

NS1 Module 2023-2024 Dr. Mohammad Odaibat

Cerebrospinal Fluid (CSF)

- Composition and formation
 - CSF is the 3rd major fluid of the body
 - Adult volume 90-150 mL
 - Neonate volume 10-60 mL
- CSF functions
 - Supplies nutrients to nervous tissues
 - Removes metabolic wastes
 - Protects / cushions against trauma

Cerebrospinal Fluid (CSF)

Four major categories of disease

- Meningeal infections
- Subarachnoid hemorrhage
- CNS malignancy
- Demyelinating disease

Indications for analysis

- To confirm diagnosis of meningitis
- Evaluate for intracranial hemorrhage
- Diagnose malignancies, leukemia
- Investigate central nervous system disorders

Sample collection and Processing of CSF samples

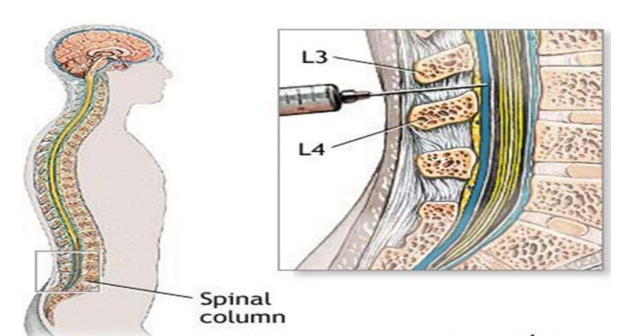
- Samples collection
- Sample transport and storage
- Gross description (Appearance)
- Counting cells
- Gram stainning and others
- · Culturing using suitable media
- Other identification techniques

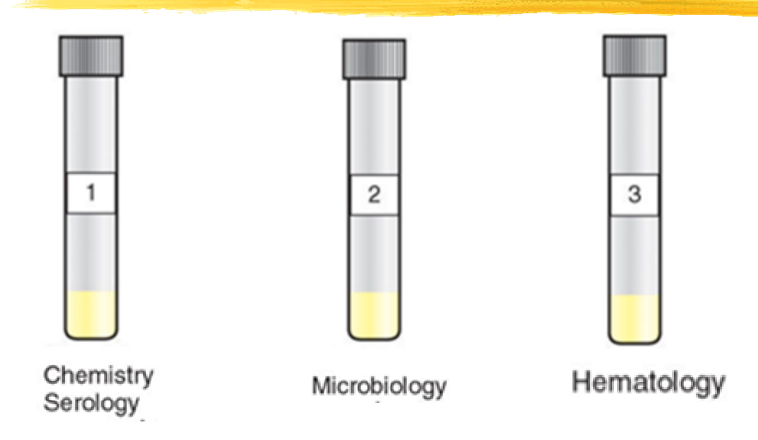
Specimen collection and handling

Routinely collected via lumbar puncture between 3rd & 4th, or 4th & 5th lumbar vertebrae under sterile conditions

Intracranial pressure measurement taken before fluid is

withdrawn.



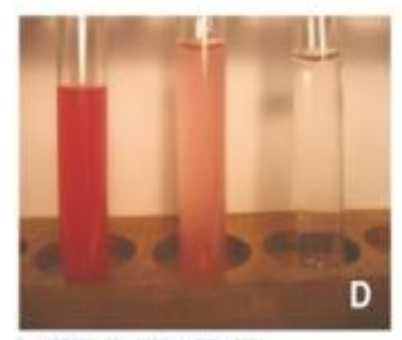


Testing considered STAT=Short TurnAround Time Specimen potentially infectious

CSF is collected by aseptically

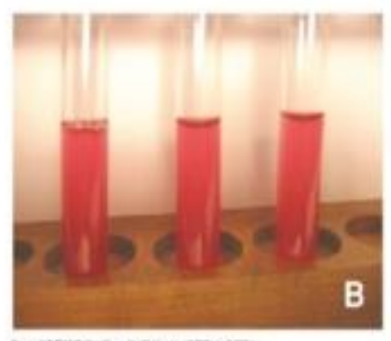
- Three tubes of CSF should be collected into sterile collection tubes that contain no additives.
- The tubes are numbered sequentially in the order in which they were collected along with the patient's name.
- tube 1 is used for chemistry studies, as well as immunology studies, as these tests are least affected by the presence of blood cells or bacteria introduced as a result of the spinal tap procedure
- tube 2 is used for culture, allowing a larger proportion of the total fluid to be concentrated.
- tube 3 is used for cell count and differential, as these tubes are least likely to contain cells introduced by the collection procedure

Traumatic Tap



Engraph & Str. Street House Built Linearch Williams & Miller

Intracranial Hemorrhage



Transport 2017 Written Classer Haatti Congraved Williams & William

Sample transport and storage

- Specimens for microbiology studies should never be refrigerated; if not rapidly processed, CSF should be incubated (35°C) or left at room temperature (< 1h).
- The CSF for viral studies may be refrigerated for as long as 24 hours after collection or frozen at -70° C if a longer delay is anticipated (should never be frozen at temperatures above -70° C).
- CSF specimen for hematology studies can be refrigerated.
- The CSF for chemistry and serology can be frozen (-20°C).
- RBC lyse in 1 hour, WBC in 2 hrs.

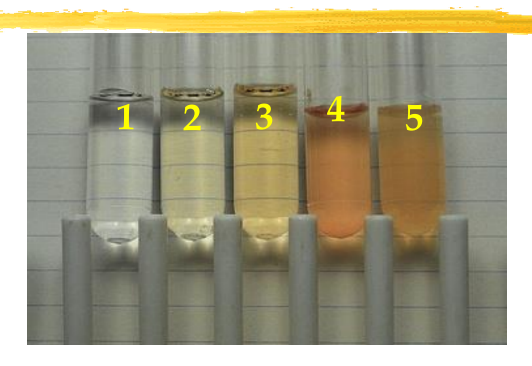
Gross description (Appearance)

Appearance

- Normal Crystal clear,
 colorless
- Descriptive Terms hazy, cloudy, turbid, milky, bloody, xanthrochromic
- Unclear specimens may contain increased lipids, proteins, cells or bacteria.
- ·Clots indicate trauma
- ·Milky increased lipids
- Oily contaminated with x-ray media



Gross description (Appearance)

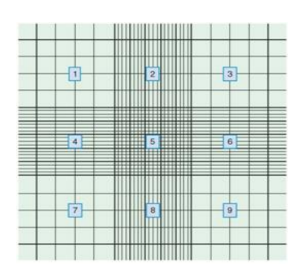


- 1- Normal CSF.
- 2- Mildly xanthochromic CSF.
- 3- Moderately xanthochromic CSF.
- 4- Red-tinged turbid CSF caused by hemorrhage
- 5- Cloudy red-tinged fluid with bacterial meningitis

Counting cells

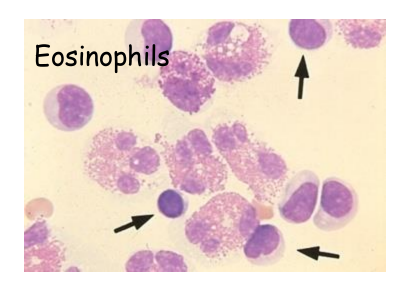
Counting cells in CSF samples

- All specimens should be examined microscopically hematology
- Manual count
- Standard hemacytometer counting chamber
- Normal CSF results
- RBCs: no cells regardless of age
- WBCs: 0 5 white blood cells (all mononuclear)

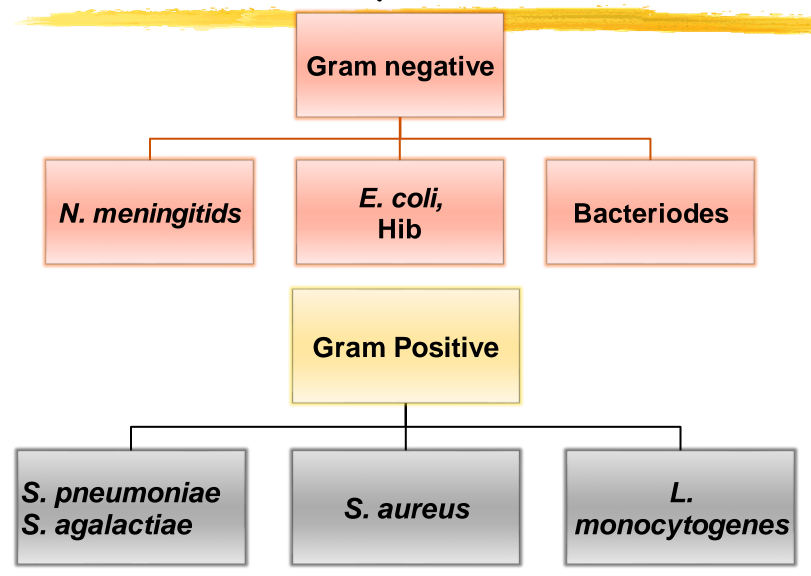


Counting cells

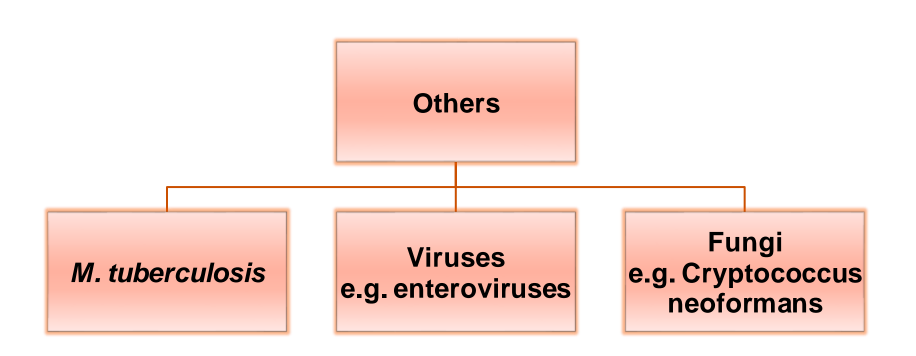
- Count and differentiate 100 nucleated cells.
- Entire smear should be evaluated for
 - abnormal cells, inclusions within cells, Clusters, Presence of intracellular organisms
- Eosinophils: often associated with parasitic / fungal infections, allergic reactions including reaction to shunts and other foreign objects



CNS - acute infectious disease



CNS - acute infectious disease

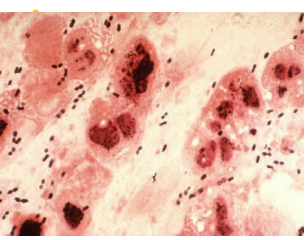


Gram stainning and others

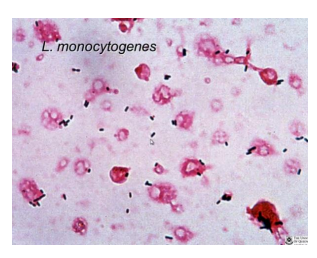
Visual Detection of Etiologic Agents

Stained Smear of Sediment.

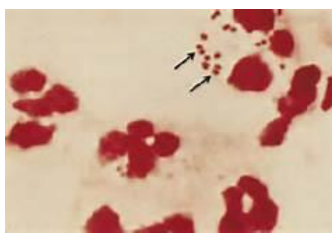
 Gram stain must be performed on all CSF sediments. The drop of sediment is allowed to air dry, is heat or methanol fixed, and is stained by either Gram or acridine orange.



Gram +ve diplococci S. pneumoniae



gram-positive, rod-shaped,

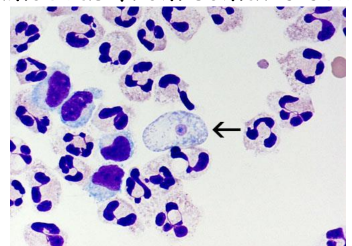


N. Meningitidis gram-negative, coffee-bean shaped diplococci.

Gram stainning and others

Wet Preparation:

- Amoebas are best observed by examining thoroughly mixed sediment as a wet preparation under phase-contrast microscopy.
- Amoebas are identifiable by their typical slow movement in one direction via pseudopodia.
- A trichrome stain can assist in the differentiation of amoebas from somatic cells.

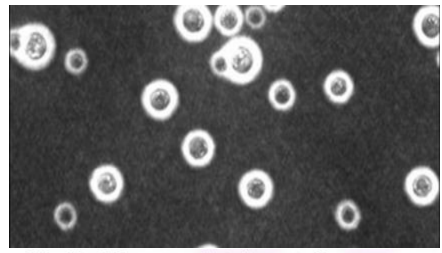


Naegleria fowleri trophozoite (arrow) stained with Giemsa-Wright stain

Gram staining and others

India Ink Stain.

- The large polysaccharide capsule of Cryptococcus neoformans allows these organisms to be visualized by the India ink stain.
- Latex agglutination testing for capsular antigen is more sensitive and specific.
- To perform the India ink preparation, a drop of CSF sediment is mixed with onethird volume of India ink
- Examination under oil immersion.

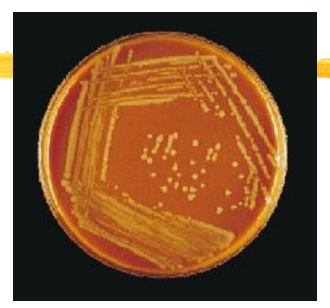


The Budding capsulated yeast Cryptococcus neoformans as shown in India ink wet mount.

Culturing using suitable media



S. pneumoniae on sheep blood agar with colistin and aztreonam (CAP)



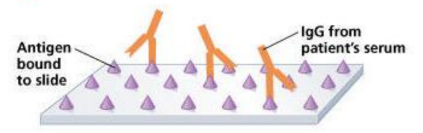
H. influenzae on a chocolate agar plate

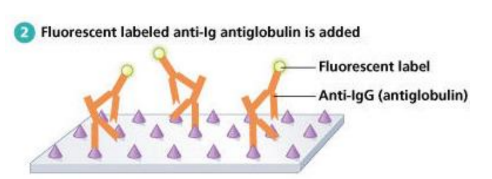


N. meningitidis on a BAP

Other identification techniques

- Serology
- ELISA
- Latx agglutination
- PCR
- IFA
- Antigen is attached to slide and flooded with patient's serum





Notes

- The volume of CSF is critical for detecting certain microorganisms, such as mycobacteria and fungi. A minimum of 5 to 10 mL is recommended for detecting these agents by centrifugation and subsequent culture.
- When the laboratory receives an inadequate volume of CSF, the physician should be consulted regarding the order of priority for laboratory tests. Processing too little specimen lowers the sensitivity of laboratory tests, which may lead to falsenegative results.