CULTIVATION OF BACTERIA

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CULTIVATION OF AEROBIC BACTERIA
For bacteria of veterinary importance, aerobic incubation is uniformly done at 37ºC.

Depending upon the workload a laboratory may have a tabletop incubator or a walk-in incubator.

For prolonged incubations, as required for the growth of *Mycobacterium tuberculosis*, screw-capped bottles should be used instead of Petri dishes or tubes to prevent the drying of medium.
CULTIVATION OF ANAEROBIC BACTERIA
Methods for cultivation of anaerobes

Evacuation and replacement of oxygen atmosphere in sealed jars

- Brewer jar
- Gas Pak system
- Chromium-sulphuric acid method

Specialized techniques where evacuated sealed jars are not used

- Solid medium
  - Shake culture method
  - Pyrogallic acid method
- Liquid medium
  - Paraffin plug method
Incorporation of reducing agents into the liquid medium

- Freshly steamed liquid media are at least temporarily anaerobic, but soon become aerobic unless a reducing agent is added.

- Reducing agents include glucose 0.5-1%, ascorbic acid 0.1%, cysteine 0.1%, sodium mercaptoacetate or thioglycollate 0.1%, or the particles of meat in cooked meat broth.

- Examples - Thioglycollate broth, Robertson's cooked meat medium

- Addition of 0.05-0.1% agar can further increase the effectiveness of reducing agents

- Liquid media should be `pre-reduced' by holding in a boiling water bath for 10 min, to drive off dissolved oxygen
Incorporation of reducing agents in Petri dish lids (Brewer’s method)

- A simple and safe method of obtaining anaerobiosis when using solid medium.

- Sodium dithionite (reducing agent), together with sodium bicarbonate and sodium carbonate (to supply CO₂), is added in the lid of a Petri dish.

- Sodium dithionite removes the oxygen to give anaerobic conditions.
Displacement of oxygen by inert gases

Anaerobic jar using hydrogen

- Most of the air from the anaerobic jar is removed and replaced with hydrogen or preferably hydrogen mixed with nitrogen in the presence of a catalyst.

- The hydrogen reacts with the remaining oxygen to form water.

- A common catalyst used is palladium, which should be completely dried.

- Anaerobiasis is checked by a chemical indicator and also a bacterial indicator.
Metal anaerobic jar
(McIntosh and Filde’s jar)

Anaerobic jar using a gas generating kit

In this system, the hydrogen required to combine with the free oxygen in the anaerobic jar is slowly released from chemicals contained in a foil envelope. The chemicals are activated by the addition of water. A low temperature catalyst is used to bring about the reaction and an indicator is enclosed to check for anaerobiosis.
Use of copper coated steel wool to remove oxygen

- uses steel wool, which is activated immediately before use by being dipping in acidified copper sulphate solution

- the metallic copper on the surface of steel rapidly absorbs oxygen.

- plates are incubated in an airtight plastic bag from which the air is removed before sealing

The Hungate procedure

- Anaerobes are grown in roll tubes in which a thin layer of agar coats the inside of the tube.
- The tubes are incubated under oxygen-free gas.
- Transparent media are preferred so that surface colonies are easily visible
CULTIVATION OF MICROAEROOPHILIC BACTERIA
For the culture of truly microaerophilic species such as *Campylobacter jejuni* and *Actinomyces israelii*, an atmosphere of 5-10% O₂ is needed.

This can be done by an evacuation replacement method with N₂ as the major replacement gas along with 5-10% CO₂.
CULTIVATION OF CAPNOPHILIC BACTERIA
A candle jar
Extra carbon dioxide @ 3-5% is needed for optimal growth of capnophilic organisms such as *Brucella abortus* and *Campylobacter*.

The simplest method for having this environment is to put the plates in a container and generate CO$_2$ inside by lighting a candle in it just before putting on the lid.

As the candle burns, the oxygen content is reduced leaving a carbon dioxide content of 3-5% by the time the candle is extinguished.

Carbon dioxide can also be generated chemically by reacting sodium bicarbonate with tartaric acid or citric acid.

Commercial CO$_2$ generating kits and special CO$_2$ incubators (also called capnoeic incubators) are available commercially, which can provide a predetermined and regulated amount of this gas.
MEDIA INOCULATION TECHNIQUES
Typical Culture Laboratory Bench
What is inoculation?

Inoculation means introduction of microorganisms into culture media and inoculum is the material used for inoculation.
INOCULATION METHODS

• Streak Plate method
• Spread Plate method
• Pour Plate method
• Spot Plate method
• Lawn or carpet culture method
• Stroke culture method
• Stab method
• Inoculation of liquid media
Inoculating wire

Nichrome wire No. 24 SWG is being used for routine inoculation. For working with the anaerobes either stainless steel or platinum-iridium is a better choice since nichrome is oxidizing. Similarly in some of the tests where oxidizing property of bacterium is to be tested (e.g. oxidase test), platinum wire, instead of nichrome should be used. The wire can be used as a:

- Wire loop that is usually of 2 mm diameter is most useful of all inoculating wires.
- Straight wire to stab the culture, picking of single colonies as well as for inoculating the liquid media.
INSTRUMENTS USED TO INOCULATE THE CULTURE MEDIA

Scalpel

- Used for making inoculations with scrapings from tissues and ulcers.

Sterile pipettes or Pasteur pipettes

- Graduated 1 ml or 10 ml glass pipettes or disposable glass or plastic pipettes are used for inoculation liquid between 0.1 and 10 ml.

- Sterilized Pasteur pipettes are also useful to collect fluids, blood, pus, etc.
STREAK PLATE METHOD

• Isolation of pure colonies (pure culture method – best method)

• The aim of a streak plate is to separate bacterial cells so that some are sufficiently separated ultimately to give rise to individually separate colonies, each derived from a single cell.
The most common way of separating bacterial cells on the agar surface so as to obtain single isolated colony.

It provides a simple and rapid method of diluting the sample by mechanical means.

As the loop is streaked across the agar surface, more and more bacteria are rubbed off until individual separated organisms are deposited on the agar.

After incubation, the area at the beginning of the streak pattern will show confluent growth, while the area near the end of the pattern should show discrete colonies.
A wore loop is sterilized by heating it until red hot in a flame and then allowed to cool for several seconds.

A loopful of liquid inoculum or bacterial growth is picked up aseptically from the broth culture tube or surface of an agar plate and streaked lightly back and forth with the flat loop, making parallel streaks back to the edge of the plate in the form of a primary inoculum (A in Fig), which is called as ‘well’.

The loop is sterilized and cooled again. Then with the edge of the loop, the successive series of strokes B, C, D and E are made with the loop sterilized between each sequence.

Each set of streaks is made in one direction, perpendicular to and crossing the second set of streaks, but avoiding the first set.

At each step the inoculum is derived from the most distal part of the immediately preceding strokes so as to gradually reduce the number of bacteria. This helps in obtaining isolated colonies.
1. Start

2. Flame & Cool

3. Flame & Cool

4. Do not flame

5. Start

6. Start

Start

1. Flame & Cool

Rotate counterclockwise.

2. Flame & Cool

Spread area 1 over area 2.

3. Flame & Cool

Spread area 2 over area 3.

4. Flame & Cool

Spread area 3 over area 4.

5. Start

6. Start

Start

1. Flame & Cool

2. Flame & Cool

3. Flame & Cool

4. Do not flame

5. Flame & Cool

6. Flame & Cool

Rotate counterclockwise.

Spread area 1 over area 2.

Spread area 2 over area 3.

Spread area 3 over area 4.

Spread area 4 over area 5.

Spread the loop over area 6.

Sterilize the loop.

Sterilize the loop.

Sterilize the loop.
SPREAD PLATE METHOD

• Isolation of pure colonies (pure culture method)

• Method for counting of bacteria in a given sample (indirect bacterial counting method – best)
In the spread-plate method of inoculation of cells to solid medium, ten fold serial dilutions of the test samples or suspensions of microorganisms are prepared in suitable diluent.

A small volume (0.1ml) of each dilution is dropped onto the surface of agar that has already hardened in a Petri dish.

The volume is then spread around the agar surface across the surface using a bent glass rod.

The plates are incubated for 24 - 48 hours at 25 - 37 °C.

After incubation, plates inoculated with a sample dilution yielding between 30 and 300 colonies are selected for counting using a colony counter.

This technique is advantageous particularly when cells are sensitive to exposure to relatively high temperatures plus the method does not require a prior melting of the solid medium.
Dilution procedure

Sample of *E. coli*

- 1 ml
- 1 ml
- 1 ml
- 1 ml
- 1 ml
- 1 ml

- 9 ml
- 9 ml
- 9 ml
- 9 ml
- 9 ml
- 9 ml

- $10^{-1}$
- $10^{-2}$
- $10^{-3}$
- $10^{-4}$
- $10^{-5}$
- $10^{-6}$

- 0.1 ml
- 0.1 ml
- 0.1 ml

- $10^{-5}$
- $10^{-6}$
- $10^{-7}$
POUR PLATE METHOD

• Isolation of pure colonies (pure culture method)

• Method for counting of bacteria in a given sample (indirect bacterial counting method).

• The bacteria are mixed with melted agar until evenly distributed and separated throughout the liquid. The melted agar is then poured into an empty plate and allowed to solidify.
A standard volume of solution is mixed thoroughly in the sterile liquefied agar (45-50 °C) in a test tube so that each organism in the solution is separated from all others (primary inoculum).

- Ten fold serial dilutions of primary inoculum are then prepared in sterile melted agar tubes.

- The inoculated agar from each tube is poured in sterile Petri plate, allowed to solidify and incubated for 24 - 48 hours at 25 - 37 °C.

- When the agar solidifies the cells are trapped in the agar and develop into colonies appear within, beneath, and on top of the agar; each colony can be counted and represents a single cell in the original solution.

- After incubation, plates inoculated with a sample dilution yielding between 30 and 300 colonies are selected for counting using colony counter.

- Disadvantages:
  a) The cells do not develop typical colonial morphology and are not easily accessible for further testing.
  b) It is not useful for temperature sensitive bacteria.
  c) It is difficult to count sub-surface colonies.
Serial dilution

1.0 ml

Original sample

9 ml H₂O (1/10 dilution)

1.0 ml

9 ml H₂O (1/100 dilution)

1.0 ml

9 ml H₂O (1/1000 dilution) (1/10,000 dilution)

1.0 ml

0.1 ml
Mix with warm agar and pour
SPOT PLATE METHOD

• Useful for procedures, such as phage typing, bacteriocin testing, haemolysin testing and antimicrobial sensitivity testing.
• A loopful of pure bacterial growth is inoculated as a spot on surface of agar plate.
CARPET OR LAWN CULTURE METHOD

• Used for antibiotic sensitivity testing (disc-diffusion method) and in bacteriocin or phage typing.

• A suspension of bacterial culture is evenly smeared on the surface of a sterile, well-dried agar plate using a sterile cotton swab. The excess material is discarded and plate is incubated, after allowing the inoculum to be adsorbed onto the surface of the media.
STROKE CULTURE METHOD

• Used for maintenance and preservation of pure cultures
• The agar slopes are inoculated using a sterile loop to streak the inoculum down the center of the slope and then the inoculum is spread in a zigzag pattern as shown in the figure.
STAB METHOD

• Used for demonstration of bacterial motility

• Usually semi-solid agar medium

• Inoculation by straight needle taking care to withdraw the wire along the line of inoculum without making further stab lines.
INOCULATION OF LIQUID CULTURE

• Broths and other fluid media are inoculated using a sterile loop, straight wire, or Pasture pipette depending on whether the inoculum is columnar growth or a fluid culture or specimen.

• If a wire loop is used to inoculate, the bottle or tube containing media held at an angle and the loop is rubbed against the side of the container below the level of the fluid.

• The neck of the bottle or tube should be flamed before and after inoculating the medium while the cap or cotton plug is held in the hand.
Important points about inoculation of culture media

- Aseptic technique is important to avoid contamination.
- When more than one medium is inoculated, follow a particular order. Inoculate media without inhibitors, followed by indicator and then selective media.
- While processing fluid specimen inoculate liquid media first to reduce the chances of carry over from contaminated solid media.
- Properly label the media to be inoculated to avoid any mix-up of the specimens.
- Inoculate the media with clinical specimens as soon as possible.
- Minimize the aerosol production by opening the caps of liquid media slowly, avoiding vigorous shaking of the specimen and avoiding the expulsion of the last drop from the pipette.
BACTERIOLOGICAL MEDIA
The composition of a medium is depended on the nutritional requirement of the microorganism to be cultivated, which are of the following two types:

- **Copiotrophic microorganisms**: These microorganisms need a growth medium that is rich in nutrients (peptone, beef or yeast extract etc.). Such a medium is called complete medium. Many pathogenic bacteria require such an environment.

- **Oligotrophic microorganism**: These microorganisms require only low concentrations of nutrients and, therefore grow on a minimal medium. These microorganisms can be isolated from soil, fresh water or marine environments.
COMMON INGREDIENTS OF CULTURE MEDIA

- **Peptone**: acid or enzymatic digest of meat, milk or soya bean meal; provides nitrogen for growing microorganisms.

- **Meat extracts**: provides organisms with a supply of amino acids, and also, with essential growth factors, vitamins and minerals salts.

- **Mineral Salts**: sulphates and phosphates; traces of magnesium, potassium, iron, calcium and other elements, which are required for bacterial enzyme activity. Sodium chloride is also an essential ingredient of most culture media.

- **Carbohydrates**: Simple or complex sugars are added to many culture media to provide bacteria with sources of carbon and energy.

- **Agar**: an inert polysaccharide extract obtained from a variety of red purple seaweed (*Rhodophyceae*); used to solidify culture media. It gels below 45ºC temperature and remain in molten form above that.

- **Water**: is essential for growth of all microorganisms; deionised or distilled water must be used in the preparation of culture media.
TYPES OF CULTURE MEDIA
(on the basis of form)

LIQUID
(Broth)

SOLID

Agar slant/slope

Agar plate

SEMI-SOLID
(Stab / Deep)
LIQUID MEDIUM

A liquid medium is commonly called as broth, e.g. nutrient broth.

After incubation, growth in broth/liquid medium may be observed as one or a combination of three forms:

- **Pellicle**: A mass of organisms is floating on top of the broth.
- **Turbidity**: The organisms appear as a general cloudiness throughout the broth.
- **Sediment**: A mass of organisms appears as a deposit at the bottom of the tube.
LIQUID MEDIUM - USES

1. When bacteria are present in small numbers in the inoculum they will grow only in liquid media, e.g., blood culture.

2. Liquid media are widely used for biochemical tests and as enrichment media.

3. Large sized inoculum can be tested easily in liquid media e.g., gauge for sterility test.

4. Presumptive bacterial count in water or milk sample is made in liquid media.

5. Isolation of bacteria in pure culture and identification of bacteria is not possible in liquid medium.
Solid media contains agar, which is a compound that goes into water solution at temperatures approaching boiling and solidifies at room (<40ºC) temperature.

Once boiled, agar-containing medium will stay liquid at 45ºC.

Solid medium are poured in molten form into various vessels, such as Petri plates, Roux flasks and test tubes, at temperatures that will not kill cells followed by a solidification.
Slant tubes (Figure 2A) are tubes containing a nutrient medium plus a solidifying agent, agar-agar. The medium has been allowed to solidify at an angle in order to get a flat inoculating surface (Figure 2B).
**Agar plates** are sterile Petri plates that are aseptically filled with a melted sterile agar medium and allowed to solidify. Plates are commonly used in the culturing, separating, and counting of microorganisms (Figure 3).
Advantages of solid media

- Single, well isolated colonies can be grown, which assist in isolation of bacteria as pure culture.
- By studying colonial morphology presumptive identification of most bacterial species can be made.
- Quantitative bacterial count and relative proportion of different bacterial species in the inoculum can be made.
SEMI – SOLID CULTURE MEDIA

These are prepared by adding agar @ 0.4 – 0.5% (w/v) to a medium and pouring in test tubes as Stab tubes (deeps). These are used mainly as transport media and also for the testing of motility of the organisms. These are inoculated by "stabbing" the inoculum into the agar using a straight wire.
TYPES OF CULTURE MEDIA
(on the basis of composition and use)

1. Synthetic media
2. Complex medium
3. Basic media
4. Enriched media
5. Enrichment media
6. Selective media
7. Differential media
8. Indicator media
9. Transport media
10. Maintenance media
11. Assay media
SYNTHETIC MEDIUM

- Also known as **chemically defined** medium.
- Produced only from well-defined, relatively **pure ingredients**; the exact chemical composition is known.
- Used for growth of **chemoautotrophs** and **photoautotrophs**, and microbiological assays.

COMPLEX MEDIA

- Also known as **chemically undefined** media.
- Produced from **ingredients** that are neither well-defined nor pure.
- Complex media may contain **extracts from animals** (e.g., beef, hearts, milk, etc.), **plants** (e.g., soy beans), or **microorganisms** (e.g., yeast); the exact chemical composition is not known.
- Used for growth of most **chemoheterotrophic organisms**.
BASIC MEDIA

- Support the growth of microorganisms that do not have special nutritional requirements, i.e. non-fastidious bacteria.

- Example are simple media such as nutrient agar and nutrient broth.

- Used in the preparation of enriched media, to maintain stock cultures of bacteria, and for sub culturing pathogens from differential or selective media prior to performing biochemical and serological identification tests.
ENRICHED MEDIA

- Designed to support the growth of fastidious pathogens that requires additional nutrients or growth stimulants.
- Basic media supplemented with whole blood, serum, extra peptones, special extracts, or vitamins.
- Examples are blood agar, serum agar and Tryptone soya agar.
- Enriched media are required for the culture of *Haemophilus influenzae*, pathogenic *Nisseria*, and several *Streptococcus* species.
**INDICATOR MEDIA**

- Contain an ingredient that give a coloured reaction following growth of a particular bacterial species.
- Examples are MacConkey agar, XLD agar, EMB agar, Brilliant green agar, and blood agar.

Clockwise from top left, XLD, brilliant green, Mac Conkey & EMB agars
ENRICHMENT MEDIA

- A fluid medium that favours the growth of a bacteria over others by containing substances that encourage the multiplication of desired bacteria.

- Useful for isolation of those bacteria that are present in very small numbers in the initial inoculum.

- Examples - selenite F broth is used as an enrichment medium for isolation salmonellae from faeces or urine, GN Hajna broth for *Shigella*, and alkaline peptone water for *Vibrio cholerae*. 
SELECTIVE MEDIA

- Inhibits the growth of some bacteria while selecting for the growth of others

- Example:
  
  Brilliant Green Agar, which contains the dye that inhibit the growth of Gram (+) bacteria and selects for Gram (-) bacteria

  XLD agar selects for salmonellae and shigellae by containing bile salts that inhibit the growth of many faecal commensals.

- In recent years, antimicrobials have become increasingly used as selective agents in culture media, e.g., York Butzler medium for isolation of Campylobacter species from faeces.

- Selective media are available for isolation of most of the important pathogens.
DIFFERENTIAL MEDIA

- Differentiates between different organisms growing on the same plate on the basis of colonial morphology.
- Contains ingredients that allow different bacteria to look reproducibly different.
- Majority of differential media distinguish between bacteria by an indicator, which changes colour when acid is produced following carbohydrate fermentation.
- Examples – EMB agar, blood agar, MacConkey agar.
DIFFERENTIAL MEDIA

Example - Blood Agar Plates (TSA with 5% sheep blood) used to differentiate different types of Streptococci

Alpha Hemolytic Streptococci - Incomplete lysis of RBC’s.
Beta Hemolytic Streptococci - Complete lysis of RBC’s.
Gamma Hemolytic Streptococci - No lysis of RBC’s
Many culture media are both differential and selective such as EMB agar, MacConkey agar, XLD agar and DCA.

These media inhibit the growth of certain microorganisms while making those organisms that it allows to grow easily produce differentiated colony morphologies.
MacConkey agar contains bile salts and crystal violet that inhibits Gram (+) bacteria, while Gram (-) enteric bacteria can grow on it. Further MCA has lactose as a sugar and neutral red as a pH indicator. Lactose fermenters (*E. coli*) produce pink colonies, while non-fermenters (salmonellae) produce pale colonies.
Enterobacter aerogenes

Escherichia coli
**Assay Media:** Media of prescribed composition used for the assay of vitamins, amino acids or antibiotics by microorganisms.

**Maintenance Media:** These media are used for preserving the viability of a bacterial culture. Such media provides minimal nutrient requirements of a bacterium.

**Transport Media:** These are mostly semisolid media that contain ingredients to prevent the overgrowth of commensals and ensure the survival of aerobic and anaerobic pathogens when specimens cannot be cultured soon after collection. Their use is particularly important when transporting microbiological specimens from field to the disease investigation laboratory. e.g., Cary-Blair medium and Amies transport medium.
## TYPES OF MEDIUM

<table>
<thead>
<tr>
<th>TYPE</th>
<th>PURPOSE</th>
</tr>
</thead>
<tbody>
<tr>
<td>Chemically defined</td>
<td>Growth of chemoautotrophs and photoautotrophs, and microbiological assays.</td>
</tr>
<tr>
<td>Complex</td>
<td>Growth of most chemoheterotrophic organisms.</td>
</tr>
<tr>
<td>Reducing</td>
<td>Growth of obligate anaerobes.</td>
</tr>
<tr>
<td>Selective</td>
<td>Suppression of unwanted microbes; encouraging desired desired microbes.</td>
</tr>
<tr>
<td>Differential</td>
<td>Differentiation of colonies of desired microbes from others.</td>
</tr>
<tr>
<td>Enrichment</td>
<td>Similar to selective media but designed to increase numbers of desired microbes to detectable levels.</td>
</tr>
</tbody>
</table>
pH indicators
pH indicators give characteristics color in different pH (alkaline, neutral and acidic) e.g., phenol red gives yellow color in acidic pH so whenever fermenter grow in sugar-containing media, they produce acid and give yellow-colored colonies

<table>
<thead>
<tr>
<th>Indicator</th>
<th>Low pH color</th>
<th>Low pH range</th>
<th>High pH range</th>
<th>High pH color</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bromothymol blue</td>
<td>yellow</td>
<td>6.0</td>
<td>7.6</td>
<td>blue</td>
</tr>
<tr>
<td>Phenol red</td>
<td>yellow</td>
<td>6.4</td>
<td>8.0</td>
<td>red</td>
</tr>
<tr>
<td>Neutral red</td>
<td>red</td>
<td>6.8</td>
<td>8.0</td>
<td>yellow</td>
</tr>
</tbody>
</table>

**pH indicators and their range**
**pH Indicators used in different culture media**

<table>
<thead>
<tr>
<th>Indicator</th>
<th>pH range:</th>
<th>Indicator</th>
<th>pH range:</th>
<th>Indicator</th>
<th>pH range:</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bromothymol Blue</td>
<td>6.0 (yellow)- 8.0 (blue)</td>
<td>Phenol Red</td>
<td>6.8 (yellow)- 8.4 (red)</td>
<td>Neutral Red</td>
<td>6.8 (red)- 8 (yellow)</td>
</tr>
<tr>
<td>Cystine Lactose Electrolyte Deficient Agar (CLED)</td>
<td></td>
<td>Mannitol Salt Agar</td>
<td></td>
<td>MacConkey Agar</td>
<td></td>
</tr>
<tr>
<td>Thiosulfate Citrate Bile Salts Sucrose (TCBS) Agar</td>
<td></td>
<td>Triple Sugar Iron (TSI) Agar</td>
<td></td>
<td>Deoxycholate Citrate Agar (DCA)</td>
<td></td>
</tr>
<tr>
<td>OF Medium</td>
<td></td>
<td>Urease Test Medium</td>
<td></td>
<td>Salmonella-Shigella Agar</td>
<td></td>
</tr>
<tr>
<td>Simmons Citrate Agar</td>
<td></td>
<td>Xylose Lysine Deoxycholate Agar (XLD)</td>
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</tr>
</tbody>
</table>
**pH indicator: Bromothymol blue (BTB)**

- It is a color indicator which turns **yellow** at acidic pH. At a neutral pH, bromothymol blue is **green**. At pH 7.5 or above, bromothymol blue turns **royal blue**.

- **Cystine Lactose Electrolyte Deficient Agar (CLED):** In CLED media, lactose fermenting colonies of *Escherichia coli* gives pale yellow colonies whereas non lactose fermenting colonies of *Proteus, Salmonella* etc gives blue colored colony.

- **Thiosulfate Citrate bile salts sucrose (TCBS) agar:** is used for the selective isolation of Vibrio spp. TCBS has very high pH (8.5 to 9.5) to supress other intestinal flora. pH indicator Bromothymol blue is used in this media, so if the organism utilizes the sucrose, it lowers the pH of the media and yellow colored colonies are seen in TCBS.

![Image of CLED agar with lactose fermenting and non-fermenting colonies](image)

**Lactose fermenting (Yellow colonies) and Lactose Non fermenting (blue colonies) in CLED**
pH indicator: Phenol red

- Yellow at pH 6.4 and below, Red at pH 8 and above.
- **Mannitol Salt Agar (MSA):**
  *Staphylococcus aureus* grows in MSA, ferments mannitol and produces (acid) yellow colonies with yellow zones, whereas most coagulase negative staphylococci (CNS) and micrococci do not ferment mannitol and grow as small red colonies surrounded by red or purple zones.
pH indicator: Phenol red

- **Urea Agar base**: Urease producing organisms gives pink red color in the media as they utilize urea with the formation of ammonia the pH of the media changed to alkaline condition.

- **Xylose Lysine Deoxycholate (XLD) Agar**: It is used for the isolation and differentiation of *Salmonella* and *Shigella spp* which appear as pink colored colonies (as they do not ferment carbohydrate) from other non pathogenic gram negative enteric bacilli which produce yellow colonies (as they ferment one or more of the sugars present in the media).
pH indicator: Neutral Red

- Red at pH 6.8 and below, yellow at pH 8 and above.
- **Deoxycholate Citrate Agar (DCA):** Lactose non-fermenter produces colorless colonies. Coliform bacteria, if present, form pink colonies.
- **Salmonella-Shigella Agar:** Lactose fermenter produces red pigmented colonies, whereas non-lactose fermenter (e.g., *Salmonella*) grows as translucent colonies (colorless) with or without black centers.
- **MacConkey Agar:** Lactose fermenting bacteria (e.g. *E. coli*) give pink colored colonies, whereas non-lactose fermenter gram negative bacilli (e.g., *Salmonella*) produces pale yellow colonies.
**pH indicator: Methylene Blue and Eosin dyes**

- **Eosin Methylene Blue (EMB) Agar:** EMB agar is used for the isolation and differentiation of lactose fermenting and non-lactose fermenting enteric bacilli.
- **Coliform bacteria** - purplish black colonies; *E.coli* with typical metallic sheen
- **Non Coliform Bacteria:** colorless colonies.
pH indicator: Malachite Green

- **Lowenstein Jensen Medium**: Lowenstein Jensen medium is used to isolate *Mycobacterium tuberculosis*. 
Common Bacterial Culture Media
Nutrient Agar

Composition of Nutrient Agar

- Beef Extract ..........3.0 g
- Peptone ................5.0 g
- Agar ....................15.0 g
- Distilled water ....1000 ml
- Final pH 6.8 +/- 0.2.

Composition of Nutrient Broth:

- Nutrient broth contains same ingredients except agar.
Blood Agar

• Blood agar is an enriched, indicator and differential bacterial growth medium.
• Blood agar is a type of growth medium that encourages the growth of bacteria, such as streptococci, that otherwise wouldn’t grow.
• Blood agar consists of a base containing a protein source (e.g. Tryptone), soybean protein digest, sodium chloride (NaCl), agar and 5% defibrinated sheep blood.
Blood Agar and Haemolysis

• Certain bacterial species produce extracellular enzymes that lyse red blood cells in the blood agar (haemolysis).

• These haemolysins (extotoxin) radially diffuses outwards from the colony (or colonies) causing complete or partial destruction of the red cells (RBC) in the medium and complete denaturation of haemoglobin within the cells to colourless products.

• Four types of haemolysis are produced in sheep blood agar by streptococci namely; alpha haemolysis, beta haemolysis, gamma haemolysis and alpha prime or wide zone alpha haemolysis.

• Haemolysis is best observed by examining colonies grown under anaerobic conditions or inspecting sub-surface colonies.
Blood Agar and α-Haemolysis

**Alpha haemolysis:**

- Partial lysis of the RBC to produce a greenish-grey or brownish discoloration around the colony.
- α-haemolysis is due to the reduction of RBC’s haemoglobin to methaemoglobin in the medium surrounding the colony.
- Many of the alpha haemolytic streptococci are part of the normal body flora, except *Streptococcus pneumoniae*
Blood Agar and β-Haemolysis

Beta haemolysis:

• Complete lysis of red blood cells causing a clearing of blood from the medium under and surrounding the colonies

• e.g. Group A beta-hemolytic streptococci—Streptococcus pyogenes and Group B, beta hemolytic streptococci—Streptococcus agalactiae.

• For group A streptococci maximal activity of both the hemolysins - oxygen labile (SLO) and oxygen stable (SLS) hemolysins is observed only in anaerobic conditions.
Blood Agar and $\gamma/\alpha'$-Haemolysis

**Gamma or non-hemolysis:**

- No hemolysis of RBC. No change of the medium under and surrounding the colonies.

**Alpha prime or wide zone alpha hemolysis:**

- A small zone of intact erythrocytes immediately adjacent to bacterial colony, with a zone of complete red-cell hemolysis surrounding the zone of intact erythrocytes.
- This type of hemolysis may be confused with $\beta$-hemolysis.
Blood Agar and Target Haemolysis

**Target hemolysis**

*Clostridium perfringens* is readily identified in the laboratory by its characteristic “double zone” hemolysis also known has target hemolysis.
Chocolate Agar

- Chocolate Agar is the lysed blood agar.
- The name is derived from the fact that RBC lysis gives the medium a chocolate-brown color.
- Chocolate agar is used for the isolation of fastidious organisms, such as *Haemophilus* and *Niessera*.
- The composition of chocolate agar and the Blood Agar is the same and the only difference is while preparing chocolate agar, the red blood cells are lysed.
- The lysis of RBC during the heating process releases intracellular coenzyme **NAD or Factor V** into the agar for utilization by fastidious bacteria.
- **Hemin of Factor X** is available from non-hemolyzed as well as hemolyzed blood cells.
Robertson’s Cooked Meat (RCM) Medium

- Robertson’s Cooked Meat (RCM) medium is used for the cultivation of aerobic, microaerophilic, and anaerobic microorganisms, especially *Clostridium species*.
- It is also known as Cooked Meat Broth (CMB) as it contains pieces of fat-free minced cooked meat of ox heart and nutrient broth.
- It supports the growth of both spore forming and non-spore forming obligate anaerobes and also differentiates between putrefactive and saccharolytic species.
- Oxygen in culture media can be reduced by various agents such as glucose, thioglycollate, cooked meat pieces, cysteine and ascorbic acid.
Robertson’s Cooked Meat (RCM) Medium

**Principle**

- Before inoculation, RCM/CMB medium is boiled to make it oxygen-free.
- After inoculation, it is covered with a layer of sterile liquid paraffin oil to prevent the entry of oxygen in the medium.
- The ingredients present in the medium help to maintain the anaerobic (reduced) environment.
- Unsaturated fatty acids present in meat utilize oxygen for autooxidation, this reaction is catalyzed by haematin in the meat.
- Glutathione and cysteine (both are reducing agents) present in meat also utilize oxygen.
- Sulphhydryl compounds (present in cysteine) also contribute to a reduced oxidation-reduction potential.
MacConkey Agar

- **MacConkey agar** is a selective, indicator and differential culture media commonly used for the isolation of **enteric Gram-negative bacteria**.
- Crystal violet and bile salts are incorporated in MacConkey agar to prevent the growth of Gram-positive bacteria and fastidious Gram-negative bacteria, such as *Neisseria* and *Pasteurella*.
- Gram-negative enteric bacteria can tolerate bile salts because of their bile-resistant outer membrane.
- MCA also helps to differentiate lactose fermenting gram-negative rods from non-lactose fermenting gram-negative rods.
- It is primarily used for the detection and isolation of members of family *Enterobacteriaceae* and *Pseudomonas spp.*
### MacConkey Agar

<table>
<thead>
<tr>
<th>Ingredient</th>
<th>MacConkey Agar (g/L)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Peptone</td>
<td>17 g</td>
</tr>
<tr>
<td>Polypeptone</td>
<td>3 g</td>
</tr>
<tr>
<td>Lactose</td>
<td>10 g</td>
</tr>
<tr>
<td>Bile salts</td>
<td>1.5 g</td>
</tr>
<tr>
<td>Sodium chloride</td>
<td>5 g</td>
</tr>
<tr>
<td>Agar</td>
<td>13.5 g</td>
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<tr>
<td>Neutral red</td>
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<tr>
<td>Crystal violet</td>
<td>0.001 g</td>
</tr>
<tr>
<td>Distilled water</td>
<td>1 L</td>
</tr>
<tr>
<td>Final pH</td>
<td>7.1</td>
</tr>
</tbody>
</table>
MacConkey Agar

**Principle behind differential capability of MacConkey agar**

- Gram-negative enteric bacteria that grow on MacConkey agar are differentiated by their ability to ferment lactose. If the lactose is fermented by the bacteria, the production of the acid drops the pH of the media. The drop in pH is indicated by the change of neutral red indicator to pink (Neutral red appears pink at pH’s below 6.8).
- Strongly lactose fermenting bacteria produce sufficient acid which causes precipitation of the bile salts around the growth. It appears as a pink halo surrounding individual colonies.
- Pink halo is not seen around the colonies of weaker lactose fermenting bacteria.
- Gram-negative bacteria that grow on MacConkey agar but do not ferment lactose appear colorless on the medium and the agar surrounding the bacteria remains relatively transparent.
MacConkey Agar

**Lactose Fermenter Organisms (CEEK)**:
- *Citrobacter spp.* - colonies are light pink after 48 hours.
- *Klebsiella spp.* - colonies are mucoid
- *Escherichia coli* - flat, dry, pink colonies with a surrounding darker pink area of precipitated bile salts.
- *Serratia spp.* - late lactose fermenter

**Non-Lactose Fermenter (NLF) Organisms (ShYPS)**:
- *Proteus spp.*: *may swarm depending on the amount* of agar in the medium; characteristic foul smell
- *Shigella spp.*
- *Yersinia spp.*: *may be colorless to peach.*
- *Salmonella spp.*
Selenite broth

• Selenite broth is used as an enrichment medium for the isolation of *Salmonella* and *Shigella* from stool, urine, water and food products.

• Selenite is inhibitory to coliforms and certain other microbial species found in faecal specimens. It is suggested that it reacts with sulphur and sulphydral groups of in critical cell components of microorganisms.

• Coliforms, fecal streptococci, and Gram-positive organisms are inhibited by sodium selenite.

• After the inoculation of the sample, selenite broth is incubated for 12-18 hours at 35°C - 37°C and then subculture is done in selective agar
Deoxycholate Citrate Agar

- Deoxycholate Citrate Agar (DCA) is a selective and differential medium, recommended for the isolation of enteric pathogens particularly *Salmonella* and *Shigella* species.
- DCA is selective for enteric pathogens owing to increased concentrations of both citrate and deoxycholate salts.
- Sodium deoxycholate at pH 7.3 to 7.5 is inhibitory for gram positive bacteria.
- Citrate salts, in the concentration included in the formulation, are inhibitory to Gram positive bacteria and most other normal microflora present in intestine.
Deoxycholate Citrate Agar

• Lactose helps in differentiating enteric bacilli (lactose fermenters produce red/pink colonies while lactose nonfermenters produce colourless colonies – neutral red is pH indicator).

• Lactose-fermenting colonies may have a turbid zone of precipitation around them caused by the precipitation of deoxycholate in the acidic environment.

• The reduction of ferric ammonium citrate to iron sulfide is indicated by the formation of black iron sulfide. If the bacteria produce H₂S, the colonies will have black centers.

• Coliform bacteria and gram-positive bacteria are inhibited or greatly suppressed due to sodium deoxycholate, sodium citrate and ferric ammonium citrate.

*Salmonella* and *Shigella* species do not ferment lactose but *Salmonella* may produce H₂S, forming colorless colonies with or without black centers.
EMB Agar

• Eosin Methylene Blue (EMB) agar is both selective and differential culture medium.
• It is a selective culture medium for Gram-negative bacteria being inhibitory to Gram-positive bacteria.
• EMB media assists in visual distinction of *Escherichia coli* from other non-pathogenic lactose-fermenting enteric gram-negative rods, and the *Salmonella* and *Shigella* genera.
• Differentiation between these gram-negative bacilli is based on the colony color.
  – Colored colonies in EMB agar: Lactose fermenter
  – Colorless colonies in EMB agar: Non-lactose fermenter
• The aniline dyes (eosin and methylene blue) inhibit Gram positive and fastidious Gram-negative bacteria. They combine to form a precipitate at acid pH, thus also serving as indicators of acid production.
EMB Agar

- Gram-negative bacteria that ferment lactose produce acid which turns the colonies to dark purple as the acid acts upon the dyes.
- Rapid fermentation of lactose & production of strong acids by bacteria such as *E. coli* causes a rapid reduction in the pH of the EMB agar, which is the critical factor in the formation of the green metallic sheen observed.
- Other lactose fermenters produce larger, mucoid colonies, often purple only in their center.
- Lactose non-fermenters are either colorless or light lavender.
Bismuth Sulfite Agar

- Bismuth Sulfite Agar (BS) is a selective as well as differential medium for isolation and presumptive identification of *Salmonella* spp especially *Salmonella Typhi*.

- Bismuth Sulfite Agar is selective due to the presence of inhibitors, and is differential on the basis of hydrogen sulfide (H₂S) production.

- The presence of Bismuth sulfite indicator and brilliant green inhibits the growth of Gram-positive bacteria and members of the coliform group, while allowing *Salmonella* to grow luxuriantly.

- Ferrous sulfate is an indicator for H₂S production, which occurs when the H₂S produced by *Salmonella* reacts with the iron salt. This reaction causes a black or green metallic colony and brown or black precipitate whilst the reduction of bismuth ions to metallic bismuth produces a metallic luster around of the colonies.
Mannitol Salt Agar

• Mannitol salt agar (MSA) is a selective, differential and indicator medium which is used to isolate and identify *Staphylococcus aureus* from the clinical specimen.

• Incorporation of 7.5% sodium chloride in the medium helps to select only those bacteria which can tolerate high salt concentrations such as *Staphylococcus* spp. This concentration inhibits the growth of most other gram-positive and gram-negative bacteria.

• Pathogenic staphylococci, i.e. *Staphylococcus aureus* is able to ferment mannitol others (coagulase-negative *Staphylococcus*) are not. *S. aureus* ferments mannitol and changes the pH of the medium to acidic. As MSA contains phenol red as a pH indicator, at pH levels below 6.9, the medium is a yellow color.

• However, if coagulase-negative staphylococcus (CONS) grows, they cant ferment mannitol, so the color of the media around the bacterial colony does not change to yellow, it appears pink.