# Microbial growth and metabolism

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# Microbial growth and microbial growth requirement

### Microbial growth

Microbial growth is the increase in number of cells, not cell size.

### **Growth in colonies**

- A pure culture contains only one species or strain.
- A colony is a population of cells arising from a single cell or spore or from a group of attached cells.
- a unit used to estimate the number of viable microbial cells is colony forming unit (CFU)

### **Bacterial Division**

The normal reproductive method of bacteria is **binary fission**, in which a single cell divides into two identical cells

#### Plasma membra Cell elongates and **Bacterial Division** DNA DNA is replicated The normal Cell wall and reproductive plasma membrane begin to grow method of bacteria inward is binary fission, in 💿 Cross-wall forms completely around which a single cell divided DNA divides into two identical cells. separate (a) A diagram of the sequence

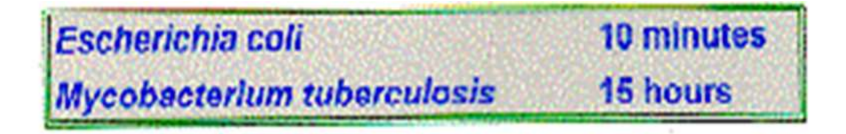
of cell division.

### **Growth rate**

The number of generation per hour.

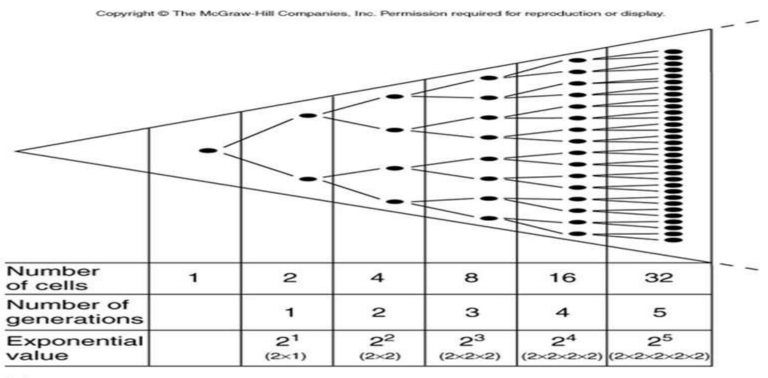
### **Generation Time**

The **time required for** a cell to divide or a **population to double** is known as the generation time.



## The generation time depend upon:

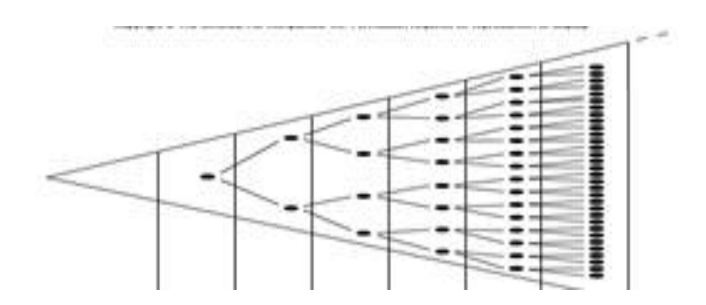
- 1- The nutrient in the medium.
- 2- Physical condition (pH, temp. etc.)



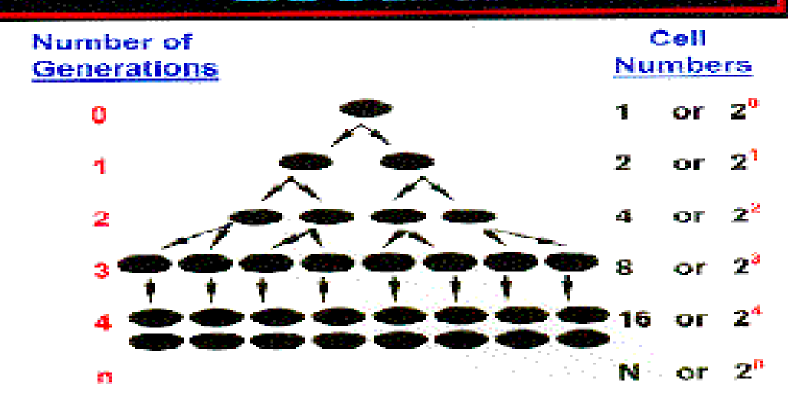
(a)

Q/- what is the generation number of bacteria when the number of cell is 8

- If the generation number of bacteria was 4 what is the Exponential value?



## Wathematical Expression of Growth



## **Bacterial growth curve**

- All microorganisms undergo similar growth patterns Each growth curve has 4 phases:
- 1- Lag phase
- 2- Log phase
- 3- Stationary phase
- 4- Death or decline phase

Between each phases there is a transitional phase is represent the time require by all the cell before get to inter the new phase.

### 1- Lag phase:

- The number of the population remains constant.
- Microorganism start to adapted itself to the environment.
- The lag phase is generally longer if the cells are taken from an old or refrigerated culture.
- If the cells are taken from young, new growing culture (microbial population) and inoculated to a fresh medium the lag phase may be short or even absent

## 2- Log phase (logarithmic phase or exponential phase):

- The bacteria multiply at the fastest rate possible under the conditions provided.
- Most research is performed on cells during log phase
- This phase is called log phase because the logarithm of the bacterial mass increases linearly with time,
- and exponential growth phase because the number of cells increases as an exponential function of 2n (i.e.  $2^1$ ,  $2^2$ ,  $2^3$ ,  $2^4$ ,  $2^5$  and so on).

<sup>-</sup> The portion of the growth curve where rapid growth of bacteria is observed is known as Logarithmic phase.

### 3- Stationary phase:

- Growth levels off.
- Cells per volume does not increase or decrease.
- Growth rate = Death rate.

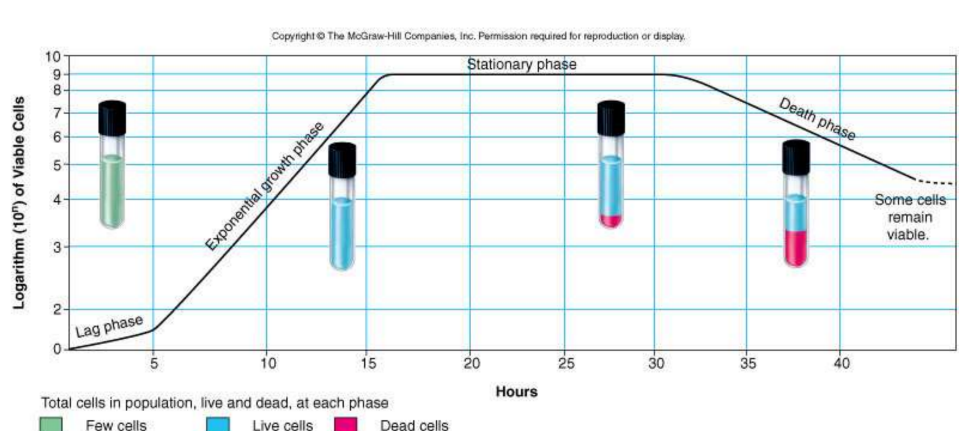
### Due to

- Depletion of nutrients
- Increase in waste products

### 4- Death phase:

- The number of deaths exceeds the number of new cells formed
- Cells per volume decreases
- Due to
- Very low concentrations of nutrients
- Very high concentrations of waste products

# Standard Growth Curve



## **Nutritional Requirements for Microorganisms:**

-Water (preservation of a microbial culture from drying)

## -Energy:

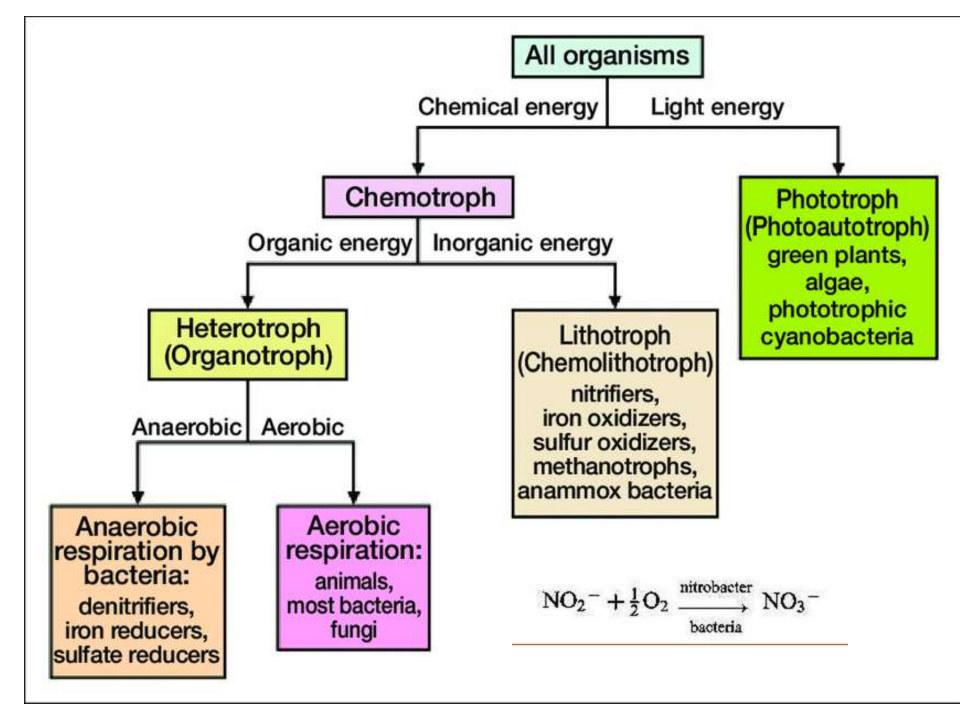
Phototroph- Energy from sun light (cyanobacteria)

Chemotroph- Energy from chemicals (Nitrobacter, Sulphur bacteria)

Heterotroph- from carbohydrate and other organic material

## -Carbon:

- Autotrophs- carbon from carbon dioxide- (inorganic carbon)
- Heterotroph- carbon from organic carbon
- e.g. Carbohydrates, lipids, protein.



- Essential elements
- Hydrogen (H), Sulfur (S), Oxygen (O), Phosphorous (P)
- Nitrogen (N)
- commonly supplied as ammonia (NH4)
- some microbes fix atmospheric nitrogen (N2)

### - Trace elements

- Required in small amounts
- Copper (Cu), Zinc (Zn), Selenium (Se)

### **Environmental requirements for growth**

 Temperature, pH, Oxygen, Carbon dioxide, Osmotic pressure, Hydrostatic pressure

### - Temperature

- Psychrophiles less than 20 °C
- Mesophiles 20 45 °C
- Thermophiles -45 80 °C
- Extreme thermophiles more than 85 °C
- Psychrotroph- 0-40 (20) °C

- pH (- log [ H+ ])
   Low pH = acid, High pH = basic or alkaline
- Acidophiles below pH 5.5 (Acidobacteriota, Mucor)
- Neutrophiles at pH 6 8
- Alkalophiles above pH 8 (Thiohalospira alkaliphila)

### - Molecular oxygen

Microbe vary greatly in sensitivity to oxygen.

- Aerobes microbes which require oxygen.
- <u>Facultative anaerobes</u> microbes which can grow in presence or absence of oxygen.
- Obligate Anaerobes which do not utilize oxygen and are killed by oxygen.
- <u>Aerotolerant anaerobe</u> is an organism that tolerates the presence of oxygen but does not require it for growth.
- Microaerophiles required 3 15 % oxygen.

# Chemical Requirements for Growth: Oxygen

# O<sub>2</sub> requirements vary greatly

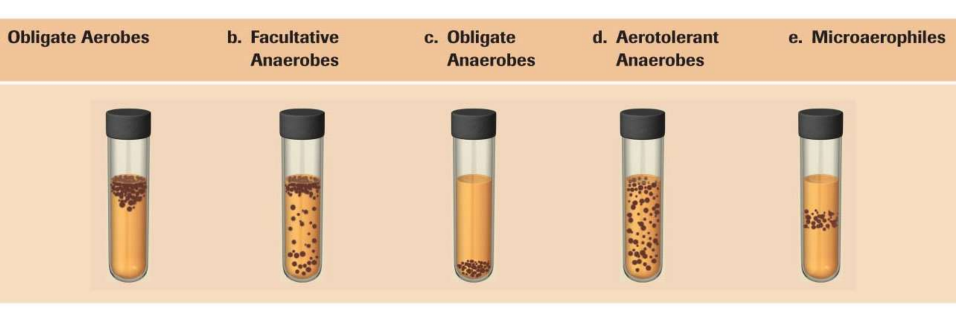


Table 6.1: The Effects of Oxygen on the Growth of Various Types of Bacteria

### - Carbon dioxide

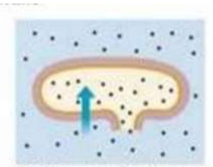
- Capnophiles : 3 10 % carbon dioxide
- Many microaerophiles are also capnophiles



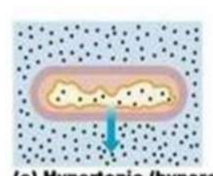
### **Osmotic Pressure**

High osmotic pressure (hypertonic) removes water causing plasmolysis – inhibits growth i.e. salt as preservative

Low osmotic pressures (hypotonic) cause water to enter and can cause lysis



(d) Hypotonic (hypoosmotic) solution—water moves into the cell and may cause the cell to burst if the wall is weak or damaged (osmotic lysis)



(e) Hypertonic (hyperosmotic) solution water moves out of the cell, causing its cytoplasm to shrink (plasmolysis)

### Quantitative methods for measuring growth of bacteria:

The growth of bacteria can be determined by numerous techniques based on one or more of the following types of measurement:

#### 1- Cell count

- a- microscopy or by using electronic particle counter.
- b- colony count or number (plate count method, MPN)

#### 2- Cell mass

- a- weighting (dry weight)
- b- Measurement of cell nitrogen
- c- Indirectly by turbidity with culture, directly without culture
- 3- Cell activity- indirectly by relating the degree of biochemical activity to the size of population Such as measurement of utilizing O2

### **Direct methods:**

With direct methods we count individual cells or colonies that are assumed to have apart or arise in through the division of a single cell.

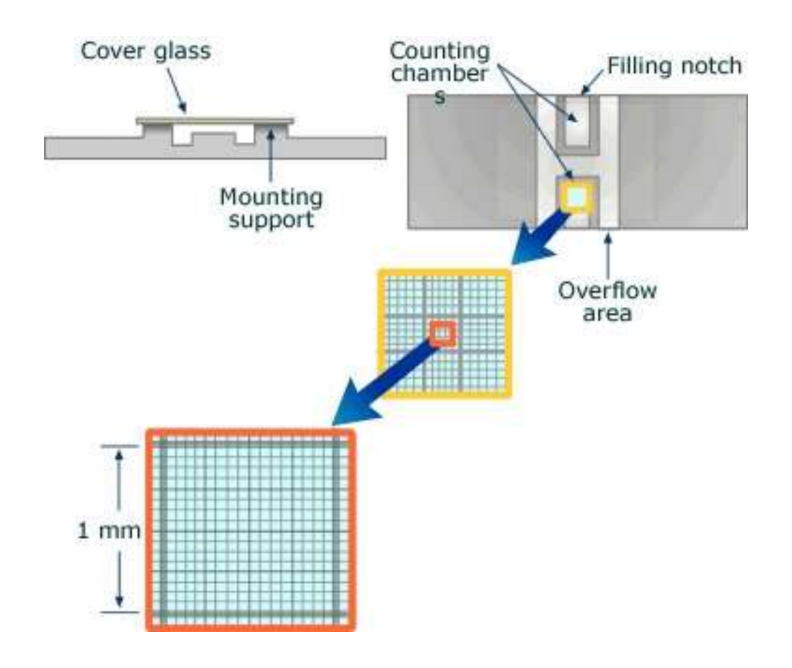
## 1- Counting Chamber (Hemocytometer):

The hemocytometer is a specialized microscope slide used to count cells.

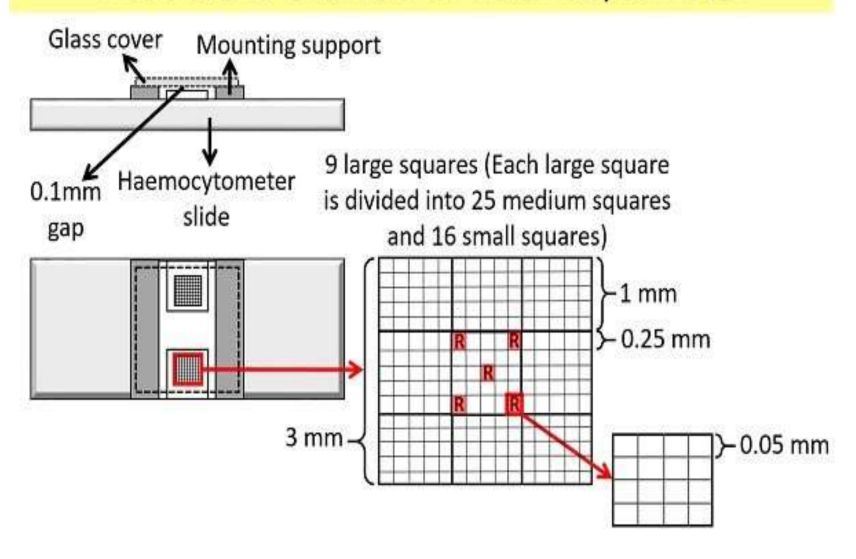
The center portion of the slide has etched grids (H) with precisely spaced lines.



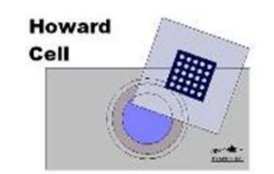


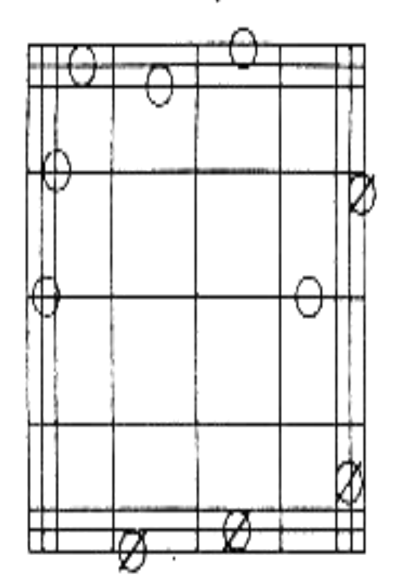


# Neubauer's Chamber or Haemocytometer



# CORNER SQUARE (ENLARGEMENT)





Count cells on top and left touching middle line (O). Do not count cells touching middle line at bottom and right (Ø).

### Steps of Counting of cell using hemocytometer

- 1- percentage of viable cell= Total viable cell/ total cell \*100
- 2-Average of cell per square = total viable/ average
- 3- Dilution factor =Final volume / volume of cell
- 4- Concentration (viable cells/ml)=Average of cell\* dilution factor\*

### 10000 (104)

Suppose you have 200 ml of sample diluted with 200 ml stain. you calculated the number of viable cell as 36 and the number of a viable cell (dead cell) was 6 and the average of cell per square was 6.

#### **Answer the following questions**

- 1. Calculate Percentage of viable cell
- 2. Find the concentration of viable cell/ml

#### **Answer**

200+200= 400 total volume

D. f= final volume / volume .of cell( 400/200=2)

Total cell= 36+6=42

Percentage of viable cell= total viable cell/ total cell\*100

concentration of viable cell/ml= Av. Of cell \* dilution factor\*10<sup>4</sup>

$$=6*2*10^4=120,000$$

### 2- Coulter Counter:

electronic counting (this machine detects the difference in current as individual microorganisms pass through a small orifice).

It is Very fast, easy to use but;

Very EXPENSIVE.



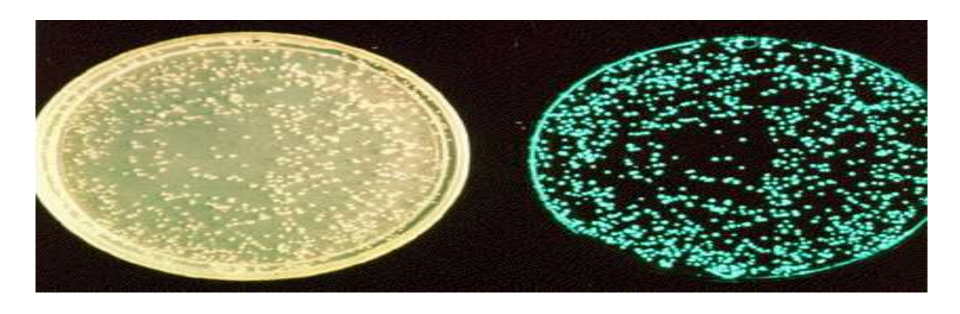
# 3- Viable count assays (Colony Counting):

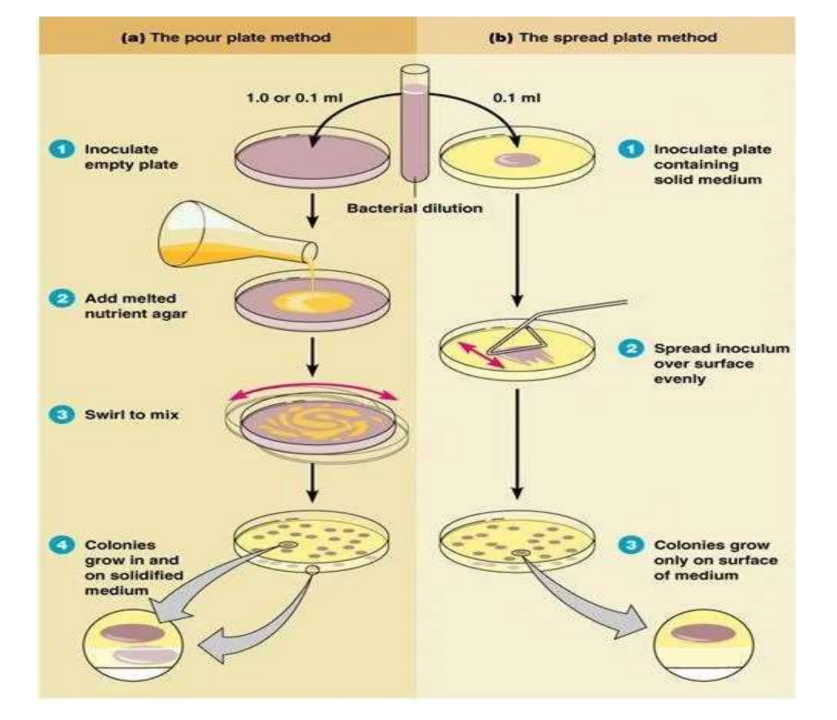
Colony counting after plating dilutions of the sample onto growth medium.

Standard plate counts using spread and pour plate techniques (cfu for "colony forming unit").

### two viable count assays:

1 - Spread plates 2- Pour plates



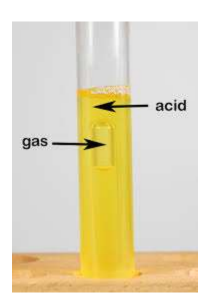


### **Indirect Method:**

Indirect methods often rely on the results of metabolic tests or other growth characteristics. And it's to:

- Measurement of metabolic activity.-Gas or Acid Production.







- Turbidity using a spectrophotometer.
spectrophotometry, using a spectrophotometer.

## These Indirect counts depend on:

- The effects of the organisms to estimate their numbers.
- As organisms grow they make the nutrient broth turbid.
- This turbidity can be measured with a colorimeter

