**LESSON 3.**

**Ultrastructure of bacteria. Acid-fast bacteria and Ziehl-Neelsen stain. Spores and Ojeshko stain. Intracellular additives and Neisser stain**

**LESSON PLAN:**

- The structure of the bacterial cell (structural features of acid-resistant bacteria).

- Sil-Nielsen staining technique of acid-resistant bacteria.

- The role of the Sil-Nielsen method in the diagnosis of tuberculosis.

- Spores, conditions and stages of spore formation.

- Spore staining technique by Ojeshko method.

- Volutin granules and their detection by Neisser's method.

**Acid resistant bacteria**

Due to the weak permeability of the cell wall, they are resistant to decolorization with acid, alcohol and alkalis. This specificity is due to the presence of the following substances in their cell wall:

Lipids

Similar substances

Oxytocin

Mycolic acid, etc.

Mycobacterium tuberculosis (TB)

M.leprae (leprosy)

Some species of Actinomyces

Ziehl-Neelsen technique (AFB Staining)

* Place slide with heat fixed smear on staining tray. Put a filter paper on the slide and add carbol fuchsin, heat the smear until vapour just begins to rise. Do not overheat (boil or dry).   Add additional stain if necessary. Allow the heated stain to remain on the slide for 5 minutes.
* Tilt the slide slightly and filter paper are discarded. Add 5% sulfuric acid or 3% acid alcohol solution for 10-15 seconds until the smear is sufficiently decolorized.
* Tilt the slide slightly and gently rinse with tap water. Add methylene blue for 1 min. Tilt the slide slightly and gently rinse with tap water. Allow the slide to dry with bibulous paper.  
  View the smear using a light-microscope under oil-immersion. Acid fast bacteria is stained red-pink, non-acid fast bacteria is blue

**Bacterial spores**

* Form of conservation of the species in adverse conditions
* Sporulation lasts 20-24 hours
* Metabolic activity is very poor
* The conductivity is very weak, they are resistant to acid, alkaline, alcoholic acid
* It is mainly found in Gram positive bacteria (clostridia and bacilli)
* Highly resistant resting stages formed during adverse environment (depletion of nutrients)
* Formed inside the parent cell, hence called Endospores
* Very resistant to heat, radiation and drying and can remain dormant for hundreds of years.
* Formed by bacteria like Clostridia, Bacillus

**Sporulation**

* Form of conservation of the species in unfavorable conditions
* Sporulation lasts 20-24 hours
* The protoplasm is compacted to the nucleoid. This part is called the spore core.
* Enzyme activity increases.
* Unique enzymes are dipicolinsintetase (5-10%).
* The core contains calcium salt of dipicolinic acid
* The core is surrounded by a peptidoglycan wall is called the prospore
* Between these layers, the peptidoglycan is formed by the cortex.
* A layer of keratin-like protein is formed over the cortex layer
* The outer membrane of the spore contains lipoprotein and a small amount of carbohydrates.

**Types of spores in bacteria**

central – causative agent of anthrax (B.antracis)  
terminal – causative agent of tetanus (C.tetani)  
subterminal – causative agent of botulism (C.botulinum) and gas gangrene (C.perfringens)

Germination

Under favorable conditions (in the human body) spores are converted into vegetative forms. This process is called germination and lasts 3-5 hours. First of all, the cortex is decomposed by the lysosome, and the vegetative form is removed. Then the process of cell growth and division is underway.

Technique of stainin of spores by Ojeshko method

* Place slide with heat fixed smear on staining tray. Add 0,5% salt acid (HCl) heat the smear keep the slide steaming. Do not overheat (1-2 min.)
* Tilt the slide slightly and gently rinse with tap water. Allow the slide to dry with bibulous paper or heat fix the smear.
* After stain by the Ziehl-Neelsen method. The spores are stained red and vegetative forms are stained blue.

**Volutin granules**

* In microorganisms, polyphosphate is a cytoplasmic additive in the form of granules. It was first described in the bacterium Spirillum volutans (hence the name)
* Volutin is an internal reserve of phosphates, and when there is a lack of phosphorus in the environment, the cell divides several times due to it.
* Many bacteria accumulate volutein when some nutrients are deficient. Yeasts, corynebacteria, and mycobacteria usually form appendages in the later stages of growth
* Polyphosphate granules-metachromatic grains (Babes-Ernest bodies) are found in cornebacteria (Corynebacterium diphteria, etc.), a sign of recognition of these bacteria
* It is determined by Neisser method.

**Neisser staining method**

* Place slide with heat fixed smear on staining tray. Add Neisser Glacial acetic acid and methylene blue for 2-3min. Tilt the slide slightly and gently rinse with tap water.
* Add iodine for 30 sec. -1 minute. Tilt the slide slightly and gently rinse with tap water.
* Add Chrysoidin for 5-7 min. Tilt the slide slightly and gently rinse with tap water
* Allow the slide to dry and then view with a 100x bright field objective.
* Because the Volutin granules are alkaline, they turn a dark blue color, taking on the color of acetate.
* Because the cytoplasm is acidic, it takes on the color of vesuvin and turns yellow