### Enterococcus (Enterococcus faecalis, Enterococcus faecium etc.)

Microscopy: G+ cocci in pairs or short chains, catalase negative

<u>Cultivation:</u> small greyish white colonies on blood agar with viridation Some of them have yellow pigment, some are mobil selective diagnostic Slanetz-Bartley (sodium azide) agar - pink to red colonies Bile-aesculin agar: black colonies

**Biochemistry:** pyrrolidonylarylamidase (PYR-positive) and leucinaminopeptidase (LAP-positive) high resistance, growing in 6,5% NaCl agar, large temperature interval

**Pathogenicity:** part of normal digestive tract flora, more frequent in long therm hospitalised patients with medical devices or patients treated with broad spectrum antibiotics

Urogenital infections, wound infections, intraabdominal infections, endocarditis – more often in drug users or seniors, catether sepsis, biliary tract infection



#### Factors of virulence:

*gelatinase, feromon* substance, *colonization factors, bacteriocins* - inhibition of other bacteria *VanA, B, C* gens causes rezistence to vancomycin (*C* is gen of primary resistence, *VanA/B* of secondary resistance, transferable through plasmides)

Treatment: primary resistant to cefalosporines

<u>Liht urinary tract infection</u>: ampicillin, ampicillin with  $\beta$ -lactamase inhibitors, nitrofurantoin, possible glycopeptides.

<u>Wound infections, sepsis and endocarditis</u>: combination of aminoglykoside + penicillin/ampicillin or glycopeptides (vancomycin, teicoplanin)

VRE (vancomycin resistant enterococci) - linezolid, quinupristin/dalfopristin

#### Laboratory dg.:

microscopy, cultivation on BA, on Slanetz-Bartley medium Latex agglutination – differentiation from streptococci, from other bacteria through PYR test and LAP Phenotypic test (production of yellow pigment, moovement) Biochemistry: fermentation of arabinosis and pyruvate:

#### E. faecium

arabinosis fermentation – change of the indicators colour pyruvate negative resistent to ampicilin EN-coccus test E. faecalis

without fermentation pyruvate fermentation susceptible to ampicillin

### G+ rods

#### Listeria monocytogenes



**Morphology:** microscopy: G+ rods, catalase positive

<u>Cultivation</u>: chromogennous media, growth in cold, on BA form grey colonies with haemolysis – looks like enterococci, streptococci or differoids

**Pathogenicity:** wound infection, new-born babies infection (meningitis or sepsis)

**Virulence factors**: lysteriolysin, internalins (intracellular alive)

Treatment: fluoroquinolons

Laboratory dg.: microscopy, cultivation on chr. medium/ BA and bile-aesculin medium, catalase detection, BBL test

# Corynebacterium difteriae



**Microscopy:** G+ rods with metachromatic granules, club-shaped looking like chineese signs, catalasa positive

Cultivation: does not grow on MH, but on BA, on telur media (Clauberg)

**Pathogenicity:** strains producing toxin (microb attacked by fag) causes diphteria with pablanes (couldn't take off without bleeding), man suffocate, arise of myocarditis etc. Non-toxic strains causes skin inflammations.

#### Factors of virulence: diphteric toxin

Therapy: vaccination, antidiphteric globulin (deserters!), PNC, tracheostomy, cortikoids

**Laboratory dg.:** microscopy, staining of specific parts - granules (Lebranc), Clauberg medium - metal shiny colonies with blue zone around colonies, Lofler medium, detection of toxins through Elek test, PCR, demonstration on guinea-pig.

### Other Corynebacteria (C. jejkeium etc.)

Microscopy: G+ rods with metachromat. granules, club-shaped form looks like chineese signs, arranged in palisades, catalase positive **Cultivation:** any growth on MH, but BA **Pathogenicity:** wound infection, sepsis, urinary tract infections **Factors of virulence**: haemolysins **Treatment:** vancomycin, teicoplanin, rifampicin, if posssible - PNC Laboratory dg.: microscopy, cultivation on BA, biochemistry, BBL test...

### Rod Bacillus

### B. antracis

Microscopy: G+rods looks like bamboo stick, spors (central terminated) – only in air <u>Cultivation</u>: on BA – large, flat, spreading through the agar surface - caput medusae, ahaemolytical <u>Pathogenicity and pathogenesis</u>: contact with ill person, dead animals or their productes (skin), spors invade into organism, germinate and produce toxin. Via entrance is disease devided into 3 forms.

- 1. skin pustula maligna
- 2. pulminal after inhalation arises hemoragic necrosis of nodes with mediastinitis ends as septic shock
- 3. intestinal via contaminated food causes bloody diarrhea, high temperature etc.
- !! spors are easy to diffuse, that's why it is discussed as a biological warfare!!

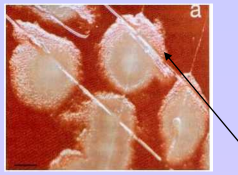
Virulence: toxin (3components)

<u>Therapy:</u> PNC, ciprofloxacin, doxycyklin, chloramphenicol <u>Prevention:</u> veterinary control of animal, vaccination of animal or people <u>Laboratory diagnosis:</u> microscopy, cultivation on BA Antigen detection - Ascoli termoprecipitation reaction, animal demonstration

!! Can do only laboratory with biosafety level III.







## B. cereus



**Microscopy:** G+rods, central terminated spores

<u>Cultivation</u>: on BA flat colonies with  $\beta$  haemolysis, PEMBA-blue colonies

<u>**Pathogenicity</u>**: component of gastrointesinal flora, contamination of food, causing diarrhea, vomitting. Diarrhea is caused by thermolabil enterotoxin (source: sauce), vomitting is caused by thermostabil toxin (source: rice). Also causes eye + wound infection</u>

#### Factors of virulence: enterotoxins

**Treatment:** rehydratation + linkosamids. Prevention: good food preparation Eye infection: linkosamids + aminoglykosides

**Laboratory dg.:** microscopy, cultivation on BA/PEMBA, detection of granules toxin detection via ELISA method or latex agglutination