

# تفريغ تعقيم

المحاضرة الأولى

محاضرة:

عَرُوب طرابشه

الصيدلانية:



لجان الرُّفَعات



# Counting, Detecting and identifying microorganisms

Chapter 15&18

# Key Facts

الصحة الحيوي

▶ **Bioburden**: the number and type of microorganisms (MO) present (in) or (on) a pharmaceutical raw material, medicine, or medical device

▶ **Pharmacopeias describe procedures for**: → *هيا إجراءات لـ non-sterile drug*

- Procedures for counting microorganisms → *إجراءات عد الكائنات الحية الدقيقة*

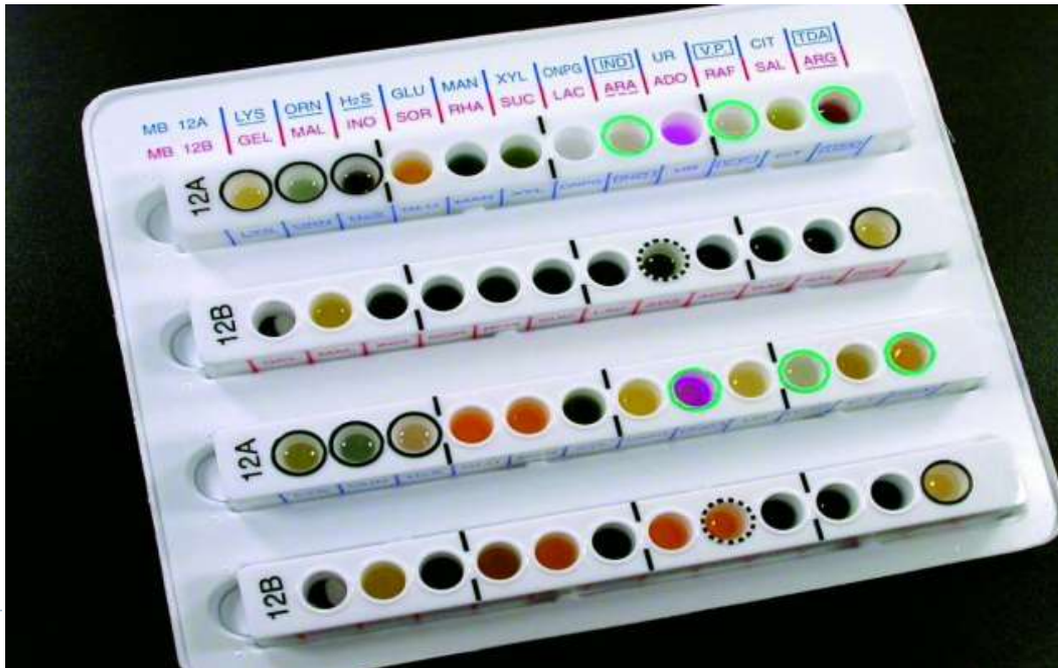
- Procedures for confirming the absence of named **objectionable MO** → *وجودهم يجب مشاكل صحية أو بأثرها على product stability*

- Specifications for the max permitted levels of bacteria and fungi for the different categories of nonsterile medicine

- Requirements for the absence of one or more of the major objectionable MO: E.coli, Salmonella species, Staph aureus, Pseudomonas aeruginosa, and Candida albicans

# Key Facts

- ▶ Commercially available test kits and automated instruments are available for identifying different categories of bacteria and yeasts



# Key Facts

▶ **Objectionable:** → غير مرغوب فيه

- Because it is a pathogen so its presence in a medicine may cause an infection (E.coli & Staph aureus are pathogens → health hazards) → مخاطر صحية

- Its presence may be indicative of poor-quality raw materials or poor manufacturing procedures يدل على وجوده على مواد خام برديته او اجراءات التصنيع السيئه

افتبار عدم وجود

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**Gelatin:** test for absence of E. coli and Salmonella

لصنع كابتسول  
او كابتسول

# Bioburden

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▶ Determination of bioburden quantitatively: to count the organisms present → Total Viable Count (TVC)

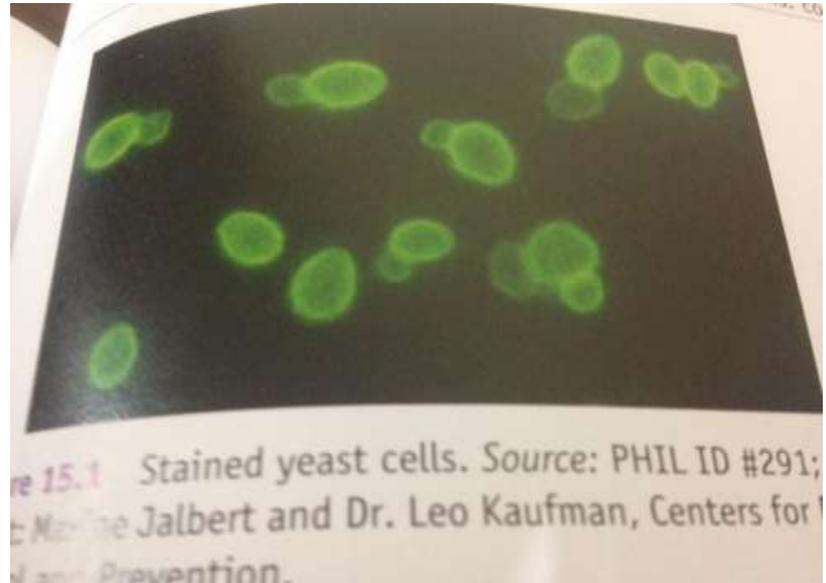
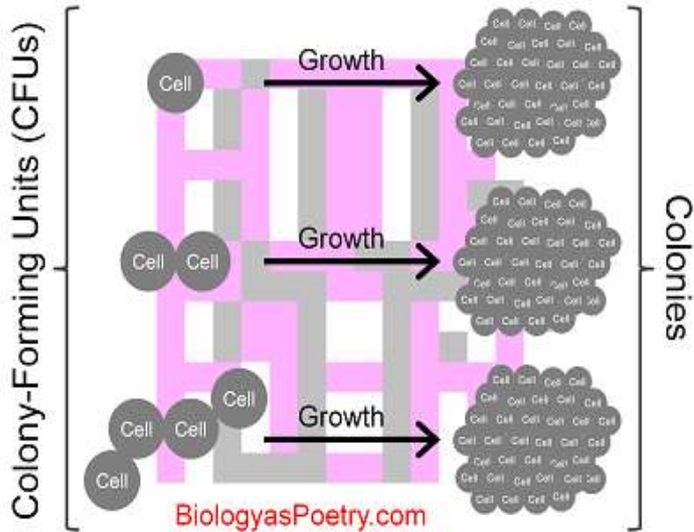
▶ Bioburden test → the most common microbiological test undertaken in the pharmaceutical industry:

- Raw material (including water)
- Finished products
- Various stages during manufacturing process

## Traditional counting methods:

- ▶ Place a sample of the material to be tested onto or into gelled culture medium in a petri dish (plate) and counting the visible colonies that arise after incubation
- ▶ A single colony may develop from an individual cell or from a group of cells attached or clumped together → count is expressed as colony forming unit (CFU) and not cells per ml or gram  
*وحدة تشكيل المستعمرة*
- ▶ USP states that petri dishes containing between 25-250 colonies produce reliable bioburden results
- ▶ High Bioburden materials:
  - Mined minerals (talc, kaolin, bentonite)
  - Vegetable origin (starches, gums, thickening agents)
  - Animal origin (gelatin)

# Traditional counting methods

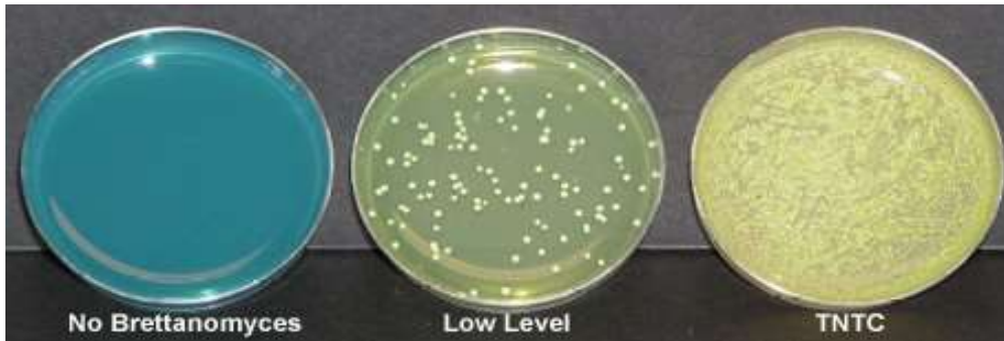


# Traditional counting methods

- ▶ Dilution is necessary for material with a high bioburden, why?

to achieve a countable number of colonies on the plate

Perform a series of dilution (tenfold increments)



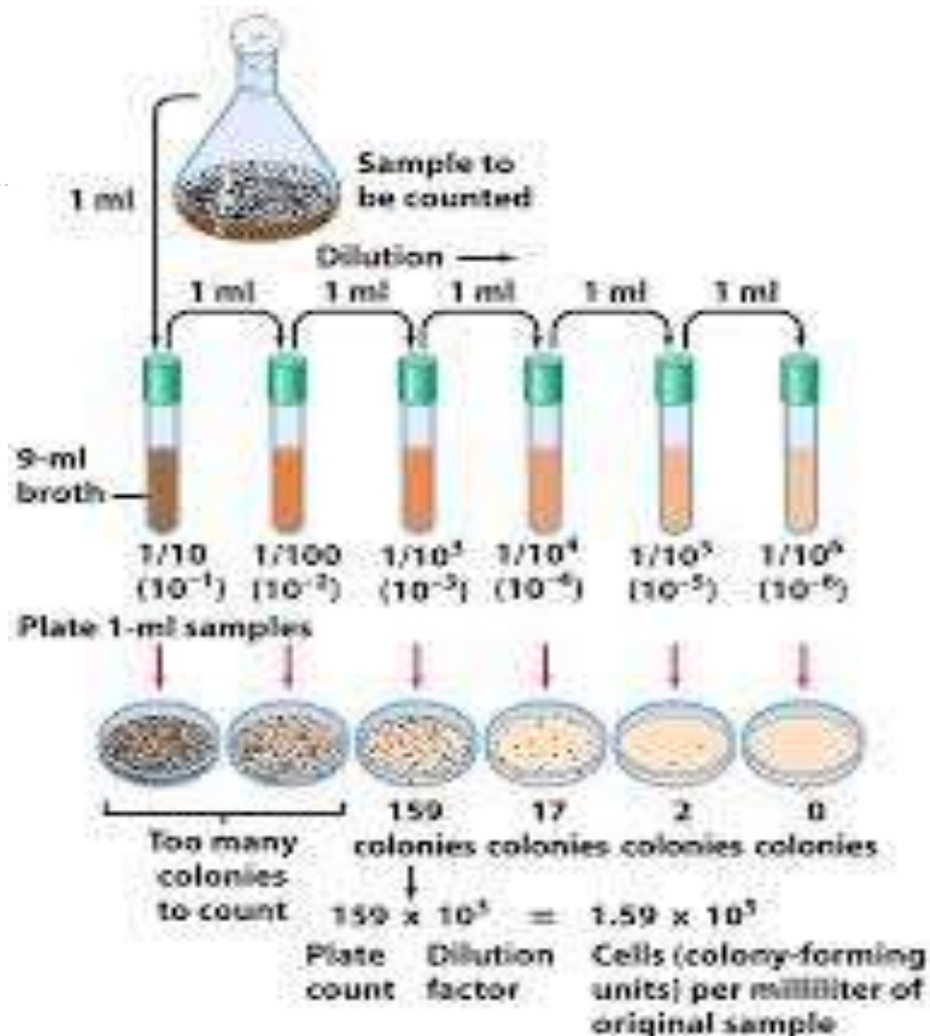


Figure 9-11 Brock Biology of Microorganisms 110e  
 © 2006 Pearson Prentice Hall, Inc.

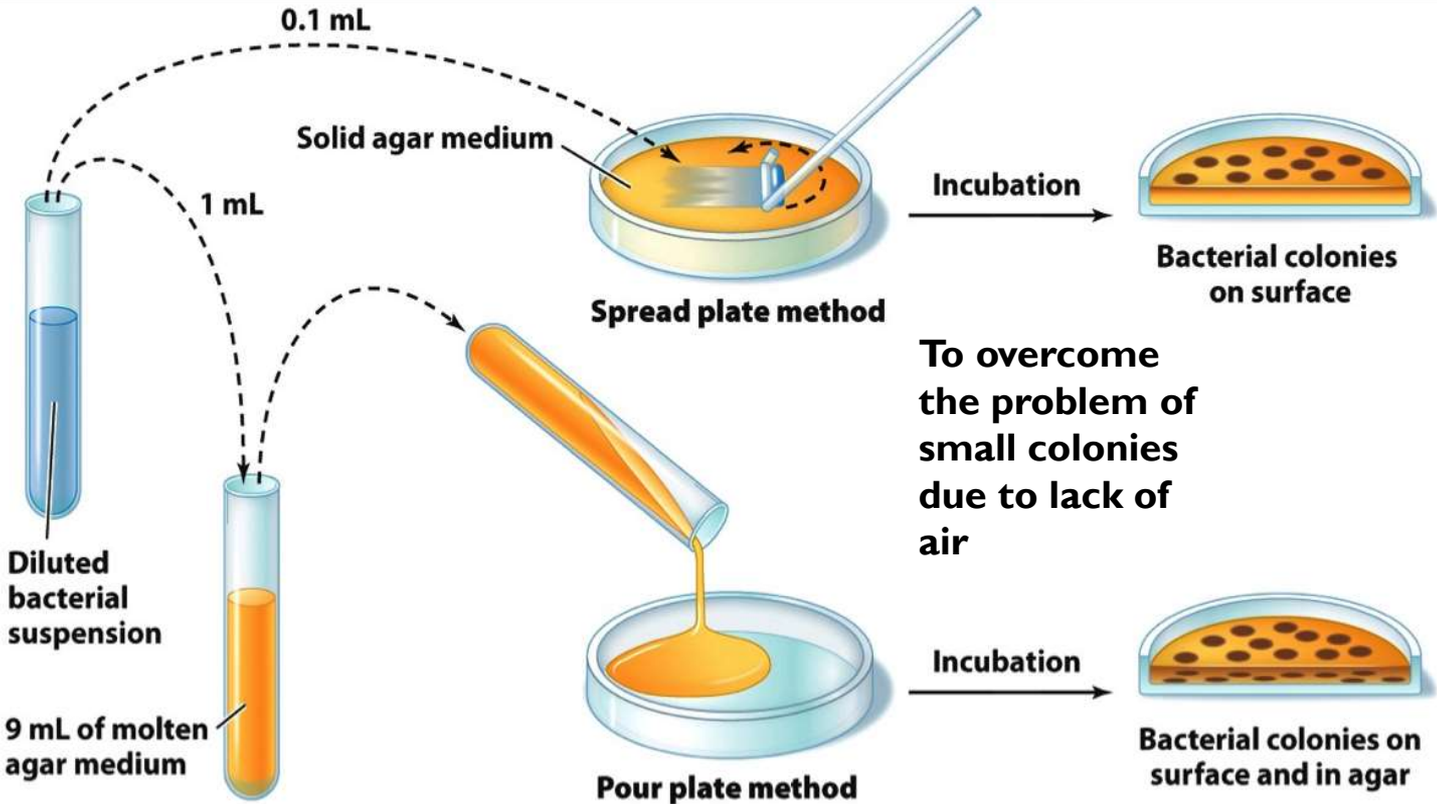
# Traditional counting methods:

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1. Pour plates (most popular)
2. Spread plates
3. Membrane filter method
4. Most probable number method



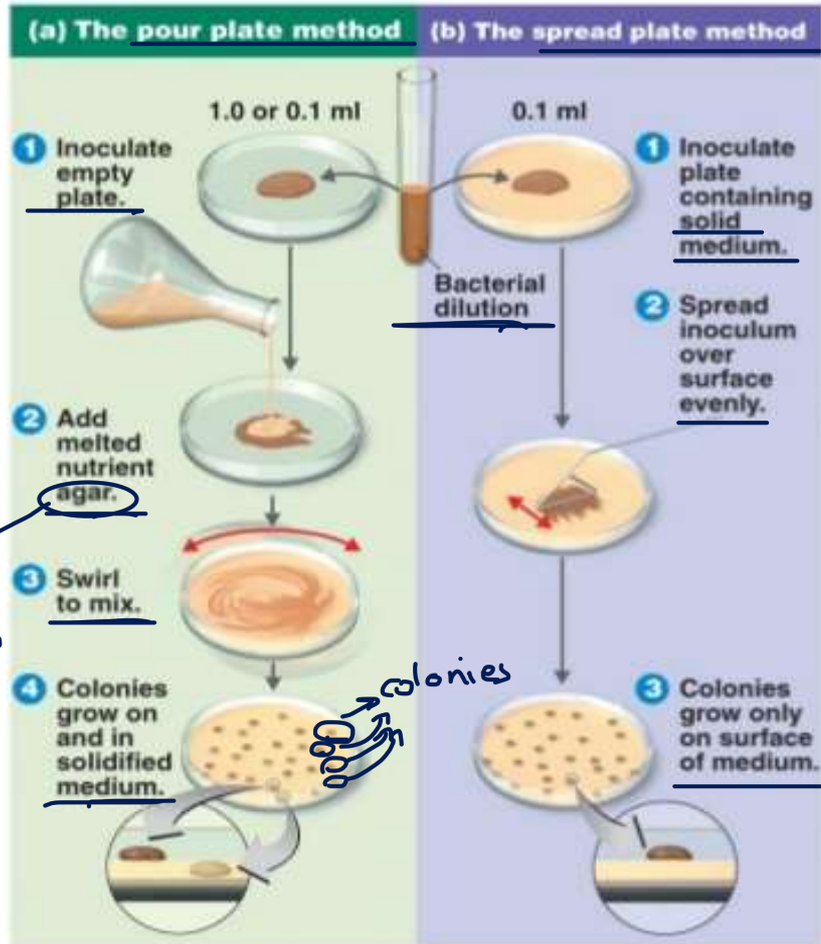
Sterile glass or plastic “hockey-stick” spreader

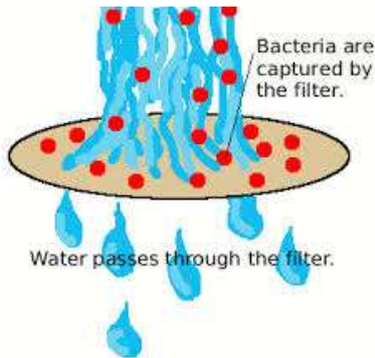
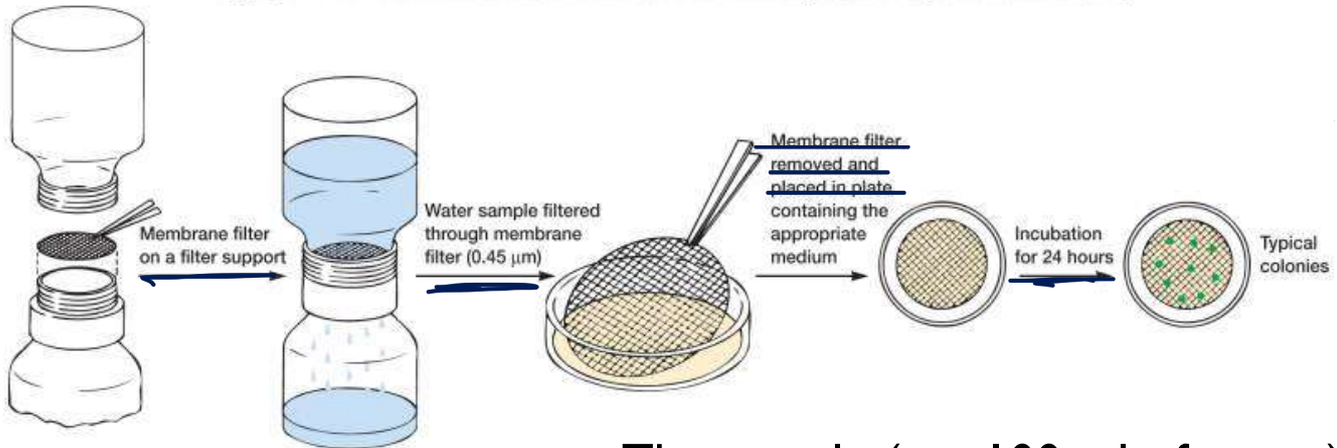


- Plating serial dilutions of the specimen

- Pour plate method
- Spread plate method

ليش بنسكده ١؟  
 تحتها بعد ما نزره البليكر يا  
 تكبير بغيره ونقدر نعددها





- The sample (e.g. 100 ml of water) is passed under vacuum through a sterile filter membrane with pore size that is small to retain all contaminating MO on its surface (usually 0.45  $\mu\text{m}$ )

# Membrane filter method

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▶ Advantages:

- For low bioburden material like Purified water EP → required to have no more than 100 CFU/ml
- Can be used to determine concentration of organisms in oils or ointments (after dispersing it in oil)
- The best method if the sample is likely to contain antimicrobial chemicals (preservatives)



Antimicrobial agent	inactivator
① <u>Quaternary ammonium compounds</u> , parabens and chlorhexidine	<u>Lecithin with or without Polysorbate (Tween) 80</u>
<u>Thimerosal</u> and other <u>mercurials</u>	<u>Sodium thioglycolate</u>
<u>Beta-Lactam antibiotics</u>	<u>Beta-lactamase</u>
<u>Phenols</u> , <u>alcohols</u> and <u>weak acid preservatives</u>	<u>Dilution alone</u>

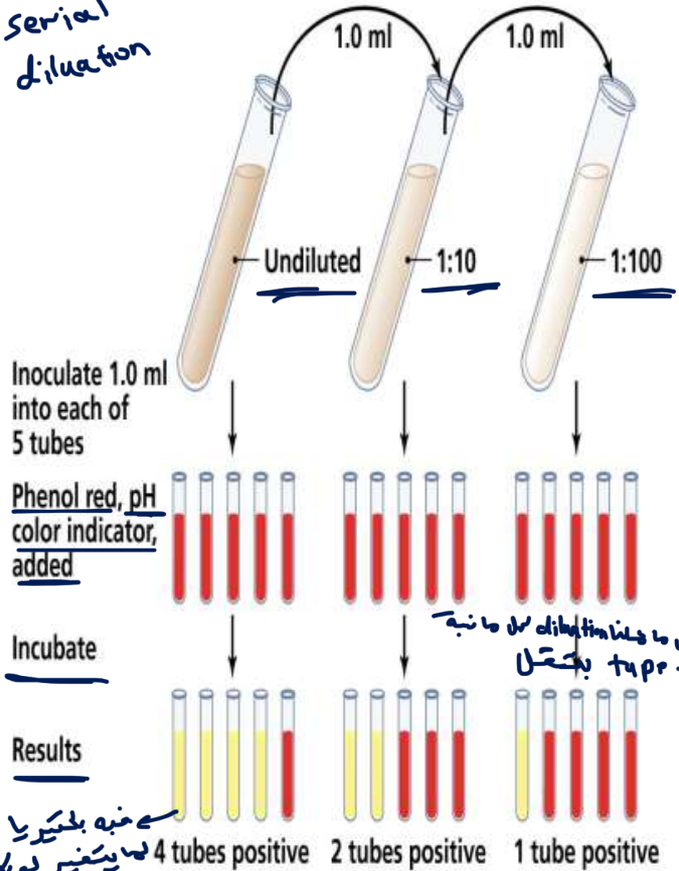
يعني ما يقتل ، انكسر مع حرارة





MPN is used when the sample contains insoluble material that would interfere with plate counts as in herbal products

Serial dilution



# INDIRECT MEASUREMENTS: MPN

- Multiple Tube Fermentation Test as measured in MPN or Most probable Number
- Count positive tubes and compare to statistical MPN table.

Combination of Positives	MPN Index/ 100 ml	95% Confidence Limits	
		Lower	Upper
4-2-0	22	9	56
4-2-1	26	12	65
4-3-0	27	12	67
4-3-1	33	15	77
4-4-0	34	16	80
5-0-0	23	9	86
5-0-1	30	10	110
5-0-2	40	20	140
5-1-0	30	10	120
5-1-1	50	20	150
5-1-2	60	30	180
5-2-0	50	20	170
5-2-1	70	30	210
5-2-2	90	40	250
5-3-0	80	30	250
5-3-1	110	40	300
5-3-2	140	60	360

# Calculation of concentration of MOs in a sample

**Table 15.3** Specimen results from a viable count.

Dilution	Dilution factor	Colony count 1	Colony count 2	Colony count 3
A	$10^1$	TNTC <sup>a</sup>	TNTC	TNTC
B	$10^2$	TNTC	TNTC	TNTC
C	$10^3$	453	521	419
D	$10^4$	85	79	81
E	$10^5$	7	6	8
F	$10^6$	0	1	0

<sup>a</sup>TNTC = too numerous to count

**Viable count of original sample =**

$$\frac{\text{Mean colony count}}{\text{Volume of dilution used}} \times \text{dilution factor}$$

# Detection of objectionable MOs

▶ Non sterile dosage forms may contain some MOs

▶ The quality of non sterile products is controlled by pharmacopeia in two ways: يتم التحكم بجودة المنتجات بواسطة دستور الأدوية بطريقتين:-

1. Limit on total number or concentration of MO that may be present الحد من العدد الإجمالي أو تركيز MO، الذي يظل موجود

2. Particular objectionable organisms must be absent in a specified weight of material: يجب أن تكون بعض الكائنات الحية خائفة في وزن محدد من المواد

e.g. EP quality gelatin → Salmonella should be absent in a 10g sample and E.coli absent in 1 g sample

# Detection of objectionable MOs

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1. **Dissolution or dispersal** of the sample in a suitable liquid culture medium and , where necessary, inactivation of any substances that might inhibit the growth of the organism under test
2. **Enrichment**: increasing the relative concentration of the test organism **by growing in a liquid medium** that inhibit other contaminants but **allows free multiplication of the organism of interest**
3. **Streaking** liquid cultures from step 2 onto **selective agar media** that usually permit easy **recognition** of any colonies of the test organism that might arise
4. **The use of specific biochemical or immunological confirmatory tests (test kits)**



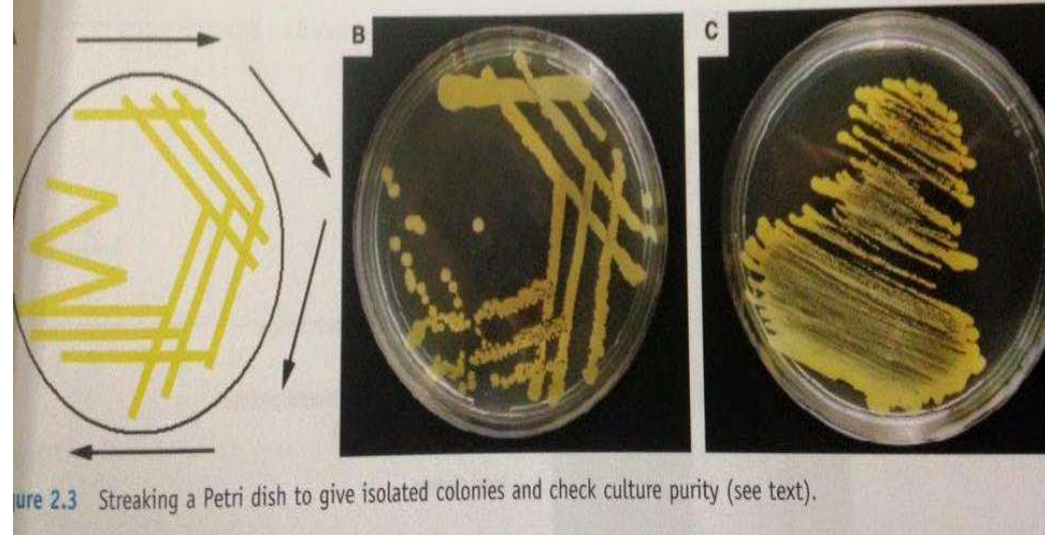
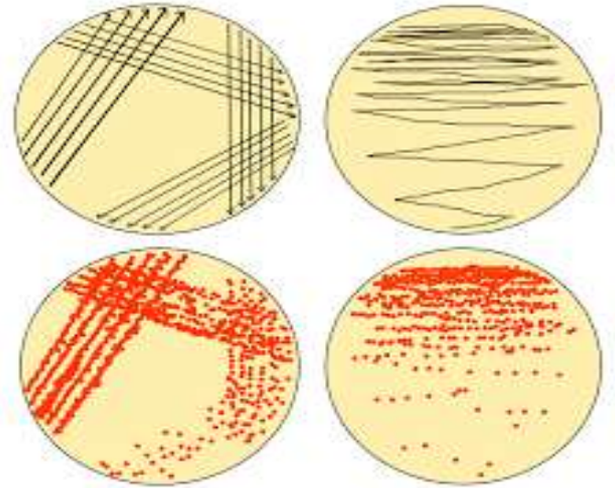
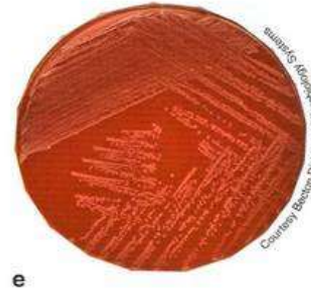
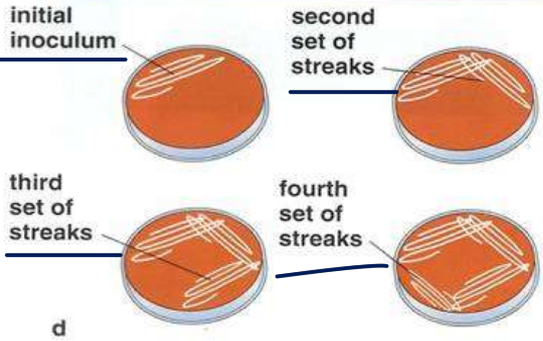
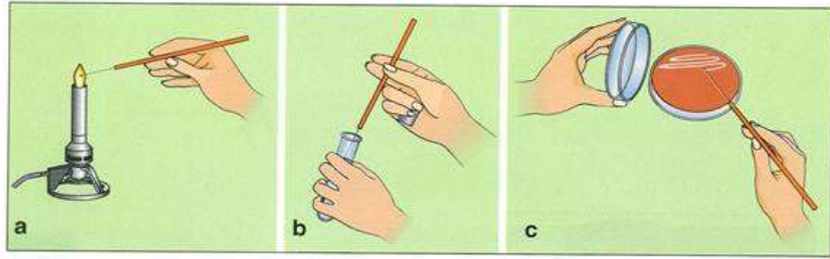
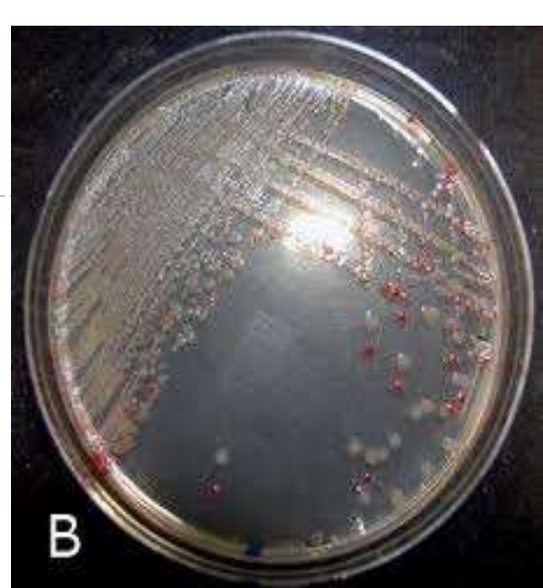


Figure 2.3 Streaking a Petri dish to give isolated colonies and check culture purity (see text).



Courtesy: Robert Dickinson, Microbiology

## Detection of objectionable MOs

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- ▶ Selective agar media recommended in the pharmacopeias:
- ▶ MacConkey's agar for E. coli
- ▶ XLD agar for Salmonella species
- ▶ Mannitol salt agar for Staphylococcus aureus
- ▶ Cetrimide agar for Pseudomonas aeruginosa

Students are required to read about each type of agar mentioned

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### General bacterial media [edit]

- Bile esculin agar (BEA)**

BEA is used for the isolation of *Enterococcus*, as well as Group D *Streptococcus* species

- CLED agar**

Cysteine, lactose, electrolyte-deficient agar (CLED) agar is used to isolate and differentiate urinary tract bacteria, since it inhibits *Proteus* species swarming and can differentiate between lactose fermenters and nonfermenters.

- Granada medium**

Granada medium is used to isolate and differentiate Group B streptococcus (GBS), *Streptococcus agalactiae* from clinical samples. GBS grows in granada medium as red colonies and most of accompanying bacteria are inhibited.

- Hektoen enteric agar (HEA)**

HE agar is designed to isolate and recover fecal bacteria belonging to the *Enterobacteriaceae* family. It is particularly useful in isolating *Salmonella* and *Shigella*.

- Lysogeny broth (LB)**

- MacConkey agar (MAC)**

MAC is a selective and differential medium used to differentiate between Gram-negative bacteria while inhibiting the growth of Gram-positive bacteria. The addition of bile salts and crystal violet to the agar inhibits the growth of most Gram-positive bacteria, making MacConkey agar selective. Lactose and neutral red are added to differentiate the lactose fermenters, which form pink colonies, from lactose nonfermenters that form clear colonies. An alternative medium, eosin methylene blue (EMB) serves a similar purpose.

- Mannitol salt agar (MSA)**

MSA is also a selective and differential medium. The mannitol indicates organisms that ferment mannitol: mannitol fermentation produces lactic acid, lowering the pH and turning the plate yellow. The salt is to select for halophiles; organisms that cannot withstand a high salt content are unable to grow well.

- Mueller-Hinton agar (MHA)**

MHA contains beef infusion, peptone, and starch, and is used primarily for antibiotic susceptibility testing. It can be in a form of blood agar.

- Nutrient agar**

Nutrient agar is usually used for growth of nonfastidious organisms and observation of pigment production. It is safe to use in school science laboratories because it does not selectively grow pathogenic bacteria.

- Önöz agar**

Önöz agar allows more rapid bacteriological diagnosis, as *Salmonella* and *Shigella* colonies can be clearly and reliably differentiated from other Enterobacteriaceae. The yields of *Salmonella* from stool samples obtained, when using this medium, are higher than those obtained with LEIFSON agar or *Salmonella-Shigella* agar (SSA).

- Phenylethyl alcohol agar (PEA)**

PEA selects for *Staphylococcus* species while inhibiting Gram-negative bacilli (e.g., *Escherichia coli*, *Shigella*, *Proteus*, etc.).



Hemolyses of *Streptococcus* spp. (left) α-hemolysis (*S. mitis*); (middle) β-hemolysis (*S. pyogenes*); (right) γ-hemolysis (= nonhemolytic, *S. salivarius*)



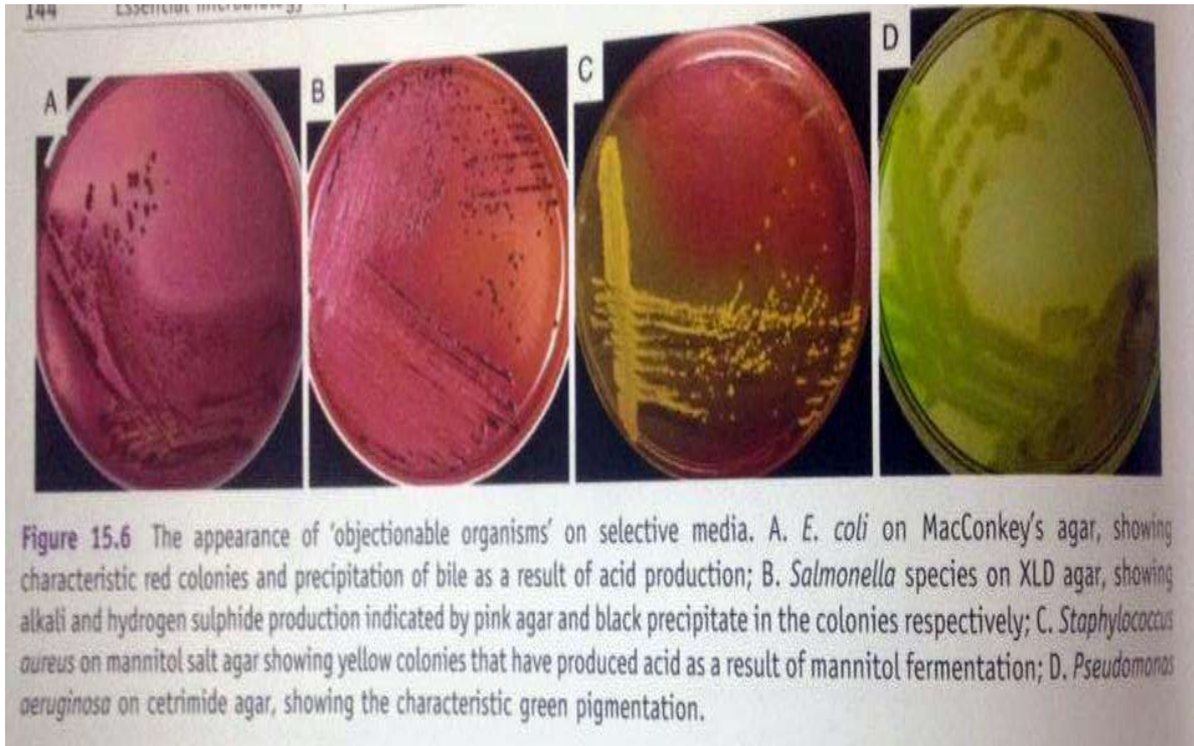
Four types of agar plate demonstrating differential growth depending on bacterial metabolism



Fungi (ascomycetes) growing in axenic cultures, each of which is a

# Detection of objectionable MOs

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# Automated bioburden determinations

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- ① Bases of the methods
- ② Limited application
- ③ Look for pg 144



# Bioburden specifications in the pharmacopeias

Table 15.4 Quality criteria for nonsterile medicines.

Route of administration <sup>a</sup>	Maximum total aerobic microbial count CFU/g or ml	Maximum total yeast and mould count CFU/g or ml	Specified microorganisms absent in 1 g or 1 ml
Nonaqueous oral products	$10^3$	$10^2$	Absence of <i>E. coli</i>
Aqueous oral products	$10^2$	$10^1$	Absence of <i>E. coli</i>
Rectal products	$10^3$	$10^2$	
Oral mucosal, gingival, cutaneous, nasal and ear products	$10^2$	$10^1$	Absence of <i>Staph. aureus</i> and <i>Ps. aeruginosa</i>
Vaginal products	$10^2$	$10^1$	Absence of <i>Staph. aureus</i> , <i>Ps. aeruginosa</i> and <i>Candida albicans</i>
Inhalation products (excluding nebulized liquids)	$10^2$	$10^1$	Absence of <i>Staph. aureus</i> , <i>Ps. aeruginosa</i> and bile-tolerant Gram-negative bacteria

