Microbiology lec. 8

Bacillus and Clostridium

By: Lect. Shaima'a Al-Salihy



The family Bacillaceae consists of rodshaped Gram-positive bacteria that form endospores. The family includes:

1- aerobic spore forming: Bacillus

2- anaerobic spore-forming: Clostridium

Spore-forming G positive bacilli

Bacillus

Bacillus

- Large G +ve rods arranged in chains.
- aerobic
- saprophytic prevalent in soil, water, air & on vegetation.
- Non motile spore forming (the location of the spore is either central, terminal, or subterminal according to species).



Human infections caused by Bacillus sp.

Bacteria		Diseases		
Bacillus anthracis		Anthrax (cutaneous, gastrointestunal, and inhalational) and anthrax meningitis		
Bacillus cereus		Gastroenteritis, intravenous catheter septicemia, and endocarditis		
Bacillus Iechniformis		Gastroenteritis		
Other sp.	Bacillus	Opportunistic infections		

Bacillus anthracis

The causative agent of **anthrax**, which is a primarily a disease of animals.

□ Morphology:

- Large G+ve rods, arranged in long chain, with square end (bamboostick) appearance.
- Capsulated (polypeptide)
- Non motile
- Non hemolytic.
- Spore-forming (central spore, the same width of the cell).

Bacillus anthracis



□ Cultural characteristics:

- On nutrient agar: medusa head.
- On blood agar: non hemolytic, ground glass appearance, and some times have tail.
- On penicillin- containing media: strings of pearl phenomenon.
- On gelatin medium: inverted fir tree appearance
- □ Antigenic structure:
- Capsular antigen
- Cell wall antigen
- Somatic antigen

Bacillus anthracis





□ Virulence factors:

- Capsule: poly- D- glutamic acid capsule, antiphagocytic.
- Anthrax toxin- complex: composed of three components:
 - **Protective antigen (PA):** is the binding domain of anthrax toxin binds to specific cell receptors forming a membrane channel that mediate entry of EF & LF into the cell.
 - Edema factor (EF): causes cellular edema within the target tissue and inhibit neutrophil function.
 - Lethal toxin (LT): the major virulence factor and causes death in infected animals.

Virulence factors of *B. anthracis*



Pathogenicity



□ Pathogenesis of B. anthracis

How the Bacterial Toxin "Lethal Factor" Results in the Fatal Spread of Anthrax



Source: Dixon et al., Anthrax. New England Journal of Medicine 341:815-826, 1999.

TOTAL VIDEO CONVERTER HTTP://EFFECTMRTRIX.COM

Clinical findings:

- In human, 95% of cases are cutaneous anthrax & 5% are inhalation anthrax.

Cutaneous anthrax: generally occur on arms or hands & less frequently on face & neck.

Stages of cutaneous anthrax:

- **Papule** at the site of entry (wound or scratches).
- Vesicle
- Eschar: malignant pustule
- Gastrointestinal anthrax:
- Ulcers
- Bloody diarrhea, due to necrosis and ulceration which produces GI hemorrhage.
- Renal failure due to anthrax toxin.
- Death in untreated cases (MR more than 50%).
- Oropharyngeal anthrax



Inhalation anthrax: wool sorter's disease

- Non specific symptoms: after 1-6 days incubation period low grade fever and non productive cough.

- Second phase: The early clinical manifestations of inhalation anthrax is marked after 1-2 days hemorrhagic necrosis & edema of the mediastinum & substernal pain, high fever, shortness of breath, tachypnea, and hematemesis.

Sepsis may occur. Spread to the meninges causing hemorrhagic meningitis. The fatality rate is 85-90%.

□ Immunity: specific antibodies to anthrax toxin (primarily against PA), and capsular antigen.



Epidemiology: soil is contaminated with anthrax spores from the carcasses of infected animals. These spores remain viable in soil for decades.

□ Prevention and control:

- Chemoprophylaxis with antibiotics: is indicated for people who have been exposed to anthrax but do not have symptoms of the disease (ciprofloxacin and tetracyclines).
- **Immuno-prophylaxis**: A noncellular human anthrax vaccine called anthrax vaccine adsorbed (AVA).
- Decontamination of animal products: Chemical disinfection, Burn carcasses.

HOW ANTHRAX ATTACKS

Anthrax is a naturally occuring bacterium that plagues farm animals and, occasionally, agricultural workers. An airborne form of the disease, however, can be harnessed as a potent biological weapon.



Source: "The World's Best Anatomical Charts"; "Zoology"; Anthrax Vaccine Immunization Program; Journal of the American Medical Association

B. cereus

- Causes food poisoning which has two types:
- **emetic type:** associated with fried rice, is manifested by nausea, vomiting, abdominal crumps, it is self-limited, recovery occur within 24 hrs.
- diarrheal type: associated with meat & sauces, (1-24 hrs incubation period) manifested by profuse diarrhea with abdominal crumps & pain
- eye infection

Spore-forming G positive bacilli

B. cereus



Differentiating features between Bacillus anthracis and Bacillus cereus

Characteristics	Bacillus anthracis	Bacillus cereus
Motility	Nonmotile	Motile
Capsule	Capsulated	uncapsulated
Medusa head colony	Present	Absent
Hemolysis on sheep blood agar	Absent	Present
Gelatin liquefaction	slow	rapid

Clostridium

Clostridium:

□ Morphology & identification:

- Large G+ve rods.
- Strict anaerobes
- Spore forming: spores wider than the diameter of the rod. They may be centrally, subterminally or terminally located.
- Most species are saprophyte, a few are commensals residing in the intestine of human and other animals
- Most *Clostridium* species with few exception are motile due to the presence of peritrichous flagella.

Anaerobic spore- forming Clostridia



Cl. tetani The causative agent of tetanus (lock jaw)

- Morphology:
- G+ve, straight rods with rounded ends.
- Has round, terminal, and bulging spores giving drumstick appearance to the bacillus.
- Most strains are motile
- Capsulated.
- Cultural characteristics:
- Robertson cooked meat: turbidity, gas production and blackening meat.
- Blood agar: a- hemolytic $\longrightarrow \beta$ hemolytic, Surface colonies tend to swarm over the entire surface of the agar
- □ Virulence factors:
- C. tetani produces tetanolysin, tetanospasmin, and neurotoxin (nonspasmogenic toxin). **Tetanospasmin** is the toxin responsible for the clinical manifestations of tetanus.



Cl. tetani

Virulence factors of Clostridium tetani

Virulence factors	Biological functions
Tetanospasmin	Potent heat labile toxin; prevents the release of neurotransmitters (e.g., GABA, glycine, etc.), hence blocks specific synaptic inhibition in the spinal cord. Motor neurons are left under no inhibitory control and undergo sustained excitatory discharge
Tetanolysin	Heat-stable hemolysin; unknown significance in pathogenesis of tetanus. Oxygen-labile hemolysin.
Neurotoxin	Nonspasmogenic and peripherally active neurotoxin of unknown significance

Pathogenesis:

- C. tetani is a noninvasive bacillus and causes disease only by production of toxins.
- The infection remains localized in the area of devitalized tissues (wounds burns, umbilical stump, surgical suture) into which the spores have been introduced.
- Germination of spores & vegetative cells produce toxin. Toxin production are aided by (Necrotic tissues, Calcium salt, Associated pyogenic infections).
- The toxin reaches the CNS & binds to receptors in the spinal cord & brain. This binding is irreversible.
- Unregulated spread of impulses, inhibited anywhere in the central nervous system (CNS) due to the action of toxin
 Tetanus toxoid is antigenic but nontoxic. Tetanus toxin is made into toxoid by treating it with formaldehyde.

Clinical findings:

- Incubation period: 5 days to many weeks.
- Rigidity & spasm of voluntary muscles of the jaw (Lockjaw, trismus, Risus sardonicus).
- Gradually, other voluntary muscles are involved result in tonic spasm (Opisthotonos).
- Death occurs due to interference with respiration.
- The mortality rate in generalized tetanus is high.

Tetanus neonatorum is an enormously important medical problem in developing countries. The cause is sepsis of umbilical stump.

Cl. tetani





□Host immunity:

Specific antibodies produced against tetanus toxin are protective. Antibodies specifically combine with free toxin and prevent the action of the toxin.

Cl. tetani

□Prevention: depends on:

- Active immunization with toxoids (during the first year of life, tetanus toxoid is often combined with diphtheria toxoid and acellular pertussis vaccine (DPT).
- Proper care of wounds contaminated with soil
- **Passive immunization:** prophylactic use of antitoxin (It neutralizes toxin that has not been fixed to nervous tissues).
- Administration of penicillin.



Elimination of maternal & neonatal tetanus

28 Countries eliminated MNT between 2000 & October 2012

*(Plus 15 States out of 35 in India, Ethiopia part and 29 provinces out of 33 in Indonesia) leaving 31 countries yet to eliminate MNT



Data Source: WHO/UNICEF database, October 2012. 194 WHO Member States. Map production: Immunization Vaccines and Biologicals, (WB), World Health Organization Date of Slide: 30 October 2012

- Cl. Botulinum: The causative agent of botulism.
- Morphology:
- Large G+ve rods, possesses subterminal and oval bulging spores.
- The bacillus is motile by the presence of peritrichous flagella,
- The bacteria are noncapsulated.
- Cultural characteristics:
- Grow on different types of culture media under an aerobic conditions. They produce spores when grown in alkaline glucose gelatin media at 20-25°C.
- Produce lipase which form iridescent film on C. botulinum colonies grown on egg yolk agar.
- □ Virulence factor:
- Botulinum toxin: autolysis It consists of two subunits A and B.
- There are many types of botulinum toxin form (A-G).



Pathogenesis:

- Botulism is an intoxication resulting from the ingestion of food in which *C* botulinum has grown and produced toxin.
- Spores of C botulinum germinate in infected canned food under anaerobic conditions, vegetative forms grow and produce toxin.
- The toxin acts by blocking release of acetylcholine at synapses and neuromuscular junctions resulting in lack of muscle contraction and flaccid paralysis.

Infant botulism:

- "floppy baby"
- One of the causes of sudden infant death syndrome.
- Honey is implicated as a vehicle of the spores.

Cl. botulinum



□ Clinical findings:

- Symptoms begin 18-24 hours after ingestion of the toxic food
- visual disturbances and blurred vision.
- inability to swallow, and speech difficulty.
- signs of bulbar paralysis are progressive, and death occurs to respiratory paralysis or cardiac arrest. The mortality rate is high. Recovered patients do not develop serum antitoxin.

□ Treatment:

- (a) Respiratory supportive therapy.
- (b) neutralizing unbound toxin by specific antitoxins.

(c) stopping toxin production by use of antibiotics (Metronidazole is the current antimicrobial drug of choice with penicillin).

Cl. perfringens: Causes gas gangrene, myonecrosis, and food poisoning.

□ Morphology:

- Large G+ve rods, occur as single, in chains, or in bundles.
- Capsulated
- Non-motile, however, the bacteria multiply rapidly, giving a characteristic spreading colony appearance on the media, resembling the growth of motile clostridia.
- They possess central or sub-terminal spores (rarely seen).
- Cultural characteristics:
- Robertson cooked meat: turn the meat pink then black.
- **Blood agar**: produce double zone of hemolysis; a narrow zone of β hemolytic due to theta-toxin and a much wider zone of incomplete a- hemolysis due to alpha-toxin of the bacteria.







□ Virulence factors:

- * Toxins:
- Alpha toxin (lecithinase): breaks down Lecithin. It is responsible for toxemia and increases vascular permeability of blood vessels, thereby causing massive hemolysis and bleeding, tissue destruction, and myocardial dysfunction.
- Theta toxin has hemolytic, cytolytic & necrotizing effect.
- Enterotoxin: heat-labile toxin can induce diarrhea within 6-18 hrs.
- DNase, hyaluronidase, collagenase
- * Enzymes:
- Neuraminidase.
- fibrinolysin, histamine,
- The "bursting factor" responsible for typical muscle lesions observed in gas gangrene.
- The "circulating factor" increases adrenaline sensitivity of the capillary membrane and also inhibits phagocytosis.

□ Pathogenesis:

- Gas gangrene (clostridial myonecrosis) is a mixed infection (Toxogenic & proteolytic clostridia with various cocci & G negative rods).
- The spores reach tissues either through contamination of area with soil or feces or from the intestinal tract.
- The spores germinate, the vegetative cells multiply & ferment carbohydrates in tissues producing gas.
- Spread of infection due to secretion of necrotizing toxin & hyaluronidase.
- Extension of tissue necrosis increased the bacterial growth result in hemolytic anemia, severe toxemia & death.

□ Clinical findings:

From contaminated wounds (Fractures, postpartum uterus) the infection spread in 1-3 days to produce:

- crepitation in the subcutaneous tissue & muscles.
- foul-smelling discharge.
- Necrosis, Fever, toxemia, shock & death.

□ Treatment:

- Removal of damaged tissue.
- Metronidazole is the antibiotic of choice. Prophylactic use of the antibiotic in association with surgery is effective. Antibiotic prophylaxis using broad-spectrum antibiotics, such as gentamicin, amoxicillin, and metronidazole, is effective, since occurrence of mixed infections with aerobic and anaerobic bacteria is frequent. Antibiotic therapy is not recommended for the treatment of C. perfringens food poisoning.



Cl. difficile: pseudomembranous colitis

- Administration of antibiotics result in proliferation of drug-resistant *Cl. difficle* causing watery or bloody diarrhea, sometimes with abdominal crumps, leukocytosis & fever.
- The most common antibiotics that associated with PMC are ampicillin & clindamycin.

Antibiotic associated diarrhea:

The administration of antibiotics frequently leads to a mild to moderate form of diarrhea termed AAD. The condition is less severe than PMC. About 25% of AAD is caused by *Cl. difficile*.

causes

Spore-forming G positive bacilli Clostridia *Cl. difficile*



