# Antibiotics

Date: 5/11/2024

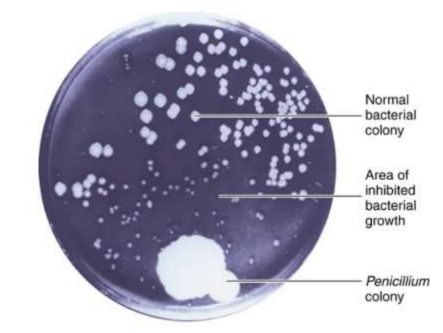
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Bachelor degree in Medicine and Surgery - Mutah university MSC Medical Microbiology – University of Manchester PhD Medical Virology - University of Manchester



# Discovery of Antimicrobial Agents

- In 1928, Alexander Fleming observed that the growth of the bacterium *Staphylococcus aureus* was inhibited in the area surrounding the colony of a mold that had contaminated a Petri plate
- The mold was identified as *Penicillium notatum*, and its active compound, which was isolated a short time later, was named penicillin.

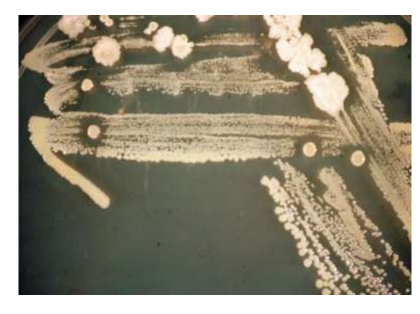


Alexander Fleming took this photograph in 1928. The colony of *Penicillium* mold accidentally contaminated the plate and inhibited nearby bacterial growth.



# Discovery of Antimicrobial Agents (cont)

- Reactions between colonies on solid media are commonly
  observed in microbiology, and the mechanism of inhibition is
  aution
  called antibiosis
  - From this word comes the term **antibiotic**, a substance produced by microorganisms that in small amounts inhibits another microorganism.
- Therefore, the wholly synthetic sulfa drugs, for example, are technically antimicrobial drugs, not antibiotics, a distinction often ignored in practice.

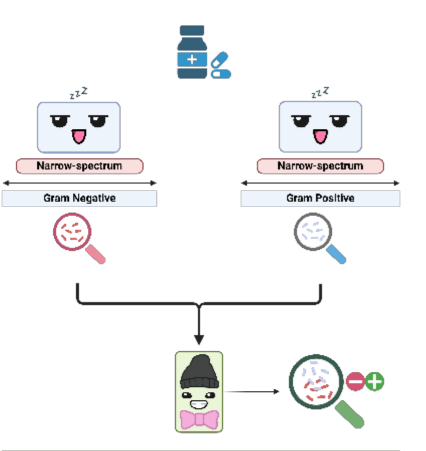




# Spectrum of Antimicrobial Activity

- Narrow-spectrum: the drugs that only act on Gram-positive OR Gram-negative bacteria.
  - Positive: such as Penicillin, and Vancomycin

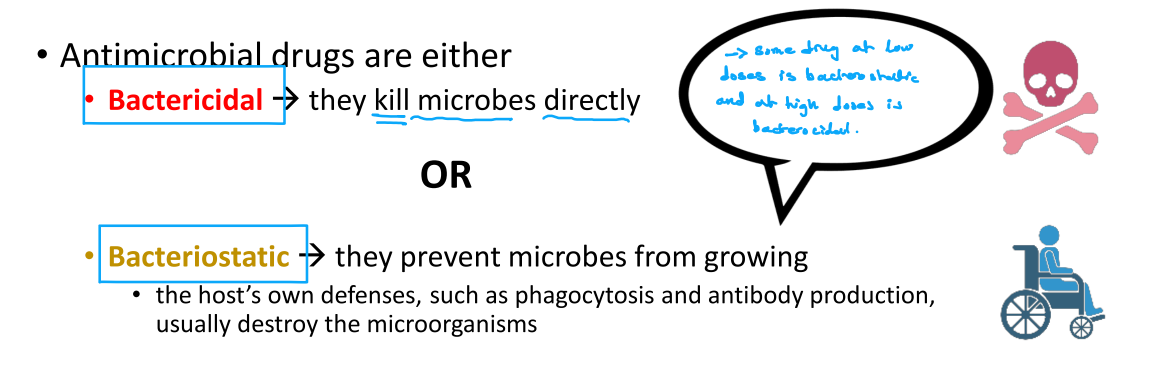
• **Broad-spectrum:** the drugs that have act on Gram-positive AND Gram-negative bacteria.



Broad-spectrum (positive & negative)



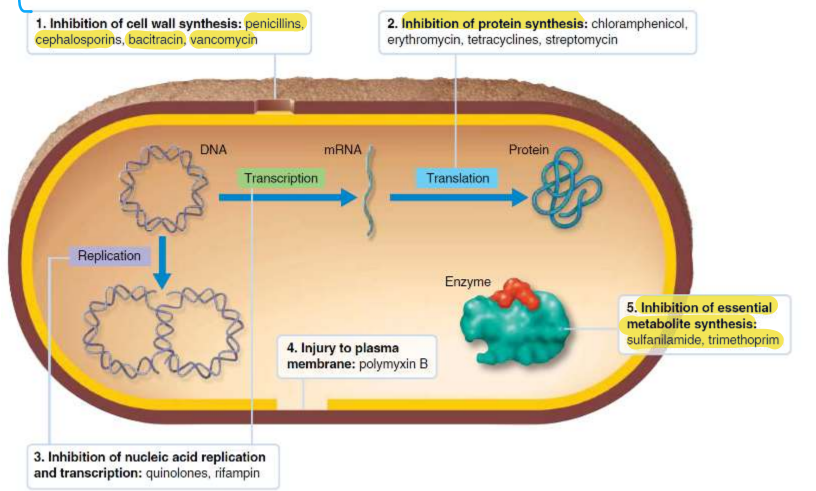
# The Action of Antimicrobial Drugs





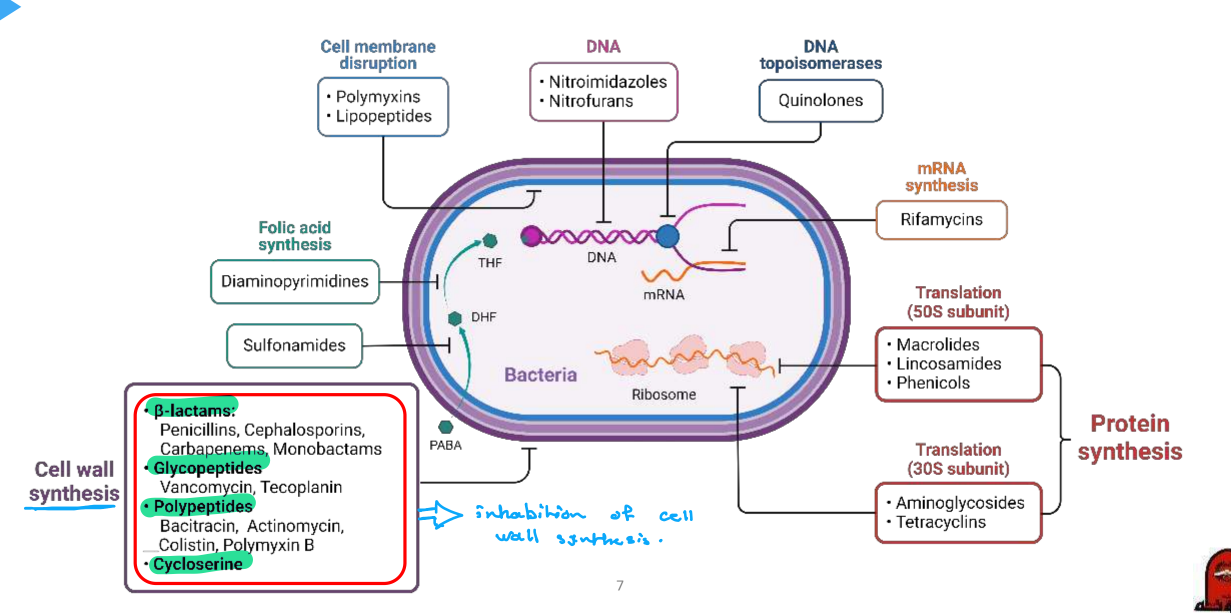
# Antimicrobial Therapy Targets

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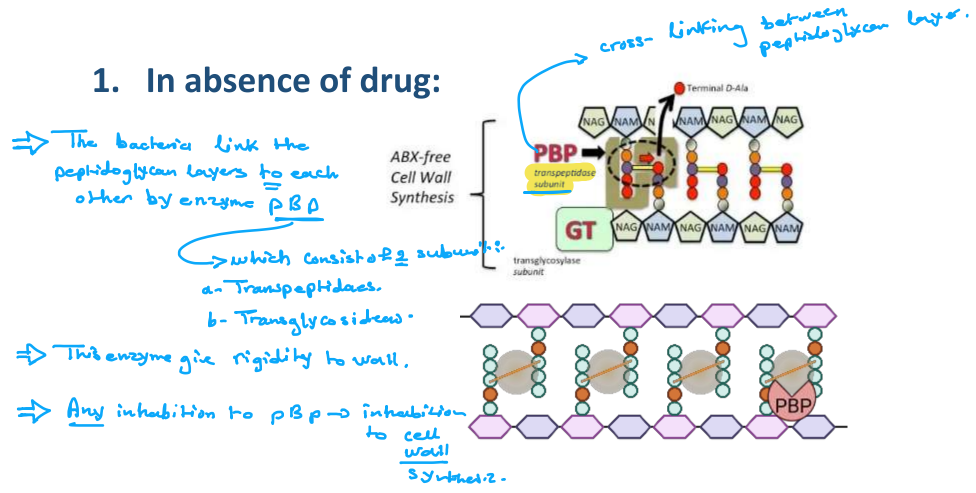




### **Antimicrobial Therapy Targets**



## Inhibition of cell wall synthesis



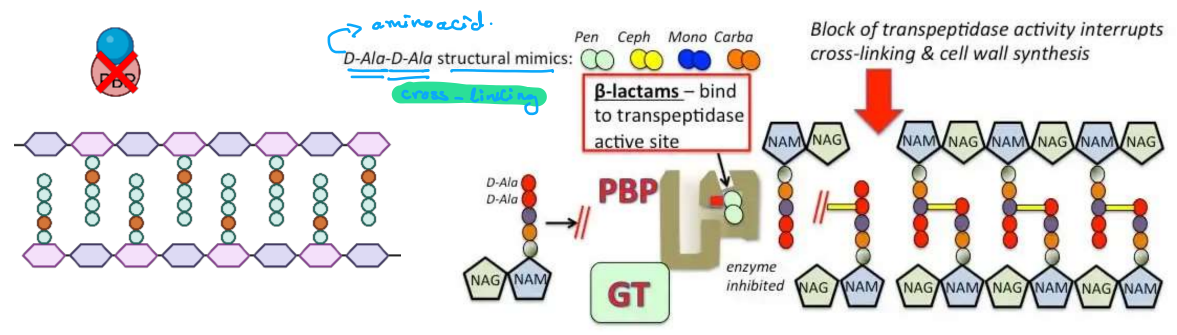


Blackum instead of D-Ada

# Inhibition of cell wall synthesis

structure in peptiogly com.

### 2. In the presence of the drug (beta-lactams):

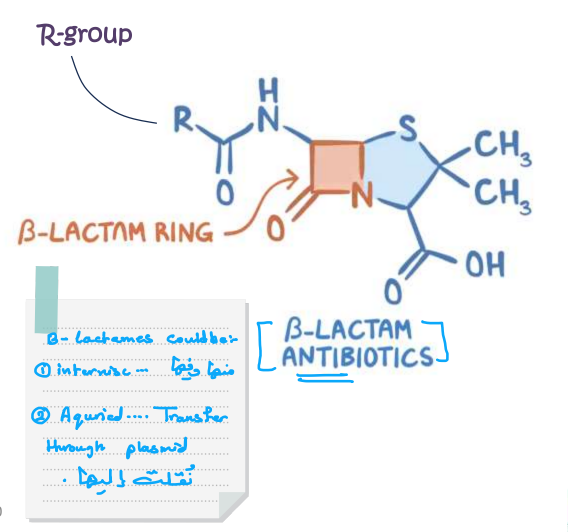


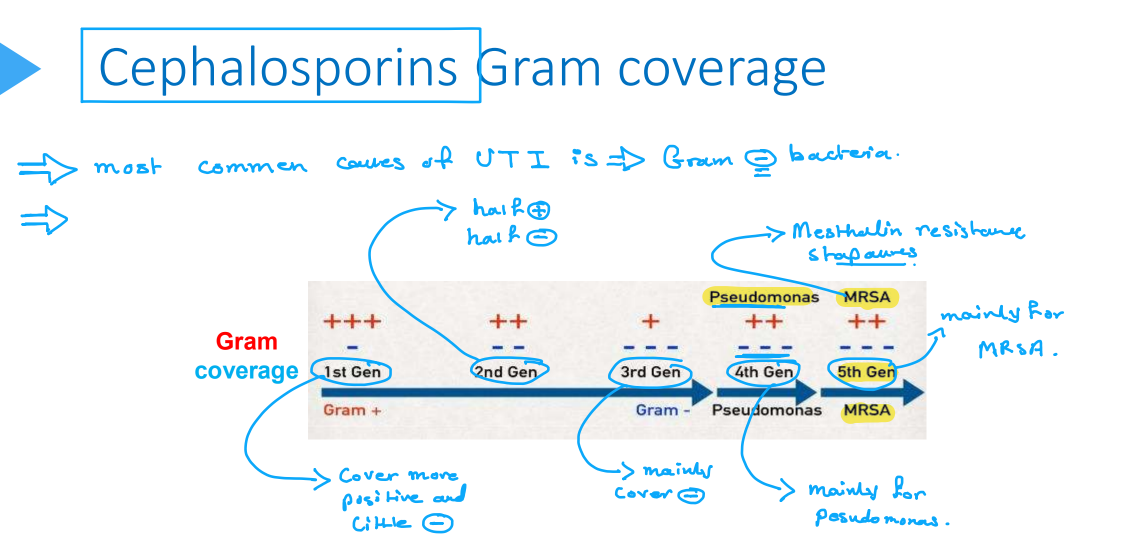
Beta lactam interaction with penicillin binding proteins (PBP) blocks cross linking and compromises cell wall rigidity.



# Beta-lactamase destroys beta-lactam ring ③

- Beta-lactamase Enzymes destroy the Beta-lactam
  Ring
  - R-group → changes the structure of the antibiotic itself
    Clavulanic Acid → inhibits beta-lactamase enzymes produced by resistant bacteria → resembles beta-lactam antibiotics structurally and "tricks" beta-lactamase enzymes into binding with it.
- R group functions in beta-lactam antibiotics:
  - Modifying Spectrum of Activity
  - Enhancing Stability against Beta-lactamase Enzymes → changes the structure of the antibiotic itself → betalactamase enzyme can not bind to the antibiotic



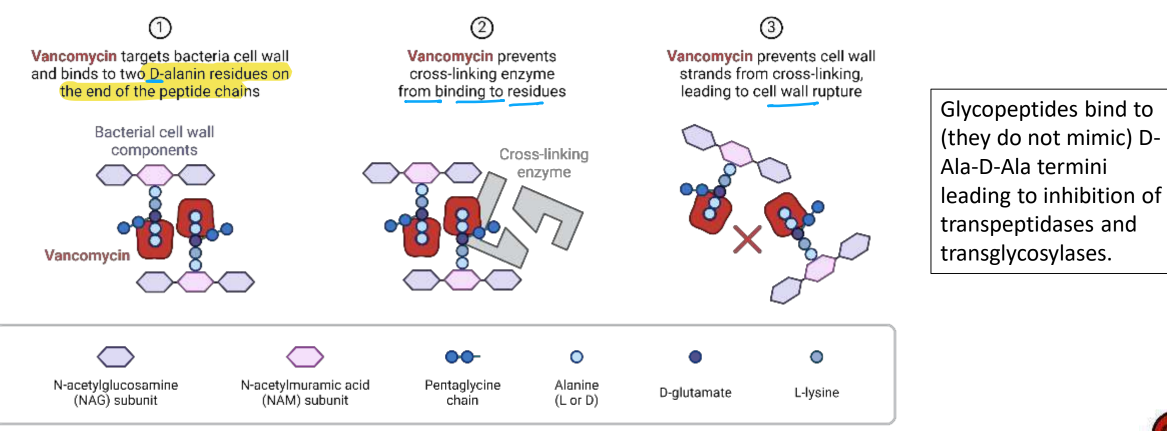




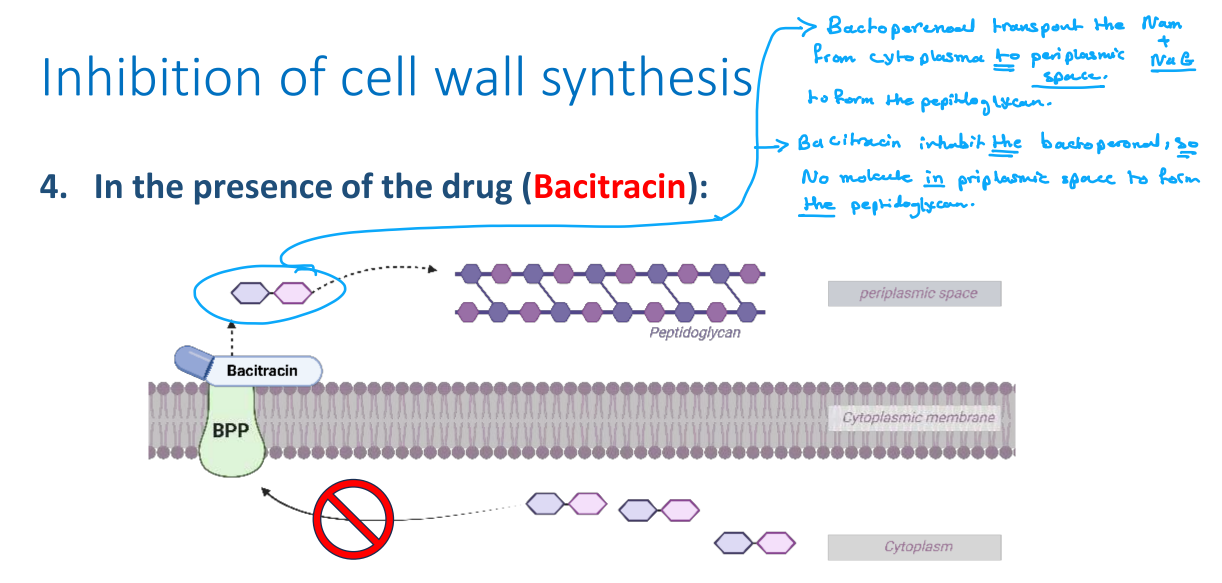
# Inhibition of cell wall synthesis

Vencomy cin bind to D- Alard ) Different way to inhabit cell wall Synthize

### 3. In the presence of the drug (Glycopeptides):



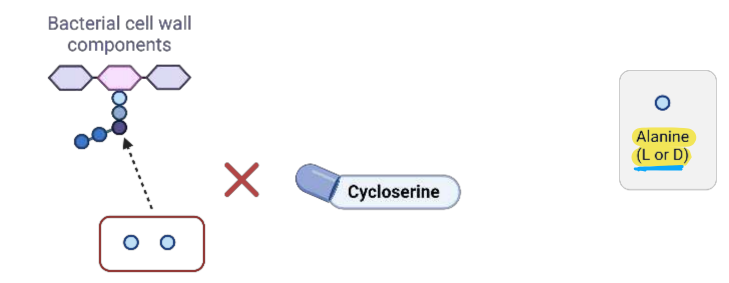




By binding to bactoprenol, Bacitracin prevents the dephosphorylation of this lipid carrier, effectively halting the transport of peptidoglycan precursors (NAG & NAM)

# Inhibition of cell wall synthesis

### 5. In the presence of the drug (Cycloserine:):



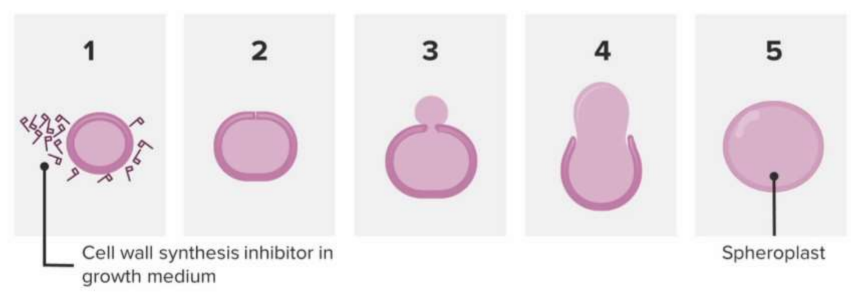
Cycloserine is a structural analog of D-alanine, an amino acid essential for peptidoglycan synthesis. It competitively inhibits two enzymes: alanine racemase and D-alanine ligase.

- Alanine racemase converts L-alanine to D-alanine, a necessary step for the formation of the D-Ala-D-Ala dipeptide.
- By inhibiting D-alanineligase, cycloserine blocks the formation of the D-Ala-D-Ala link in peptidoglycan precursors, stopping cell wall synthesis.

# Inhibition of cell wall synthesis

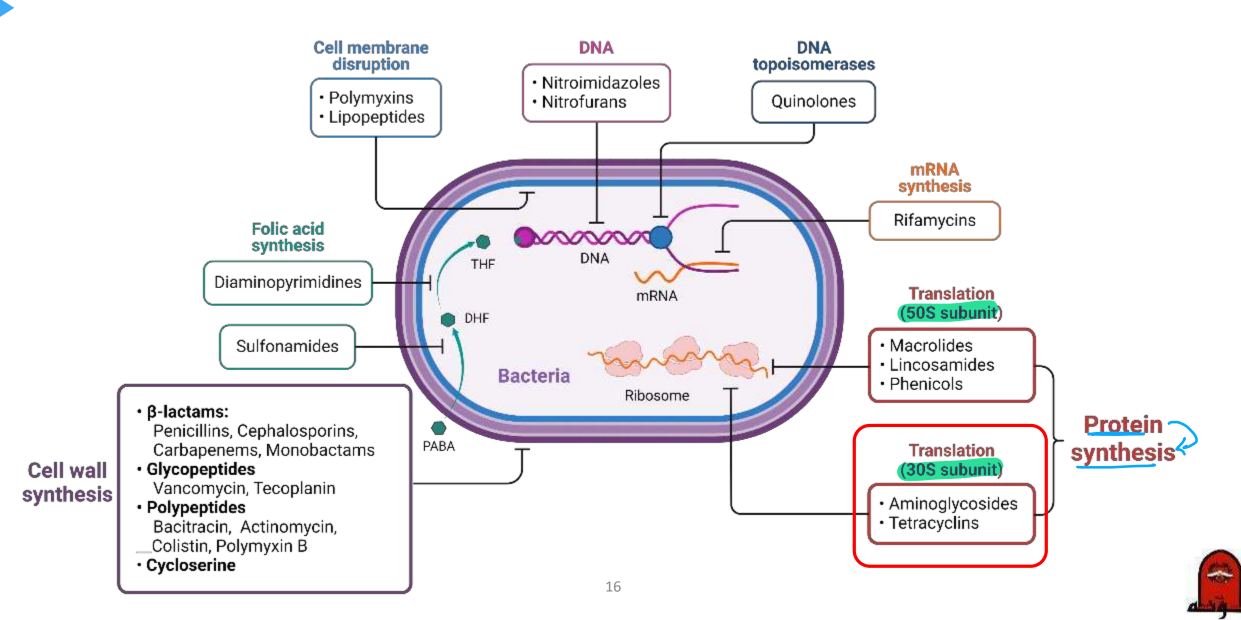
### So, no cell wall synthesis $\rightarrow$ what next? without cell wall.

The dividing cell can not build new cell wall  $\rightarrow$  Spheroplasts will autocatalyze and die





### **Antimicrobial Therapy Targets**



# Inhibition of protein synthesis (30S) الأمثلية (30S)

Antimicrobials that Bind to the 30S Ribosomal Subunit

Aminoglycosides

Streptomycin Kanamycin Gentamicin

Tobramycin

Amikacin

neomycin (topical)

**Tetracyclines** Minocycline doxycycline



Aminoglycosides

Major classes of protein synthesis–inhibiting antibacterials

### Chloramphenicol, macrolides, and lincosamides

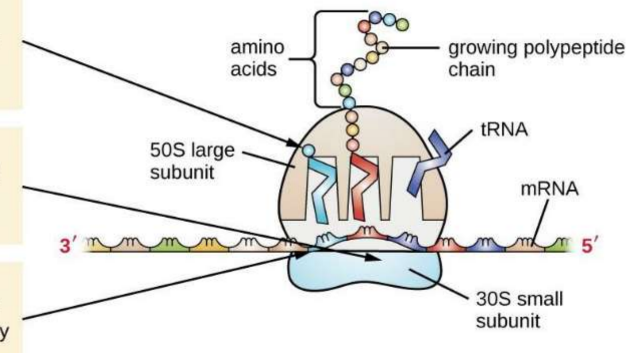
- Bind to the 50S ribosomal subunit
- Prevent peptide bond formation
- Stop protein synthesis

#### Aminoglycosides

- Bind to the 30S ribosomal subunit
- Impair proofreading, resulting in production of faulty proteins

#### Tetracyclines

- Bind to the 30S ribosomal subunit
- Block the binding of tRNAs, thereby inhibiting protein synthesis





# Inhibition of protein synthesis (30S) Aminoglycosides

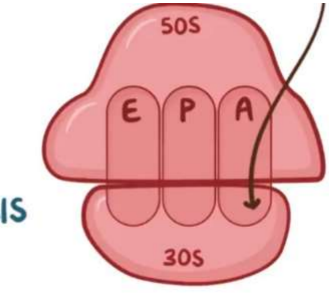
- Irreversibly bind to the 30S subunits
  - Interfere with the proofreading process, thus causing errors in the protein's amino acid sequence
    - These faulty proteins will eventually lead to the death of the bacteria.
  - Prevent the formation of the ribosome-mRNA complex
    - reducing the amount of proteins being synthesized.
  - Inhibiting the initiation of protein synthesis
- Gram positive bacteria have a thicker cell wall compared to Gram negative bacteria, so aminoglycosides can't penetrate them.
- Resistance Common



مشق و اختلق .

# Inhibition of protein synthesis (30S) Tetracyclines

\* BIND to A-SITE on 305 INHIBITS BINDING of tRNAs to mRNA-RIBOSOME COMPLEX SHUTS DOWN PROTEIN SYNTHESIS

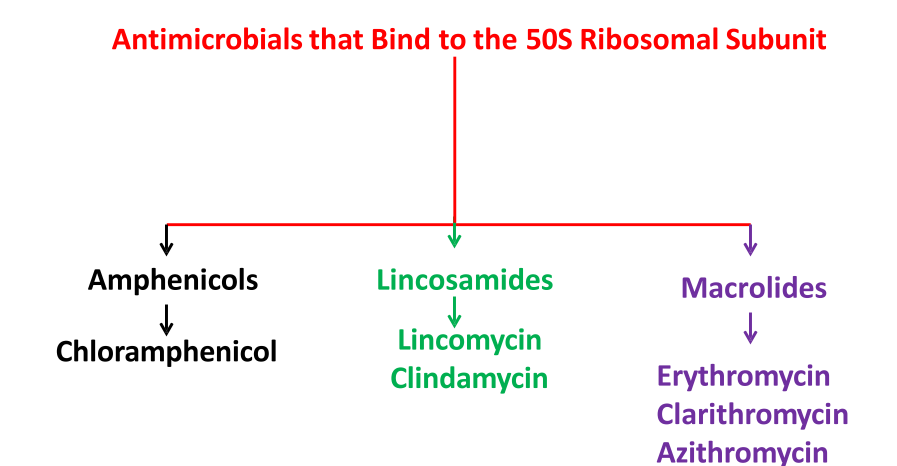


**Mode of action** - The tetracyclines reversibly bind to the 30S ribosome and inhibit binding of aminoacyl-t-RNA

**Spectrum of activity** - Broad spectrum; Useful against intracellular bacteria **Resistance** – Common

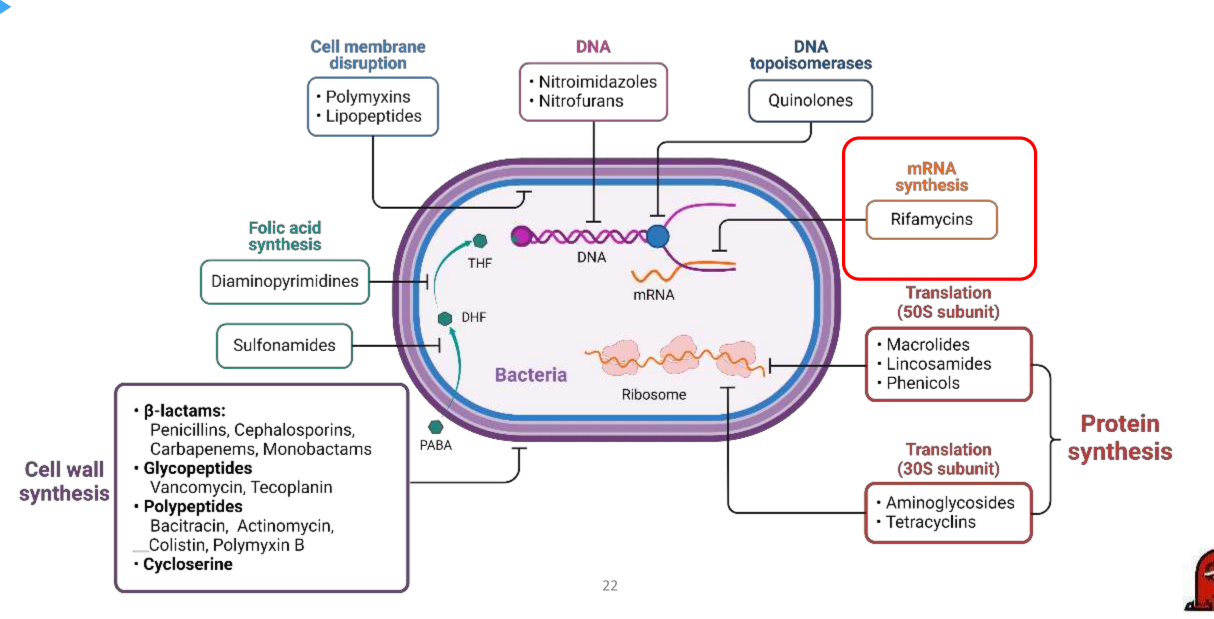


# Inhibition of protein synthesis (50S)





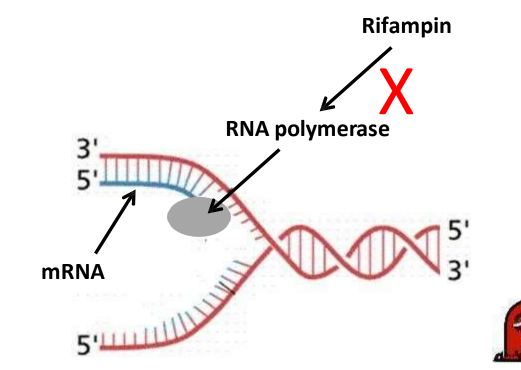
### **Antimicrobial Therapy Targets**



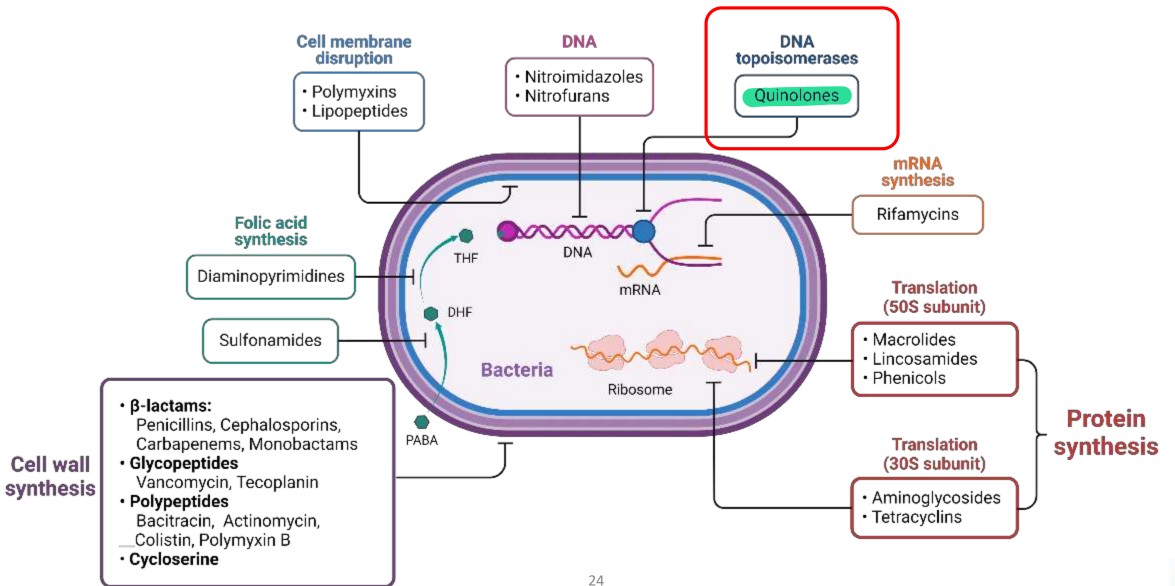
# Inhibitors of mRNA Synthesis

TB **Rifamycins group:** Rifampin, Rifampicin, Rifabutin

- Selectivity due to differences between prokaryotic and eukaryotic RNA polymerase
- Mode of action: these antimicrobials bind to DNA-dependent RNA polymerase and inhibit initiation of mRNA synthesis. No protein.
- Resistance: Common



### **Antimicrobial Therapy Targets**



# Inhibitors of DNA Synthesis

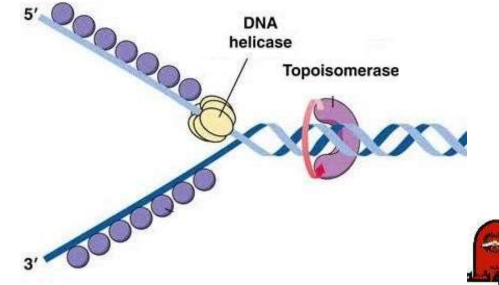
### **Fluoroquinolones:**

nalidixic acid, ciprofloxacin, ofloxacin, norfloxacin, levofloxacin.

**Mode of action** - These antimicrobials bind to the A subunit of DNA gyrase (topoisomerase II) and prevent supercoiling of DNA, thereby inhibiting DNA synthesis.

#### Resistance - Common for nalidixic acid

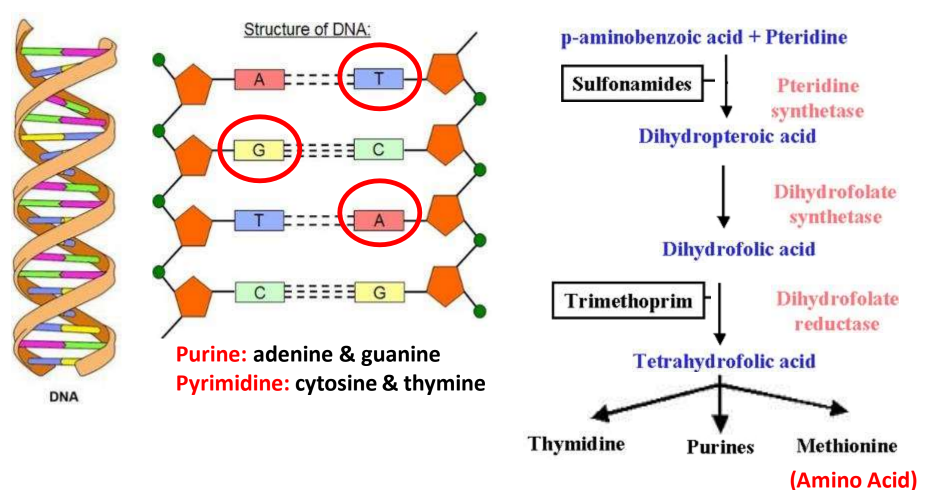
**Supercoiling** refers to the extra twisting or coiling of DNA that compacts it and helps fit the long DNA molecules within the limited space of a bacterial cell. 25



### Inhibition of nucleic acid synthesis

### Nucleic acid synthesis is inhibited by:

- 1. Trimethoprim
- 2. Sulfonamide group: Sulfamethoxazole, Sulfadiazine Sulfathiazole, Sulfamerazine





Interference with cell membrane integrity

- Polymyxin B: binds to membrane of Gram negative bacteria and alters permeability
- This leads to leakage of cellular contents and cell death
- These drugs also bind to eukaryotic cells to some extent, which limits their use to topical applications



# Antimicrobial Drug Resistance Principles and Definitions

- Antimicrobial resistance refers to development of resistance co an antimicrobial agent by a microorganism. It can be of two types:
  - 1. Acquired
  - 2. Intrinsic.

- Resistance provides a selective advantage
- Resistance can result from single or multiple steps



## Intrinsic Resistance

- Intrinsic resistance is the innate ability of a bacterial species to resist the activity of a particular antimicrobial agent through inherent structural or functional characteristics, allowing tolerance of a particular drug or antimicrobial class.
- Such natural resistance can be caused by the following:
  - 1. inability of the drug to enter the bacterial cell (**Impermeability**)
  - 2. innate production of enzymes that inactive the drug (Enzymatic Inactivation).
  - 3. lack of affinity of the drug for the bacterial target

**Example 1:** The intrinsic resistance of gram-negative bacteria to vancomycin is an example of their outer membrane being impermeable to the large, rigid, and hydrophobic glycopeptide molecule vancomycin.

**Example 2:** β-Lactamases hydrolyze β-lactam antibiotics (*Enterobacter species* and *Pseudomonas aeruginosa*)



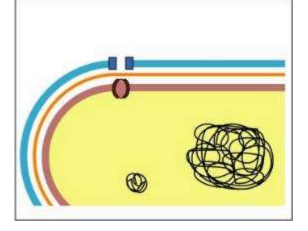
## Acquired Mechanisms of Resistance

- **1. Efflux:** Although efflux plays a major role in intrinsic resistance, changes in the cell wall proteins can also result in novel acquired traits
- **2. Target Site Modification:** Mutations in these PBPs lead to alteration of PBPs, which result in reduced affinity for β-lactam antibiotics
- Enzymatic Inactivation: Examples are enzymes that mediate hydrolysis of the β-lactam ring of β-lactam antibiotics (Staphylococcus aureus have acquired genes that produce beta-lactamase (e.g., via plasmids)



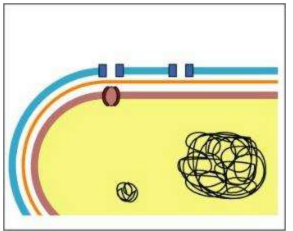
# Principles of Antimicrobial Drug Resistance

- Altered permeability
  - Altered influx
    - Gram negative bacteria

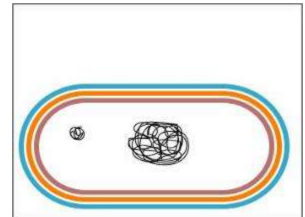


### Altered permeability

- Altered efflux
  - tetracycline



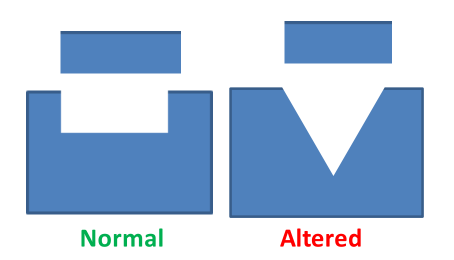
- Inactivation
  - Beta-lactamase

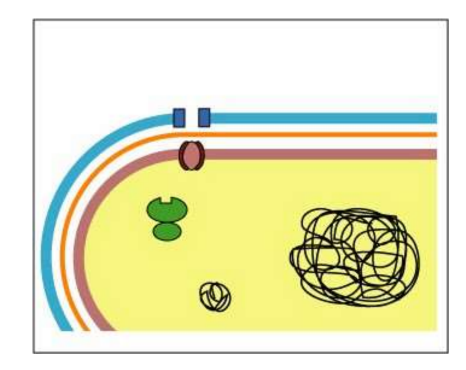




# Antimicrobial Drug Resistance Principles and Definitions

- Altered target site
  - Penicillin binding proteins
  - RNA polymerase
  - 30S ribosome









- Why beta-lactams are not effective against mycoplasma pneumonia?
- Resistance? Which type?

