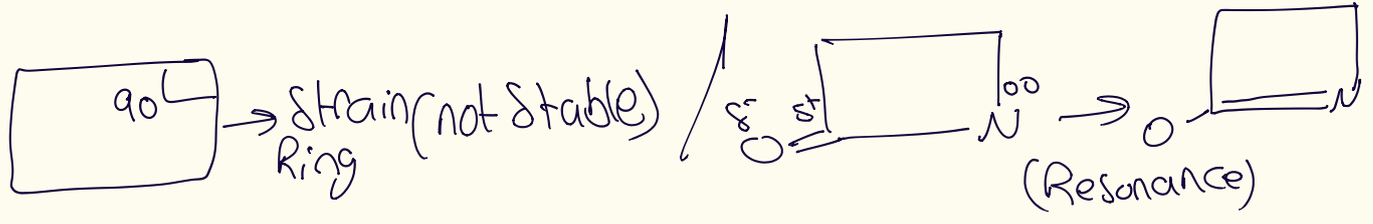
- Penicillin mimic the structure of D-ala-D-ala, because of that the transpeptidase mistakenly bind to it instead of D-ala-D-ala. ← (Selective Toxic) دواء انتقائي (X)
- Also this explains the lack of penicillin toxicity, since D-amido acids are not present in human, only the L-amino acids present.
- Also targeting the cross linking in the peptidoglycan biosynthesis which is only present in bacteria explains the selective toxicity.

⊗ β-lactam ← هي حلقة، باقية، وهي حلقة مشغولة فيها (Strain)  
 زواياها 90°، الوضع طبيعي لا يتم تكون زوايا بين C 2 تقريباً 107°  
 فتعتبر not stable

⊗ why amide bond more strong than ester?



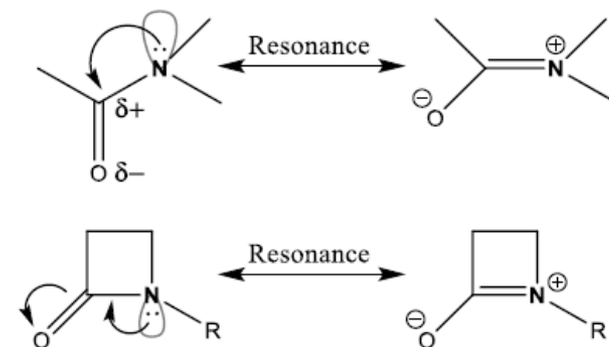
⊗ بجوابه سؤا ال امتحان هو (all of the above)



⊗ Strong bond → ester < amide < five-membered Ring  
 ← سبب Resonance  
 ← سبب انق off setting of N

# chemical properties of $\beta$ -Lactam ring:

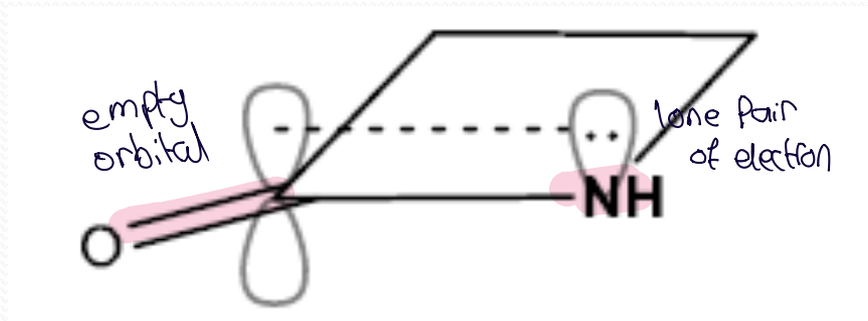
- In the original amide group (Linear amide), the Nitrogen atom that is <sup>stable</sup> beside the carbonyl group has a pair of unshared electrons which causes the Resonance between the N and the Carbonyl group (as we see in the periodic table, the O is more electronegative/electrophilic than C so the O in the carbonyl group withdraws electrons to have partial negative charge and C partial positive charge) So this carbonyl system is "Di-polar" and withdraws electrons from N.
- this Resonance is also applied for isolated  $\beta$ -Lactam ring (alone, no other substitutions) and the pair of electrons of N are not far from the carbonyl group



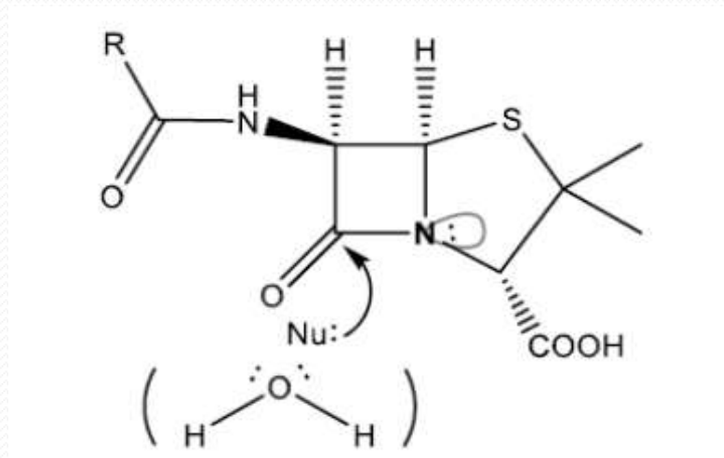
electrophilicity				
increase				
→				
C	N	O	F	↑
Si	P	S	Cl	
			Br	
			I	

## chemical properties of $\beta$ -Lactam ring:

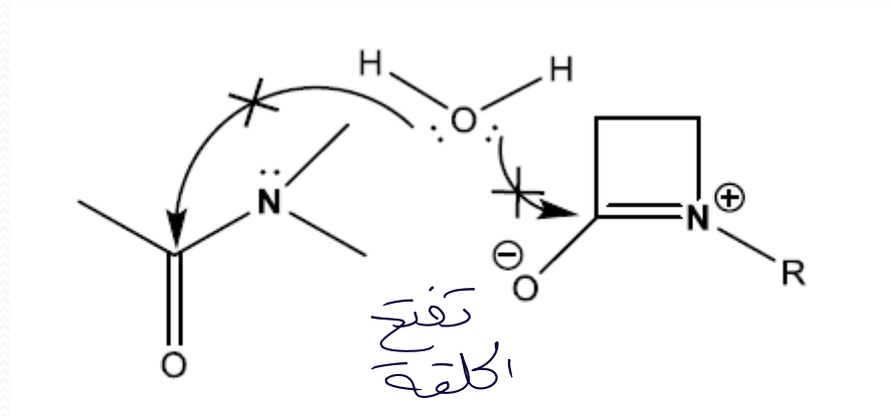
- Look at the pair of electrons of N in the  $\beta$ -Lactam ring alone
- (**NOTE** that the empty orbital of carbonyl group accepts the pair of electrons of N to make resonance)



- BUT when the  $\beta$ -Lactam is involved in the Penam system, and look at it in 3D view,
- the pair of electrons of nitrogen is far from the carbonyl group because the
- Thiazolidine ring bends this pair of electron (**Off-setting**) and the N becomes
- pyramidal (very close to the amines).
- So the Penam is completely different from amide, the carbonyl group is very electronegative and easily attacked by a nucleophile such as water (**No resonance** to make this carbonyl rich in electrons and resistant to hydrolysis as in amides that need enzymes to be hydrolyzed).



- **NOTE:** the  $\beta$ -Lactam alone is also resistant to hydrolysis as the amides due to resonance but easier than amides (the Nucleophilic attack is more easy).
- For that reason, when one of the penicillins is given orally as solution, it can be used only for one week after dissolving in water due to hydrolysis and must be stored in the fridge to slow down the hydrolysis rate by water (which is a nucleophile, contains two pairs of electrons).
- So all penicillins are unstable in water and tend to breakdown quickly in the aqueous conditions, and for that reason mostly these preparations are made as suspensions not solutions to reduce the water activity. This is also applied for parental penicillins that are given as re-constitutable powders.

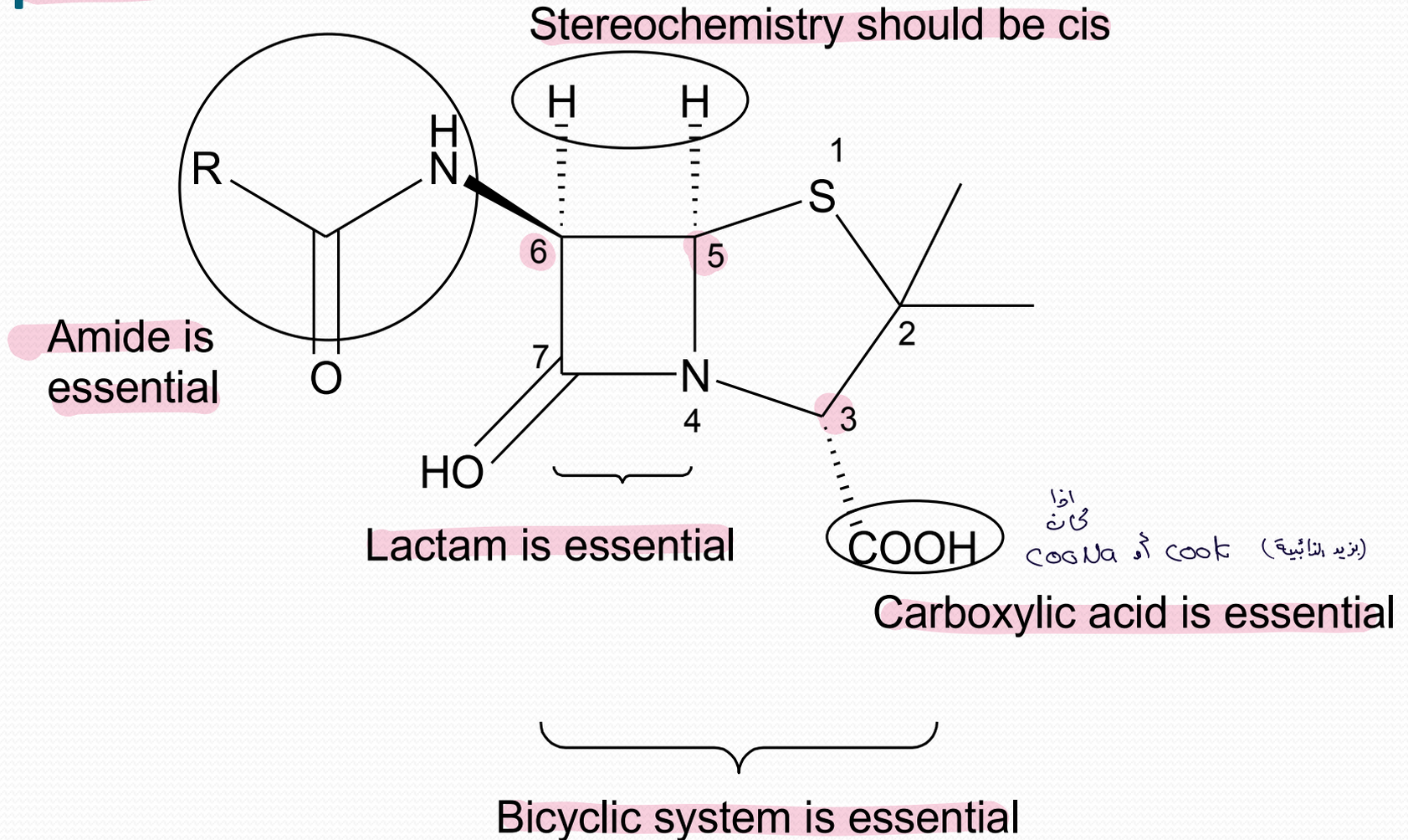


# Structure-activity relationships of penicillins (SAR)

مفوطا (90°)

- The strained  $\beta$ -lactam ring is essential.
- The free carboxylic acid is essential (the carboxylate ion binds to the charged nitrogen of the lysine at the active site).
- The bicyclic system is essential. → monocyclic ال ٨  
وهي مجموعة  
مختارة جدا
- The acylamino side chain is essential.
- Sulfur is not essential. → Carba penem  
لانه في بعضه الامكان تكون  
ببدال S فيك
- The stereochemistry of the bicyclic ring with respect to the acylamino side chain is important.

# Structure-activity relationships of penicillins



# Reasons for the acid sensitivity of penicillin G:

الهكرك (وجوده في الحلقة تمنعها من الف)

- Ring strain: due to the large angle and torsional strain exist, acid catalyzed ring opening will relief these strains.
- A highly reactive  $\beta$ -lactam carbonyl group:
- This amide bond is exceptionally unstable compared to the normal amide, because it is a 4-membered ring this will increase the angular and torsional strain

(\*) Problems of Penicillins G-

- (1) acid sensitive ( only parenteral administration)
- (2) Resistance
- (3) narrow spectrum only on Gram (+)

# Chemical degradation and properties

- The early penicillins were yellow to brown in color and very unstable so refrigeration was required to maintain activity for short time مفعلة nucleophilic attack  
مفعلة لبيته
- Unpleasant taste
- Strongly dextrorotary مفعلة ديفي
- Most penicillins are acidic  $Pka=2.5-3.0$  some are amphoteric acid / base
- The free acids are not suitable for parenteral or oral administration. Na and K salts are suitable to allow both oral and parenteral administration.
- Some salts of penicillins with organic bases such as Benzathine, Procaine and Hydrabamine have limited water solubility and so they are suitable as depot to provide effective conc. Of penicillin for long time to treat chronic infections → ابرة الروفاتبزم  
بعطيها بالحقن  
مبصير (low release)

⊗ Na/K → more soluble

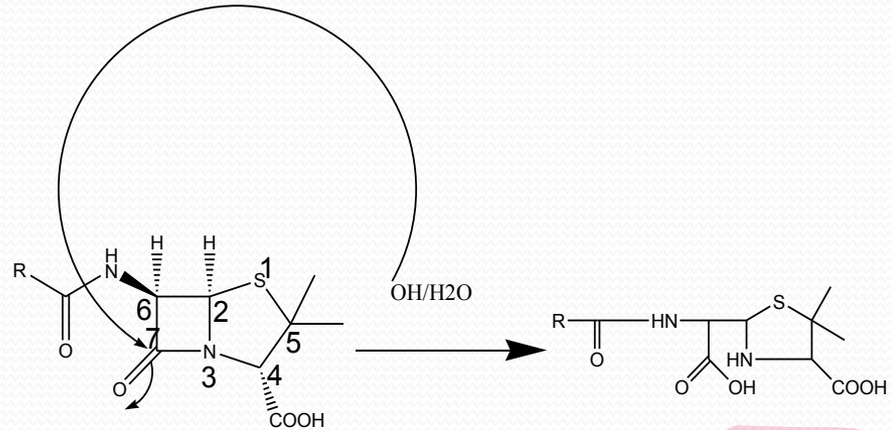
⊗ Benzathine/procaine/hydrabamine → less soluble

# Chemical reactivity

## 1. reactions with nucleophiles $\text{-OH}$ , $\text{H}_2\text{O}$ , $\text{NH}_2\text{-OH}$ , $\text{R-NH}_2$ , $\text{R-OH}$ , and body proteins

### 1. reaction with $\text{-OH}$

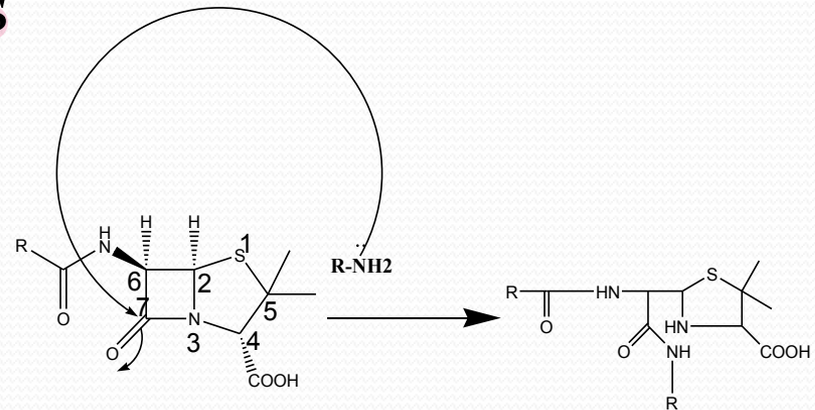
↳ give Penicillinoic acid



**Penicilloic acid**  
In active, stable in alkaline, not stable in acid

### 2. Reaction with alkyl amines

↳ give Penicillinoic amide

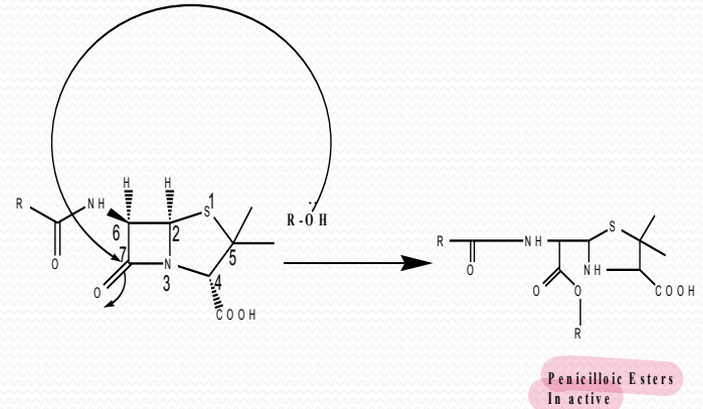


**Penicilloic Amid**  
In active

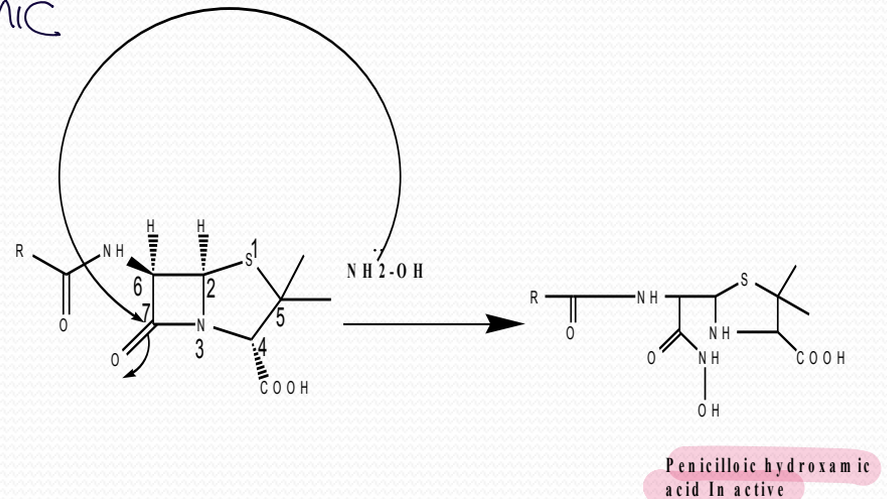
# Chemical reactivity

1. reactions with nucleophiles  $\text{-OH}$ ,  $\text{H}_2\text{O}$ ,  $\text{NH}_2\text{-OH}$ ,  $\text{R-NH}_2$ ,  $\text{R-OH}$ , and body proteins

- 3. reaction with Alcohol  
↳ give Penicillinoic ester



- 4. Reaction with hydroxyl amine  
↳ give penicillinoic hydroxamic acid



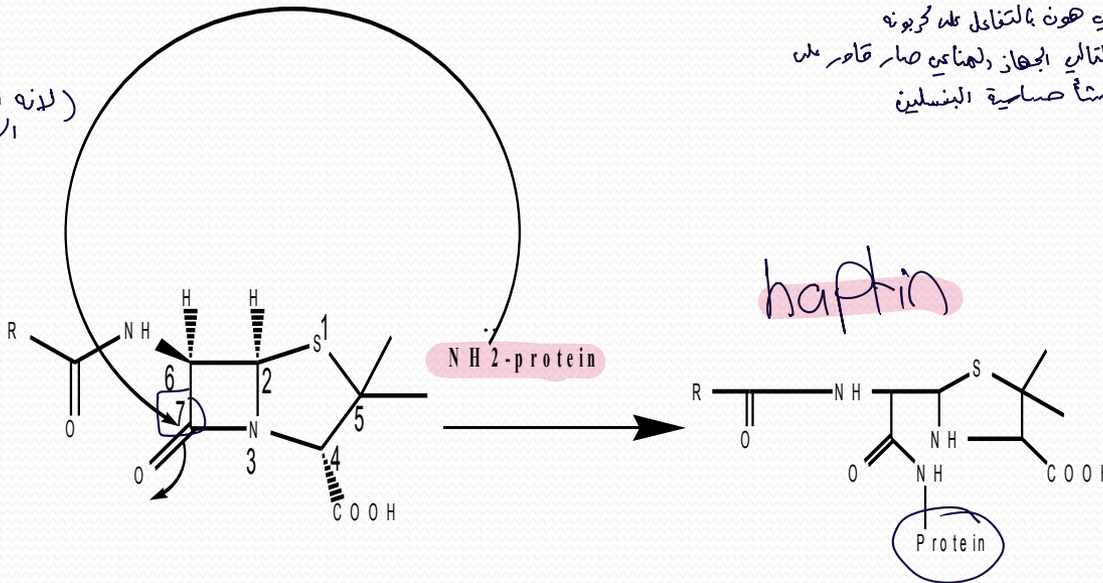
# Chemical reactivity

## 1. reactions with nucleophiles -OH, H<sub>2</sub>O, NH<sub>2</sub>-OH, R-NH<sub>2</sub>, R-OH, and body proteins

- 5. reaction with body proteins: the nucleophilic attack on B-lactam rings by body proteins (specific) generate penicilloyl proteins that are suspected to be the reason for the allergic reactions to penicillins

في الواقع الطبيعي ال Penicillin جدا صغیر لدرجة انه جهاز المناعة غير قادر على رؤيته ولكن اذا ال nucleophiles الموجودة بالبروتينات حاصلة اسيامنا ها جمة ال Penicillins زي هون بالتفاعل على كربونه رقم (7) ال بييس حجمه كثير كبير وبالتالي الجهاز المناعي صار قادر على تمييزه بهايه طريقة تنشأ صلاية البنسلين

علاج حساسية هذه allergic  
cortisone  
+ epinephrine (لانه نزل الحفظ)

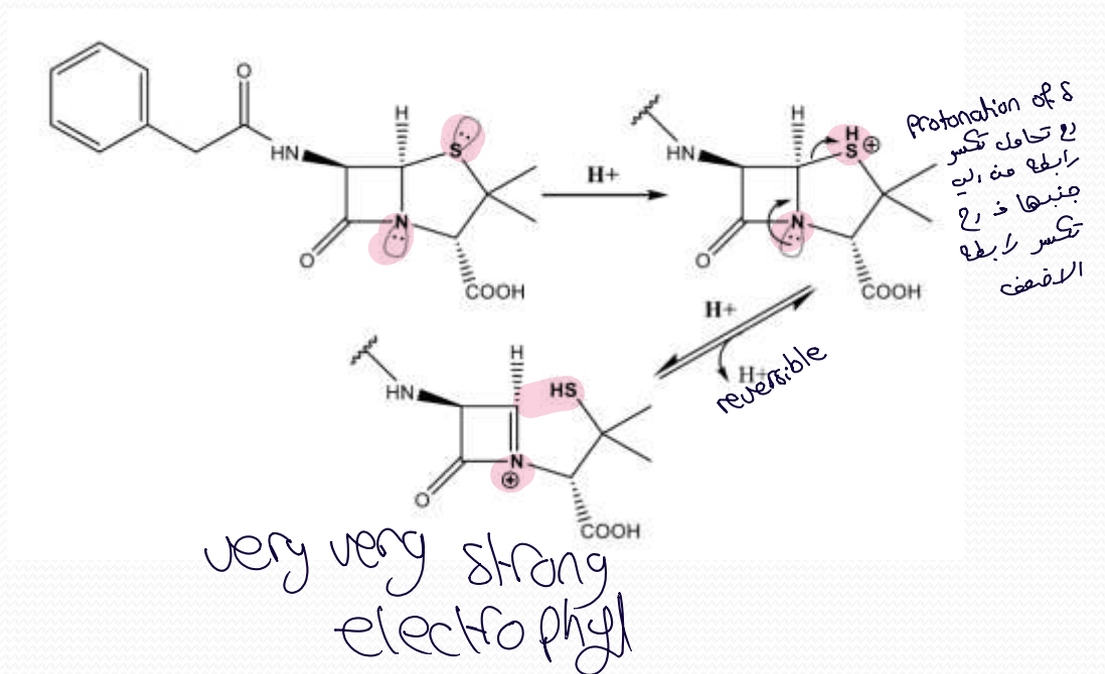


# Chemical reactivity

## 2. reactions with Acids

- In strongly acidic media ( $\text{pH} < 3$ ), penicillins undergo complex series of reactions as in the following:
- First step (reversible)

protonation of the sulfur atom will occur then a partial positive charge will be formed on S and becomes a good leaving group, the pair of electrons of N will move causing the structure to become with a thiol group and a double bond, and the N becomes quaternary with a positive charge.

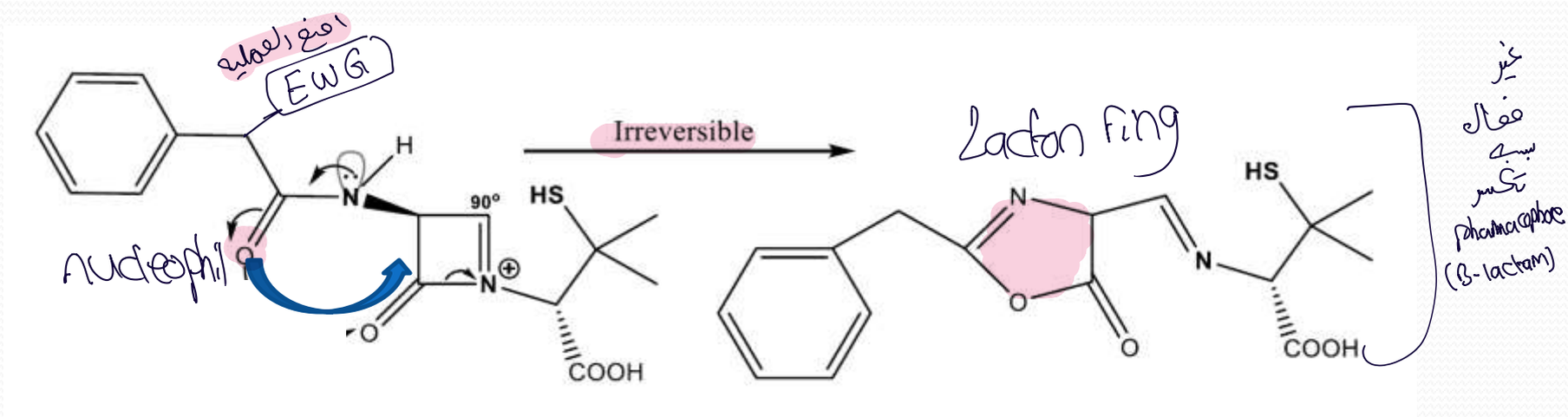


المشكلة الاولى هي  
acid Resistance

# Chemical reactivity

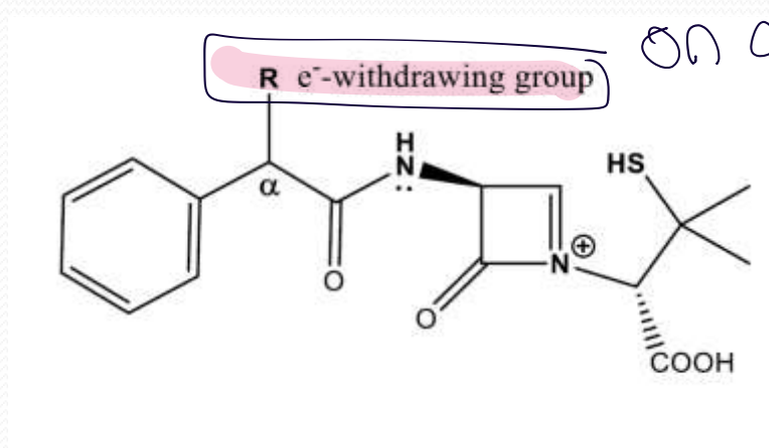
## 2. reactions with Acids

- Second step (irreversible)



the oxygen becomes a strong nucleophile

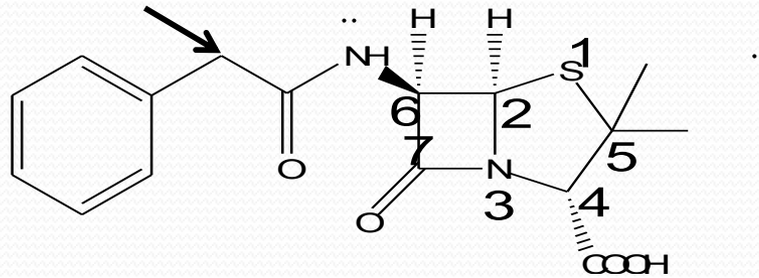
- And by knowing this mechanism, we can actually make "**Orally active penicillins**" How?
- if we take the electrons from this oxygen and prevent this attack we will prevent this step, then you have to add an electron withdrawing group at the  $\alpha$  carbon to the amidic carbonyl ( oxygen, amine).



# Acid sensitivity of penicillins

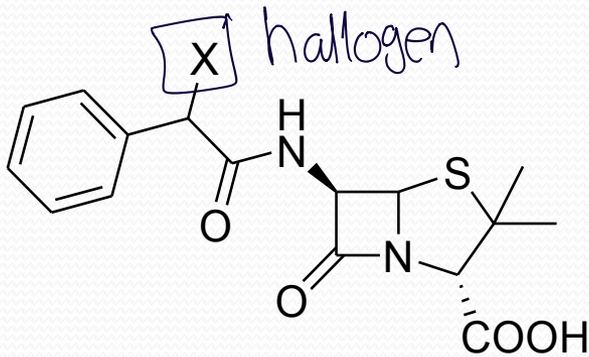
- Electron withdrawing groups in the  $\alpha$ -position of Benzyl penicillin will improve acid stability clearly

$\alpha$ -position

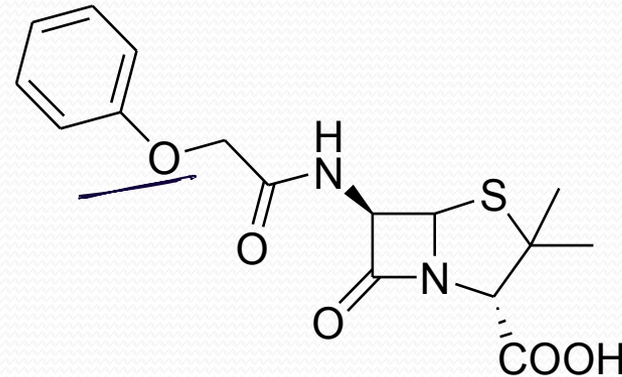


# Acid sensitivity of penicillins

Accordingly the following compounds are significantly more stable than Benzyl penicillin:

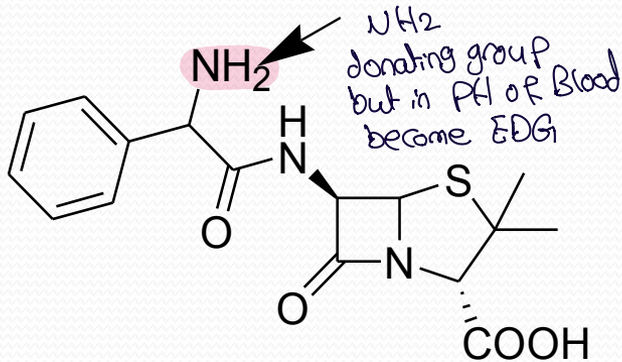


alpha - Halo benzyl penicillin (X=Cl, Br, I)

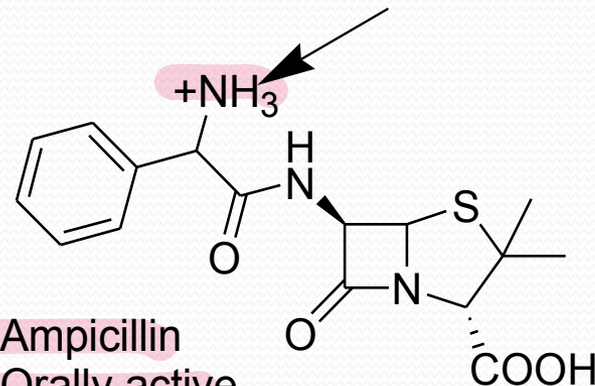


Phenoxy methyl penicillin: Penicillin V  
Acid resistant  
can be given orally

In plasma PH  $\text{NH}_2$  will become ionized to  $\text{NH}_3^+$  so it will become electron withdrawing

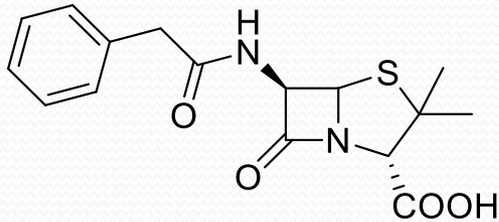


Plasma PH

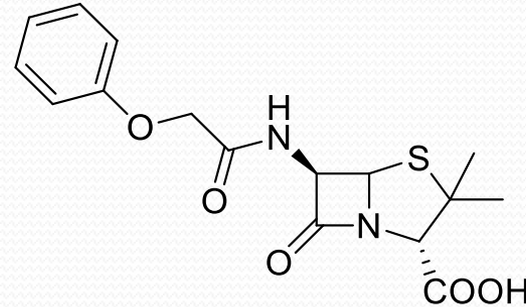


Ampicillin  
Orally active

# Acid resistant Penicillins

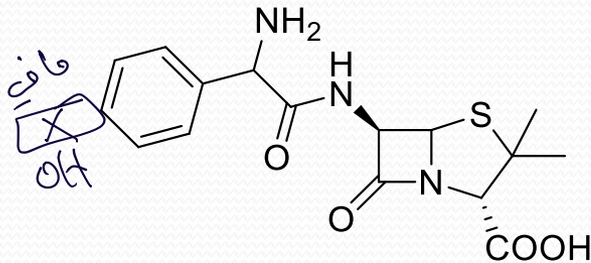


Penicillin G  
Acid labile  
Can not be given orally

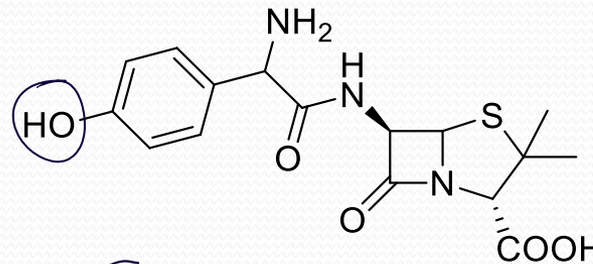


Penicillin V  
Acid resistant  
can be given orally

⊛ aminopenicillin (Amoxicillin) Ampicillin) more soluble than Benzylpenicillin



Ampicillin  
Acid resistant  
Orally active



Amoxicillin  
Acid stable  
Orally active

Amoxicillin: given once each 8 hours (longer half life).  
Ampicillin: given once each 6 hours

الاقطار بينهم  
OH



انواع المقاومة، المقاومة  
Resistance

# Bacterial Resistance:

- Two types:
- 1. Natural (innate) resistance, this is particularly important in gram negative (G-) bacteria mediated by the reduced permeability of the outer cell envelope of Gram negative bacilli which is linked to the cell wall via the peptidoglycan, such cell envelope is not present in gram positive bacteria
- 2. Other normally resistant bacteria can develop resistance by generating resistance enzymes by mutation or natural selection

# Bacterial Resistance:

- The second type of enzymatic resistance is the most common resistance mechanism.
- The resistant enzymes are collectively known as “penicillinases” and are of two general types:
  - 1. B-lactamases (most important)
  - 2. Acylases

\* Gram (+) → extra cellular  
\* Gram (-) → Peri plasma

# $\beta$ -lactamase (Penicillinase):

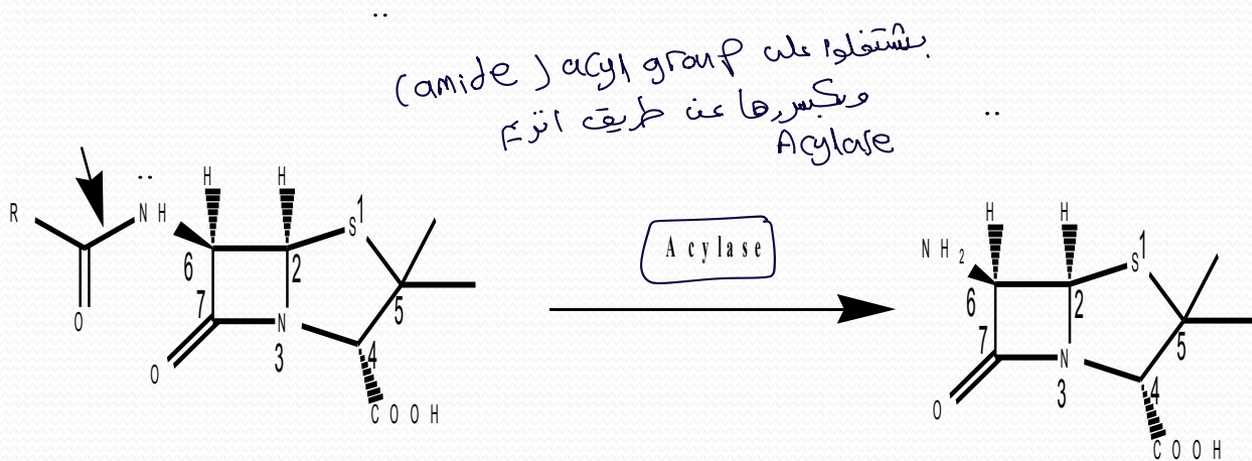
- $\beta$ -lactamase break the  $\beta$ -lactam ring, they are either chromosomal or plasmid, constitutive or inducible depending on particular species: التي بتتوفرها  
التي الاثرية
  - Gram +ve *S.aureus*: 1. inducible  $\beta$ -lactamase  
2. synthesized at cell wall and released extracellularly  
95% of *S.aureus* became resistant to penicillins
  - Gram -ve bacillic 1. constitutive R-factor  
2. Cytoplasmic enzymes
- Again  $\beta$ -lactamase from different species are different in structure and properties.

# $\beta$ -lactamase (Penicillinase):

- Gram +ve bacteria normally release  $\beta$ -lactamase to outside of the cell that will cleave penicillin before reaching the bacteria.
- Gram -ve bacteria release  $\beta$ -lactamase into the periplasmic space, which again will cleave penicillin before reaching the plasma membrane.
- Penicillin has to reach the plasma membrane where the transpeptidase present to do its antibacterial action.
- Most of gram -ve bacteria are  $\beta$ -lactamase producing bacteria

## 2. Acylases

- These enzymes can hydrolyze the acylamino-side chain of penicillins



6-Amino penicillanic Acid (6-APA)

# Rules to create Penicillinase resistant Penicillins

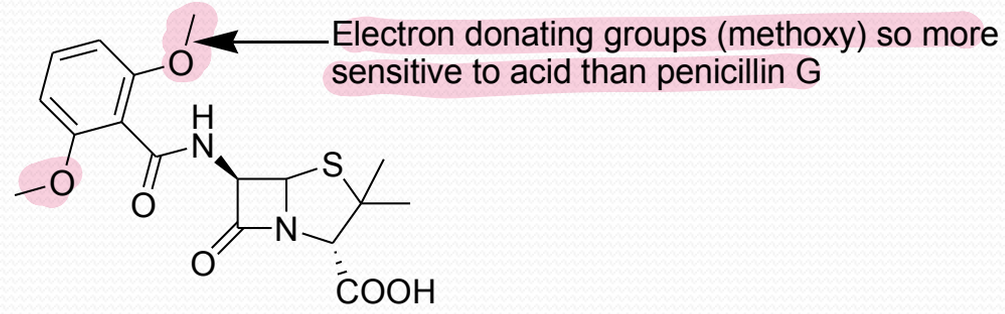
- Increase the steric bulk at the  $\alpha$ -carbon on the acyl-amino group enhances good penicillins activity
- Antibacterial activity is enhanced when the  $\alpha$ -carbon is part of aromatic ring, so based on these points we can conclude that Ortho substituted aromatic ring should produce excellent  $\beta$ -lactamase resistance
- See the following slide for examples.

لكن هون حل المشكلة اعطاهم الثلاثة بين  
 (Amoxicillin + clavulanic acid) cephalosporin (عنهم القدرة انه يكون ذلك) Beta lactamase مع Covalent bond  
 Parenteral oraly Resistance  
 Bulky group (α-position) أو امينع عليه Clavulanic acid أو Sulbactam بعطاهم

# β-lactamase resistant Penicillins

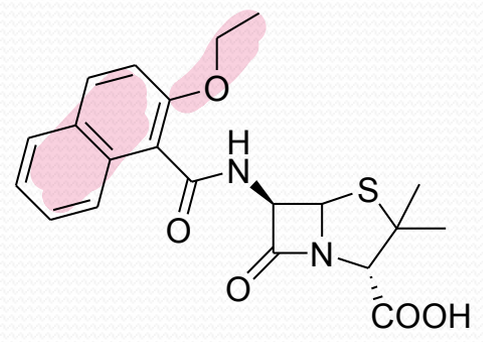
(\*) α Carbon →

اذا عليها (EWG) يكون المركب  
 oraly active من ameno penicillin groups  
 اما اذا عليها (Bulky group) وكان جزء من  
 aromatic Ring يكون ← Beta lactamase Resistance

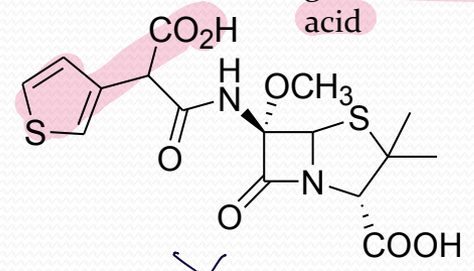


**Methicillin**  
 β-lactamase stable  
 Acid sensitive (why?)

**Ticarcillin** Orally  
 inactive, Broad  
 spectrum and β-  
 lactamase sensitive  
 given with clavulanic  
 acid



**Nafcillin**  
 β-lactamase stable  
 Acid sensitive



~~Temocillin~~  
 ticarcillin

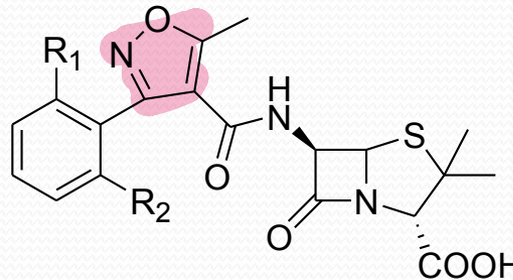
EWG + aromatic

هون ان انا لبت مسكتين و لكن فلتة

narrow spectrum

# Isoxazolyl Penicillins

حلقة فو-ع فيها (O+N)



R<sub>1</sub>, R<sub>2</sub> = H

R<sub>1</sub> = Cl, R<sub>2</sub> = H

R<sub>1</sub> = Cl, R<sub>2</sub> = F

R<sub>1</sub>, R<sub>2</sub> = Cl

Oxacillin orally active

Cloxacillin + Resistance

Flucloxacillin but limited used

Dicloxacillin

## Flucloxacillin: (how to think and analyze)

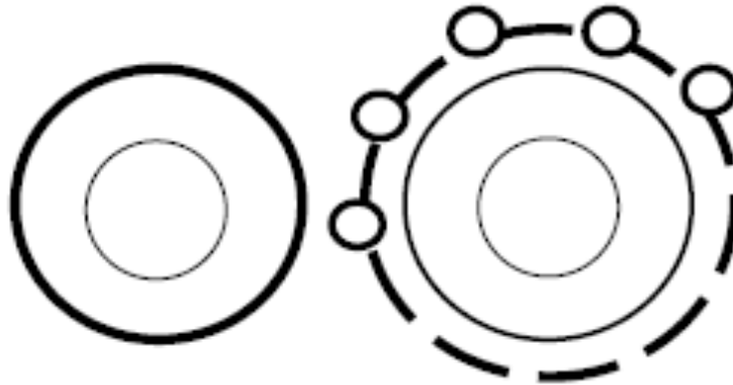
- Bulky which means  $\beta$ -lactamase resistant
- E-withdrawing which means acid stable
- $\alpha$ -carbon is part of an aromatic ring which means good activity

Penicillin dip. (⊗) من فعاليتها اقل

- Bulkier substituents are required for small sized heterocycles to give good anti- $\beta$ -lactamase activity
- Acid stable due to the electron withdrawing effect of the isoxazole ring.
- Used against *S.aureus* resistant bacteria.
- Most penicillinase-resistant penicillins are less active than Penicillin G or Phenoxymethyl penicillin (V) against most non- $\beta$ -lactamase procedures that are normally sensitive to penicillins, increasing  $\alpha$  carbon bulkiness is with price
- Penicillinase-resistant penicillins tend to be bulky and lipophilic with poor penetration into Gram negative bacteria cell envelope

## الفوق Structure بن Gram (-) / (+)

- G+ve
- Thick cell wall
- Cell Membrane



- G-ve
- Thinner cell wall
- Inner Cell Membrane
- Outer membrane with porins

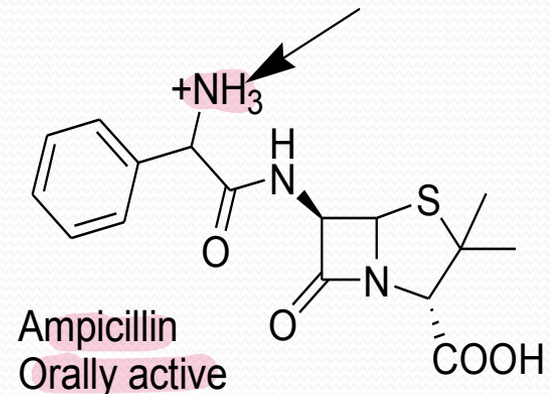
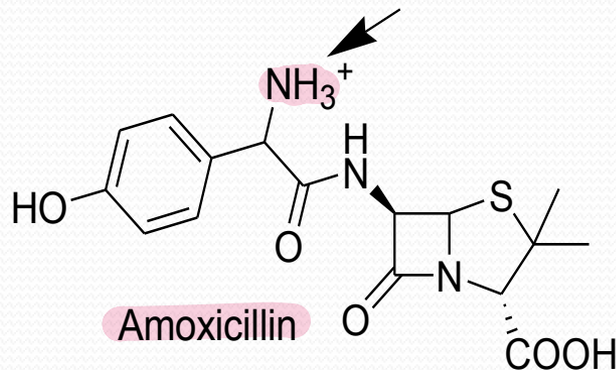


$\alpha$  position at  $\alpha$  carbon ←  $\alpha$  narrow spectrum  $\alpha$  polar or ionized group  $\alpha$   $\alpha$

# Broad Spectrum Penicillins

- Very important finding is the fact that substitution of  $\alpha$ -carbon with polar or ionized group will produce wide spectrum of activity including Gram negative bacteria. It works against (Pseudomonas, E.coli, Haemophilus influenza, ... )  
very resistance gram- bacteria UT

In plasma PH NH<sub>2</sub> will become ionized to NH<sub>3</sub><sup>+</sup>



(Amoxicillin)  
Ampicillin) is orally active ✓  
broad spectrum ✓  
but not  $\beta$ -lactamase resistance X

# Broad Spectrum Penicillins

But is not Beta Lactamase Resistance

Ampicillin and Amoxicillin are effective against Gram negative genera E.Coli, Klebsiella, Haemophila Salmonella, Shiegella and some proteus

Ampicillin and Amoxicillin largely retain Gram positive activity

D-isomer is more active than the L-isomer

The extended activity of  $\alpha$ -amino-benzyl-penicillin is not due to the anti- $\beta$ -lactamase resistant activity rather it is due to the hydrophilic nature of the molecule which enables it to penetrate the outer cell-envelope through the porin channels

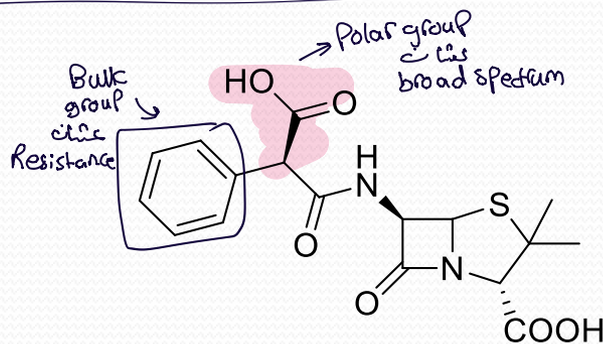
# Broad Spectrum Penicillins

- $\alpha$ -OH also expand the activity but they are less active than the  $\alpha$ -NH<sub>2</sub> derivatives and less acid-stable than the  $\alpha$ -amino group
- $\alpha$ -COOH has wide spectrum activity including all the bacteria that are  $\alpha$ -NH<sub>2</sub> sensitive as well as gram negative bacilli of the genera “Pseudomonas, Klebsiella, Enterobacter and Proteus” however its potency against Ampicillin-Sensitive bacteria is lower than Ampicillin

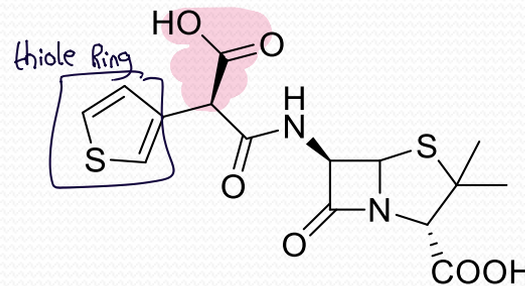
only good group  
effect on  
Pseudomonas

# Carboxypenicillins

- They have a carboxylic acid at the  $\alpha$ -carbon of the acyl side chain.
- They have broad spectrum activity.



Carbenicillin



Ticarcillin

Carbenicillin / Ticarcillin

is a broad / not resistance

but not give orally

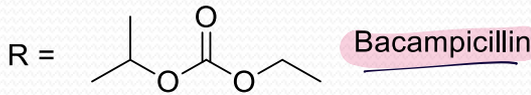
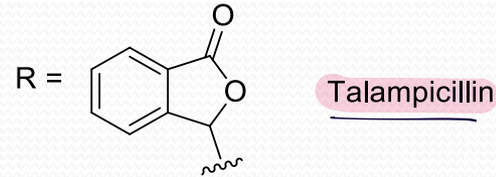
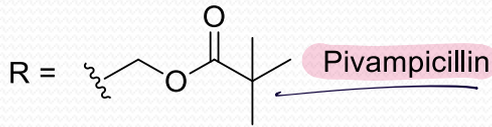
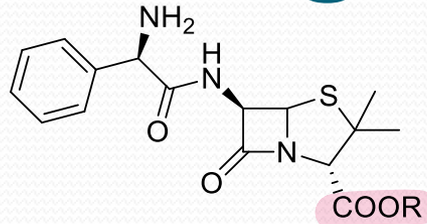
ليس تعطى فمويًا (لا تعطى فمويًا)، (تسكنة الوريد)

# Ampicillin and amoxicillin prodrugs

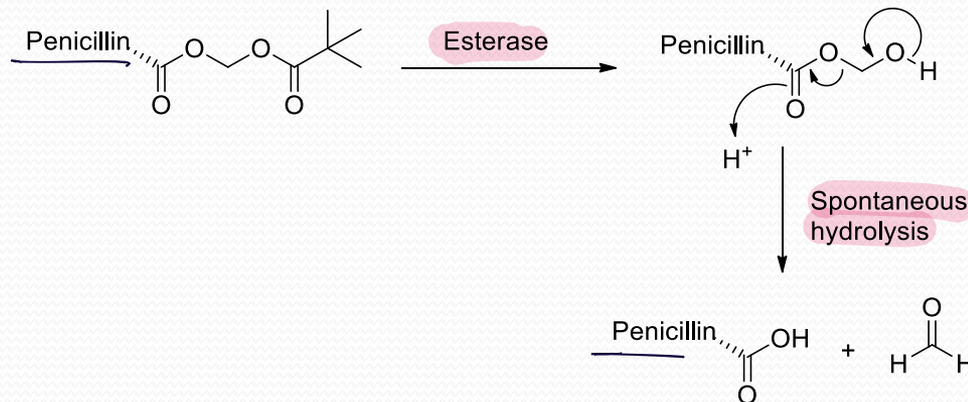
- Ampicillin and amoxicillin are Poorly absorbed through the mucus membrane, this is due to the fact that they formed a zwitter ionic molecule at physiological pH (they contain a carboxylic acid and an amino group in their structure).
  - The oral bioavailability can be improved by masking one of them, mainly the carboxylic acid.... By preparing a prodrug esters.

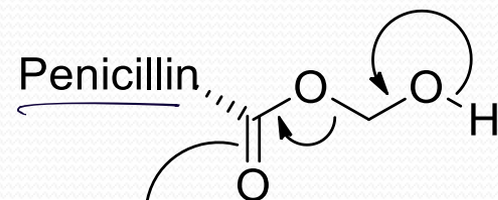
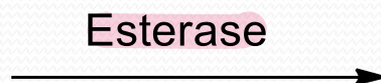
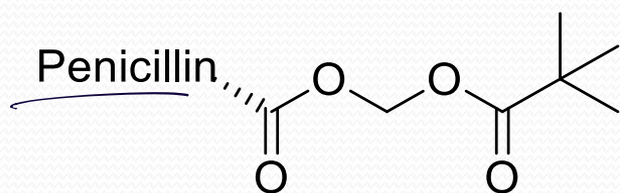
# Ampicillin prodrugs

⊗ الفكرة انو ابيح مركب ← (amino penicillin) Ampicillin او Ampicillin  
 في cool group بحوله ل ester group يعني اظنه اعلى امتصاص  
 ودرنة بروتينات الدم هي (more esterase) بتكسر بواسطة  
 رابطة ال ester وتتحول الى الدواء الاصلي الي هو (cool)  
 (prodrug)



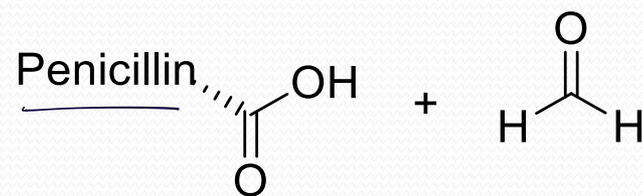
- The methyl ester did not give the same improvement in absorption and activity (why?).





H<sup>+</sup>

Spontaneous  
hydrolysis

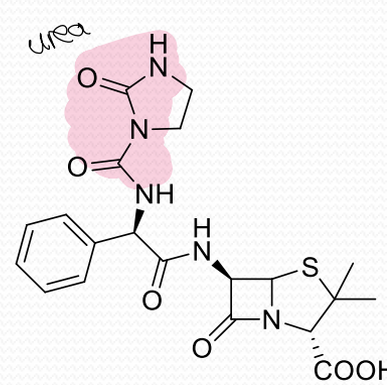


active against Pseudomonas

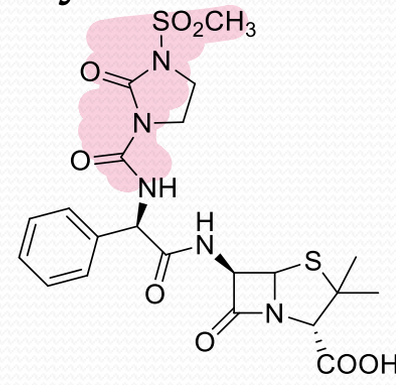
# Ureidopenicillins

- Parenteral (not orally)
- extended spectrum (*P. aeruginosa*)
- Not resistance to Beta lactamase

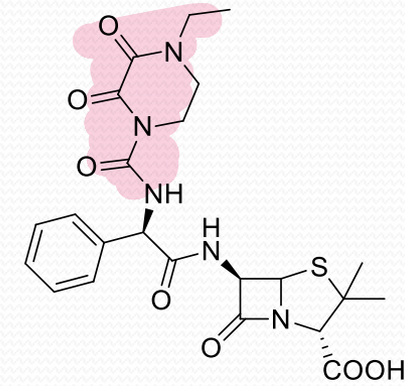
- They all have a urea group at the  $\alpha$ -carbon in the acyl side chain.
- They have better activity compared to amoxicillin and they are more resistant to  $\beta$ -lactamase.  
*هون غلط الجملة وتبجها هي*  
*زائيتها عالية*  
*No resistance*
- Used parenterally for gram -ve infections especially *P. aeruginosa*.
- the ureido group though to mimic some of the peptidoglycan structure, which means that it can bind to penicillin-binding protein



Azlocillin



Mezlocillin



Piperacillin



orally and Resistance *S. aureus*?  
 Oxacillin / cloxacillin / Dicloxacillin

all extended orally  
 Poor oral Absorption  
 بسبب اضافة water soluble group

Amoxicillin → No resistance

TABLE 8.3 Classification and Properties of Penicillins

Penicillin	Source	Acid Resistance	Oral Absorption (%)	Plasma Protein Binding (%)	$\beta$ -Lactamase Resistance ( <i>S. aureus</i> )	Spectrum of Activity	Clinical Use
(Penicillin G)							
Benzylpenicillin	Biosynthetic	Poor	Poor (20)	50-60	No	Intermediate	Multipurpose
Penicillin V	Biosynthetic	Good	Good (60)	55-80	No	Intermediate	Multipurpose
Methicillin	Semisynthetic	Poor	None	30-40	Yes	Narrow	Limited use
Nafcillin	Semisynthetic	Fair	Variable	90	Yes	Narrow	Limited use
Oxacillin	Semisynthetic	Good	Fair (30)	85-94	Yes	Narrow	Limited use
Cloxacillin	Semisynthetic	Good	Good (50)	88-96	Yes	Narrow	Limited use
Dicloxacillin	Semisynthetic	Good	Good (50)	95-98	Yes	Narrow	Limited use
Ampicillin	Semisynthetic	Good	Fair (40)	20-25	No	Broad	Multipurpose
Amoxicillin	Semisynthetic	Good	Good (75)	20-25	No	Broad	Multipurpose
Carbenicillin	Semisynthetic	Poor	None	50-60	No	Extended	Limited use
Ticarcillin	Semisynthetic	Poor	None	45	No	Extended	Limited use
Mezlocillin	Semisynthetic	Poor	Nil	50	No	Extended	Limited use
Piperacillin	Semisynthetic	Poor	Nil	50	No	Extended	Limited use

يستخدمون  
 Resistance  
 tazobactam  
 Resistance

effective against *P. aeruginosa*

\* Penicillin G → not orally + No resistance + Intermediate  
 \* Penicillin V → orally + No Resistance + Intermediate  
 (هون طليت مشكلة acid sensitive لم صفت له EWG) α position رجوع لشكل الهوكس سلايد (37)  
 \* Ampicillin → give orally + No resistance + broad spectrum  
 Amoxicillin (هون طليت مشكلتين - acid sensitive - صفت له EWG at PH 6.2) broad spectrum رجوع لشكل سلايد (37)

\* Methicillin / Nafcillin → not give orally Resistance but narrow spectrum رجوع لشكل سلايد (45)

\* Oxacillin / cloxacillin / flucloxacillin / Dicloxacillin → orally active + Resistance + narrow spectrum رجوع لشكل سلايد (46)

\* Carbenicillin / Ticarcillin → not orally (cool break on stomach) + extended spectrum + Resistance رجوع لشكل سلايد (51)

\* Methicillin / nafcillin / oxacillin / cloxacillin / Dicloxacillin / flucloxacillin →  $\beta$ -lactamase Resistance → narrow spectrum رجوع لشكل سلايد (51)  
 (هون طليت مشكلتين - acid sensitive - صفت له EWG at PH 6.2) رجوع لشكل سلايد (37)