Gastrointestinal Tract Module Bacterial infections Practical session

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Objectives

To become familiar with:

- ✓ the selective and differential media used to isolate the
 GIT infections associated bacteria
- ✓ the biochemical tests used to identify these bacteria

Types of specimen

- ✓ Stool (direct culture)
- ✓ Stool in fecal transport system (in delay longer than 1 hour)



- ✓ Rectal swab
 - Pass swab beyond anal sphincter, carefully rotate, and withdraw

Criteria of specimen rejection

- ✓ Specimen contaminated with urine, residual soap, or disinfectants.
- ✓ Specimens received in grossly leaking transport containers
- ✓ Dry specimens
- ✓ Specimens submitted in fixative or additives

Patient preparation

- ✓ Instruct the patient on how the specimen should be collected and transferred to the container
- ✓ provide him with sticks and containers

Specimen collection

- ✓ A sample is transferred with the sticks to the container
- √ The specimen should contain at least 5 g of feces
- ✓ Those parts that contain blood and/or mucus should be selected.
- ✓ Close the lid properly

Who will collect the specimen?

The patient. If stool is unobtainable, nursing staff or physician will collect fecal swab

Time relapse before processing the sample

Stool samples should be examined and cultured as soon as possible after collection.

Indications for stool culture include:

Doctors are most likely to order a stool culture for patients with any of the following characteristics:

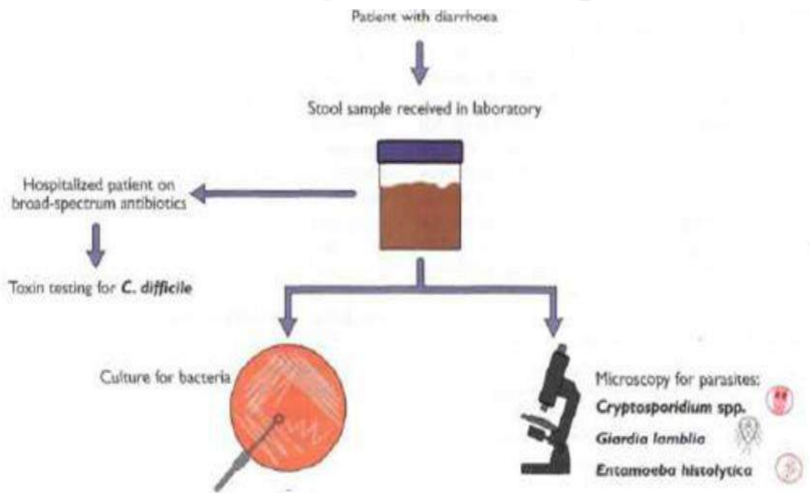
- ✓ Bloody diarrhea and/or fecal leukocytes
- ✓ Tenesmus (is the constant feeling of the need to empty the bowel, accompanied by pain, and cramping)
- ✓ Severe or persistent symptoms
- ✓ Recent travel to a third world country
- ✓ Severe dehydration

Enterobacteriaceae

General Characteristics

- Gram-negative bacilli
- Oxidase –ve
- Catalase +ve
- Ferment glucose with or without gas production
- Reduce nitrate to nitrite (with few exceptions)
- facultative anaerobes

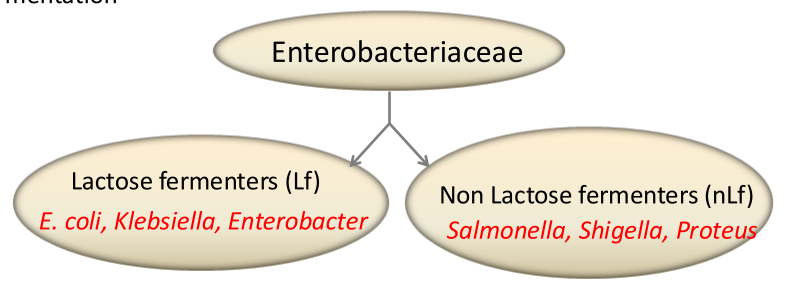
Processing of stool samples



Identification of *Enterobacteriaceae*

1- Using selective and differential media

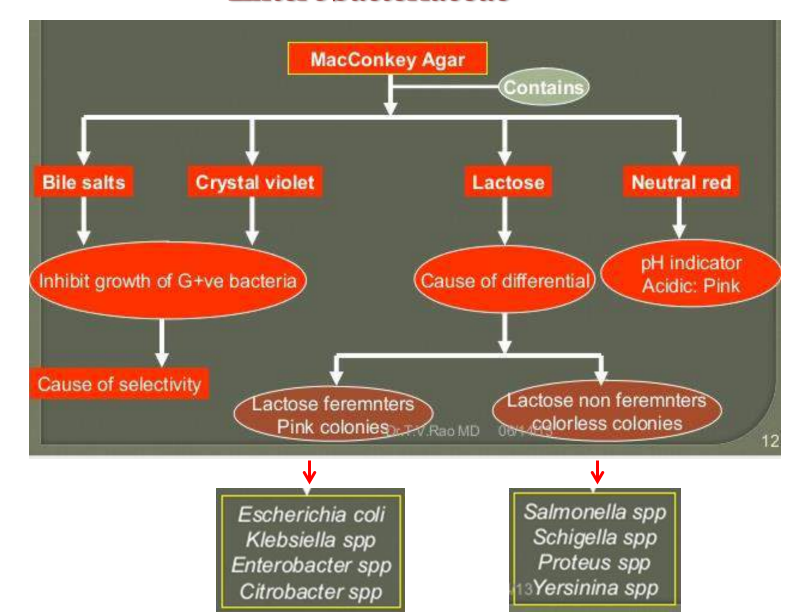
Enterobacteriaceae are divided into two main groups according to lactose fermentation



There are several selective and differential media used to isolate and distinguish between Lf & nLf including

- ✓ MacConkey agar
- √ Salmonella Shigella agar (SS agar)

MacConkey agar is a selective and differential media for Enterobacteriaceae



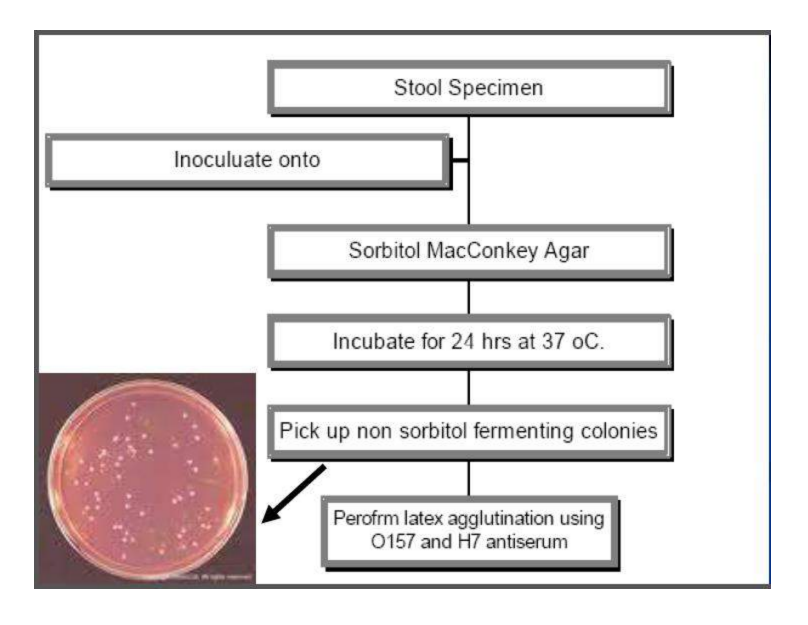
MacConkey agar



Lactose fermenter

Non Lactose fermenter

Suspected EHEC (O157:H7)



Diagnosis of Salmonella & Shigella

1- Salmonella Shigella agar (SS agar)

Purpose

For isolation and differentiation of Salmonella & Shigella

Components

- ✓ Brilliant green dye & sodium citrate: inhibit the growth of most intestinal flora
- ✓ Lactose
- ✓ Neutral red: pH indicator, red in acidic conditions
- ✓ Sodium thiosulfate (Na₂S₂O₃): sulfur source
- ✓ Ferric citrate: H2S indicator

Diagnosis of Salmonella & Shigella

1- Salmonella Shigella agar (SS agar)

Results

- ✓ Lactose fermenters: pink to red colonies (few can grow)
- ✓ Non lactose fermenters: translucent, colorless colonies with or without black centers

Shigella colorless colonies without black centers



Lactose fermenter flora: pink to red colonies

Salmonella:
colorless colonies



Diagnosis of Salmonella & Shigella

2- Hektoen enteric agar (cat eye apperance)



Identification of Enterobacteriaceae

2- Using special biochemical reactions

The differentiation of the principle groups of Enterobacteriaceae Can be accomplished on the basis of <u>their biochemical prosperities</u> and enzymatic reactions in <u>the presence of the specific substrate</u>

One important group of biochemical reactions is:

Indole, Methyl red, Vogus proskaur, Citrate utilization tests (IMViC)

E. Coli

Glucose	Lactose	Maltose	Mannite	Sucrose	Indole	MR	VP	Citrate	urease	H2S
A/G	A/G	A/G	A/G	A/G	+	+	-	-	-	-

SH. dysenteriae

Glucose	Lactose	Maltose	Mannite	Sucrose	Indole	MR	VP	Citrate	urease	H2S
Α	-	-	-	-	-	+	-	-	-	-

SH. flexneri

Glucose	Lactose	Maltose	Mannite	Sucrose	Indole	MR	VP	Citrate	urease	H2S
Α	-	-	A	-	-	+	-	-	-	-

SH. sonnei

Glucose	Lactose	Maltose	Mannite	Sucrose	Indole	MR	VP	Citrate	urease	H2S
Α	Α	-	Α	-	-	+	-	-	-	-

A/G produce Acid and Gas **A** produce Acid only

Cholera identification

Identification

- ✓ Gram negative curved rods (comma shape).
- ✓ Motile (single polar flagellum)
- ✓ Oxidase positive, catalase positive

culture

- ✓ Thiosulfate citrate bile salt sucrose agar or TCBS agar
- ✓ The medium is alkaline (pH 8.6) which enhances the growth of Vibrio species

Important components

- ✓ Sucrose: sugar source
- ✓ Bromothymol blue: pH indicator
- pH<6.0 yellow pH>7.6 -blue

Cholera identification

Results

- ✓ Vibrio cholera: Ferment sucrose smooth yellow colonies
- ✓ Vibrio parahemolyticus: non-sucrose fermenter, green colonies

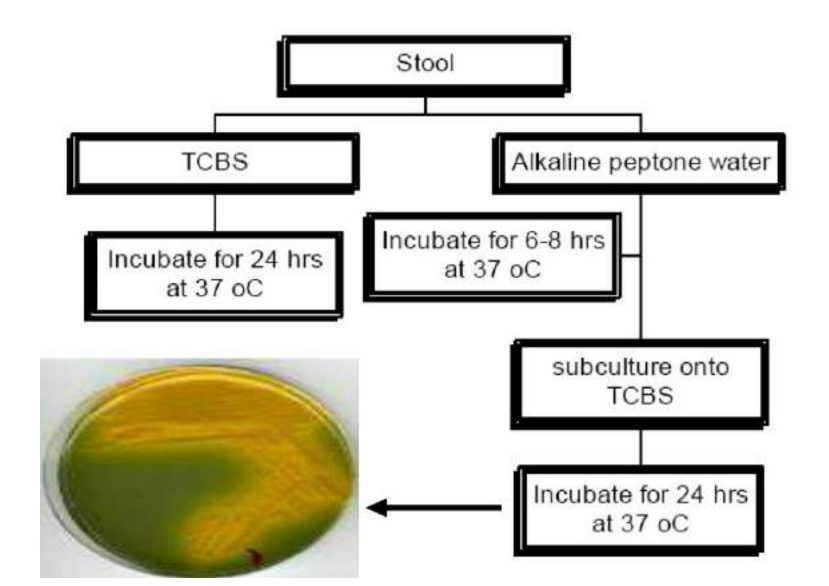


TCBS media

V. cholera

V. parahemolyticus

Cholera identification



S. aureus associated food poisoning

- > 25% of healthy people are carriers
- ➤ Mainly *S. aureus* food poisoning is diagnosed in case of outbreaks
- Diagnosis based on
 - ✓ gram positive cocci
 - √ catalase & coagulase positive
 - ✓ DNase positive
 - √ ß-hemolysis on blood agar
 - ✓ Grow on MSA with mannitol fermentation



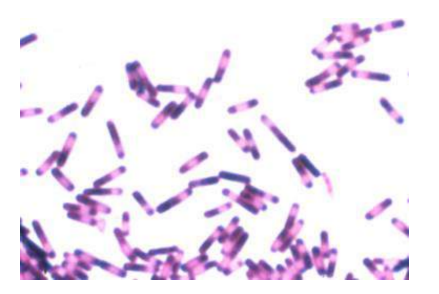
Blood agar

MSA

Clostridia

- Rod-shaped
- > Gram positive anaerobic
- > spore-forming
- Diagnosis
- Clinical picture
- Gram stain
- Toxin detection by serological techniques
- Cycloserine-Cefoxitin Fructose Agar CCFA

is recommended as a selective and differential medium for the primary isolation of *C. difficile* from fecal specimens.



Diagnosis of C. difficile infection

Routine Laboratory Diagnosis of CDI

CURRENT STRATEGY

Toxin detection Fresh Stools Toxin Culture EIA Colony

NEW RECOMMENDED STRATEGIES



Molecular testing

> Fresh Stools



Diagnosis of *C. difficile* infection Culture

- Colonies of C. difficile bacteria after 24-48 hours growth on Cycloserine-Cefoxitin Fructose Agar — CCFA.
- Results: Clostridium difficile will appear as yellow colonies with a ground glass-like appearance and a slightly filamentous edge.



