

2017 MICROSCOPE REVIEW

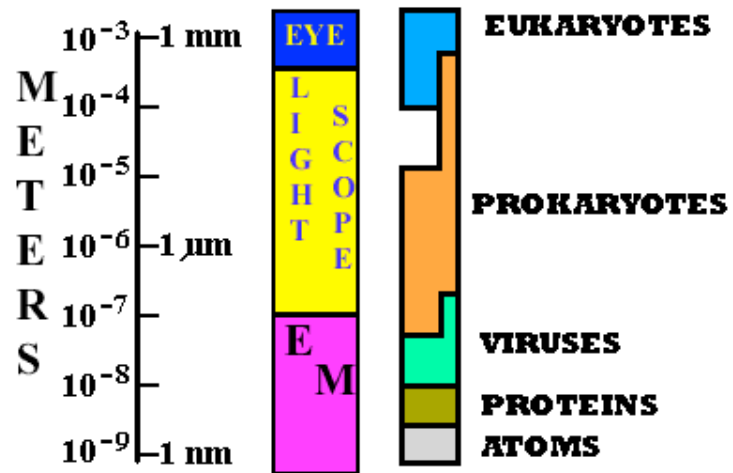
by Karen L. Lancour

RELATIVE SIZE OF MICROBES

1000 millimeters (mm) = 1 meter (m)

1000 micrometers (µm or mcm) = 1 millimeter (mm)

1000 nanometers (nm) = 1 micrometer (mcm)



Size of Commonly Known Microbes in micrometers (µm or mcm)

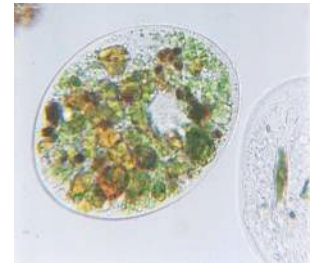
mcm	Type	Name	How known
500	Protozoa	Amoeba	Common in freshwater – have pseudopods
250	Protozoa	Paramecium	Common in freshwater – have cilia
200	Algae	Diatoms	Marine with silicon shells – important plankton
130	Protozoa	Euglena	Common in freshwater – have flagella & chloroplasts
10	Fungi	Yeast	Baker's yeast – makes dough rise
2	Bacteria	E. coli	Helps in digestion – one form causes serious food poisoning
2	Bacteria	Lactobacillus	Used to make yogurt – helps in our digestive system
1	Bacteria	Cyanobacteria	Has chlorophyll – carries on photosynthesis
1	Bacteria	Staphylococcus	Causes staph infections
0.3	Virus	Small pox virus	Causes small pox
0.15	Virus	Rabies virus	Causes rabies
0.10	Virus	Influenza virus	Causes the flu
0.03	Virus	Polio virus	Causes polio
0.03	Virus	Rinovirus	Causes the common cold

TYPES OF MICROSCOPES

Visible Light is the source of illumination: (up to 2000X)

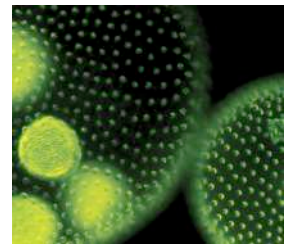
Bright Field Microscope (Lab – Light Compound Microscope)

- Most widely used
- Forms its image when light is transmitted through the specimen
- The specimen produces an image that is darker than the surrounding illuminated field
- Can be used with live, unstained and preserved, stain specimens



Dark Field Microscope

- A bright-field microscope can be adapted to a dark-field microscope by adding a stop to the condenser
- The stop blocks all light from entering the objective lens except for peripheral light
- The specimen produces an image that is brightly illuminated against a dark field
- Effective for visualizing living cells that would be distorted by drying or heat or that can't be stained with usual methods
- Does not allow for visualization of fine internal details of cells



Phase Contrast Microscope

- Used for live specimens
- Specimen is contrasted against a gray background
- Excellent for internal cell details



Differential - Interference Microscope

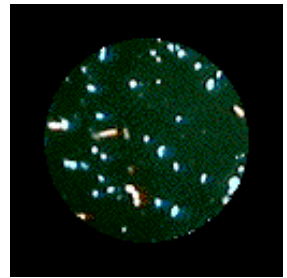
- Allows for detailed view of live, unstained specimens
- Includes two prisms that add contrasting colors to the image
- The image is colorful and three-dimensional



Ultraviolet rays are the source of illumination: (up to 2000X)

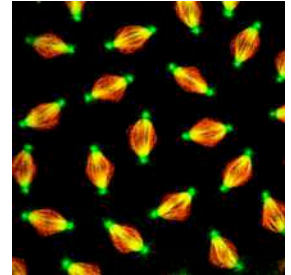
Fluorescence Microscope

- Includes a UV radiation source and a filter that protects the viewer's eyes
- Used with dyes that show fluorescence under UV rays
- Forms a colored image against a black field
- Used in diagnosing infections caused by specific bacteria, protozoa, and viruses using fluorescent antibodies



Confocal Microscope

- Allows for viewing cells at higher magnifications using a laser beam of light to scan various depths in the specimen
- Most often used on fluorescently stained specimens

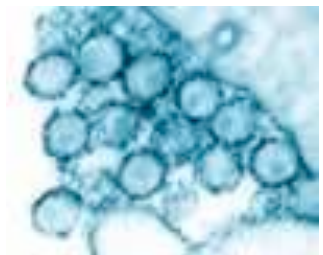


Electron beam forms image of specimen:

- Originally developed for studying nonbiological materials
- Biologists began using it in the early 1930s
- Forms an image with a beam of electrons
 - Electrons travel in wavelike patterns 1,000 times shorter than visible light waves
 - This increases the resolving power tremendously
- Magnification can be extremely high (between 5,000X and 1,000,000X for biological specimens)
- Allows scientists to view the finest structure of cells

Transmission Electron Microscope - TEM (up to 100,000X)

- Often used to view structures of cells and viruses
- Electrons are transmitted through the specimen
- The specimen must be very thin (20-100 nm thick) and stained to increase image contrast
- dark areas of a TEM image represent thicker or denser parts



TEM of HIV virus

Scanning Electron Microscope - SEM (up to 650,000X)

- Creates an extremely detailed three-dimensional view of all kinds of objects
- Electrons bombard the surface of a whole metal-coated specimen
- Electrons deflected from the surface are picked up by a sophisticated detector
- The electron pattern is displayed as an image on a television screen
- Contours of specimens resolved with SEM are very revealing and surprising
- The images may be computer enhanced to give them color



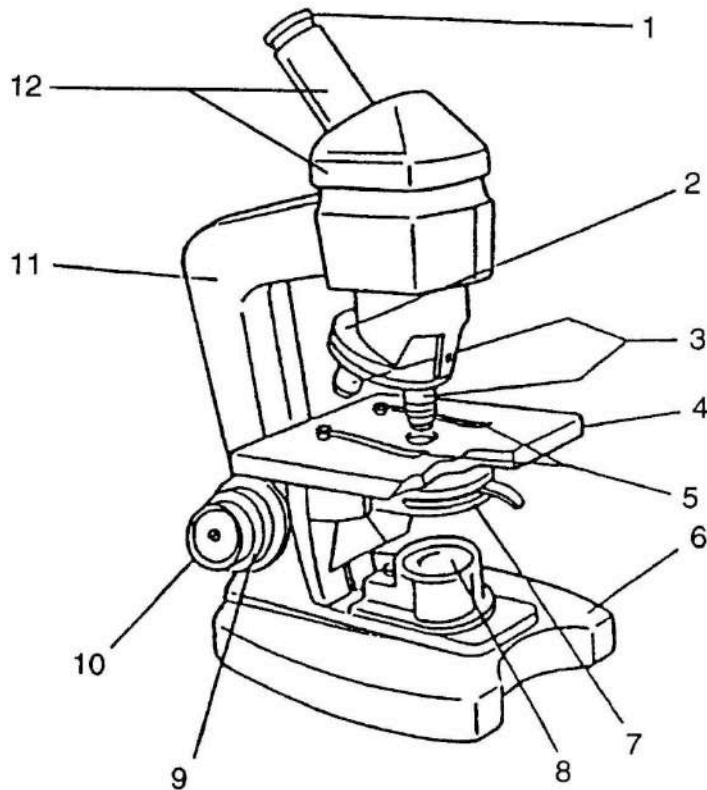
SEM of HIV virus

Note: Color can be added to the TEM & SEM images by computer

MICROSCOPY REVIEW – Light Compound Microscope

Parts of the microscope and their function: The number in front of each part of the microscope represents its number on the diagram.

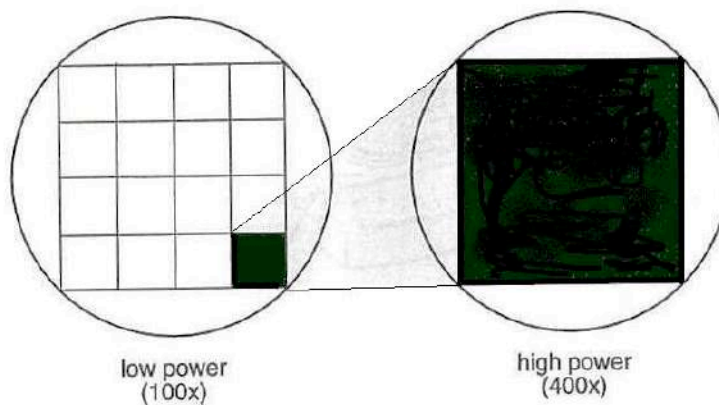
1. **ocular** - magnifies the image formed by the objective.
2. **nosepiece** - holds the objectives.
3. **objectives** - lenses that receive the light from the field of view and form the first image.
4. **stage** - supports the slide and the specimen.
5. **stage clips** - hold the slide in place.
6. **base** - foundation which supports the scope & keeps it stable.
7. **diaphragm** - controls the amount of light reaching the specimen.
8. **illuminator** - source of light.
9. **course adjustment** - used for initial or low power adjustment.
10. **fine adjustment** - used for fine tuning & high power focusing.
11. **arm** - supports the ocular, objectives and body tube.
12. **body tube** - tube or barrel between the ocular and the objectives.



PRINCIPLES OF MICROSCOPY

- A. Appearance of objects
1. Inverted and reversed (upside-down & backwards)
 2. If an "e" is placed in the stage in its normal position, it will appear as an "ə".
 3. Only a thin layer of the specimen is in focus at any level (depth of focus).
- B. Movement of specimens
1. Actual movement is opposite to appeared direction of movement.
 2. If an organism is actually moving (↗), it will appear to be moving (↙).
- C. Total magnification
1. Multiply ocular magnification times objective magnification.
 2. Oculars are normally 10X or 12X.
 3. Objectives are typically as follows:
 - a. scanning power - 4X or 5X or 6X
 - b. low power - 10X or 12X
 - c. high power - 40X or 43X or 45X
 4. Sample problem: If the ocular is 10X and the objective is 43X, the total magnification is 430.
- D. Changing objectives
1. When changing objectives from scanning power to lower power to high power, the following changes will occur:
 - a. the size of the field of view decreases.
 - b. the field of view is darker.
 - c. the size of the image increases..
 - d. the resolution (ability to separate small details) increases.
 - e. the working distance (distance between coverslip & objective) decreases.
 - f. the depth of focus (thickness of the specimen which may be seen in focus) is reduced.
 2. The relationships of magnification and the fields of view diameter and area ratios are approximately (10 X ocular and listed objective)

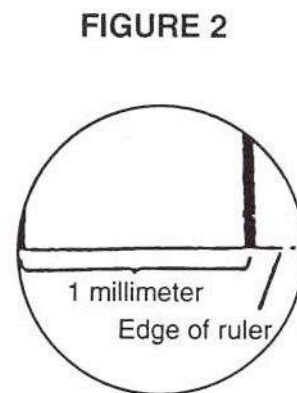
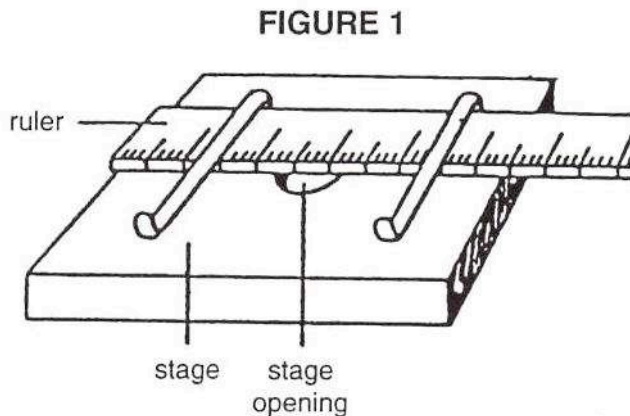
<i>objective</i>	<i>total magnification</i>	<i>diameter</i>	<i>area</i>
scanning 5X	50	-	-
low 10X	100	1/2 scanning	1/4 scanning
high 40X	400	1/4 low	1/16 low



Principles of advanced microscopy

A. Measuring the diameter of the field of view.

1. Place a transparent millimeter ruler on the stage, hold it down with the stage clips, and observe the ruler in the desired field of view - scanning or low. (See Fig. 1)
2. Focus on the metric edge of the ruler. Hint: applying gentle pressure to the free end of the ruler will help adjust for the thin ruler and allow for better focus.
3. Place the center of one millimeter marking at the left edge of the field of view (see Figure 2) and measure the diameter of the field in millimeters. For reference: Scanning power (50X) is about 3.0 to 3.2 mm and low power (100X) is about 1.5 to 1.6 mm. Sizes will vary with changes in magnification and manufacturer.



4. Convert the measurement in millimeters to micrometers by multiplying by 1000.

Samples:

low power field 1.6 mm X $\mu\text{m}/\text{mm}$ = 1600 μm or mcm

scanning power field 3.2 mm X 1000 $\mu\text{m}/\text{mm}$ = 3200 μm or mcm

5. The diameter of the high power field is less than one millimeter. It can be calculated from the diameter of the low power field by using the following formula:

$$\frac{\text{high power field diameter}}{\text{lower power field diameter}} = \frac{\text{low power magnification}}{\text{high power magnification}}$$

Sample:

$$\frac{\text{h.p. field}}{1600 \mu\text{m}} = \frac{100}{400} \quad \text{to} \quad \text{h.p. field} = \frac{100 \times 1600}{400} = 400 \mu\text{m or mcm}$$

B. Estimating the size of an object.

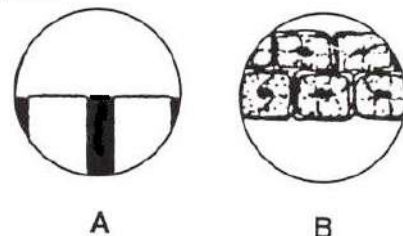
1. Determine the number of cells or objects that would fit across the diameter of the field of view.
2. Estimate the size of the object by dividing the diameter of the field by the number of the objects that would fit across the field.

Sample: a. diameter of the field is

$$2 \text{ mm} \times 1000 \mu\text{m}/\text{mm} = 2000 \mu\text{m}$$

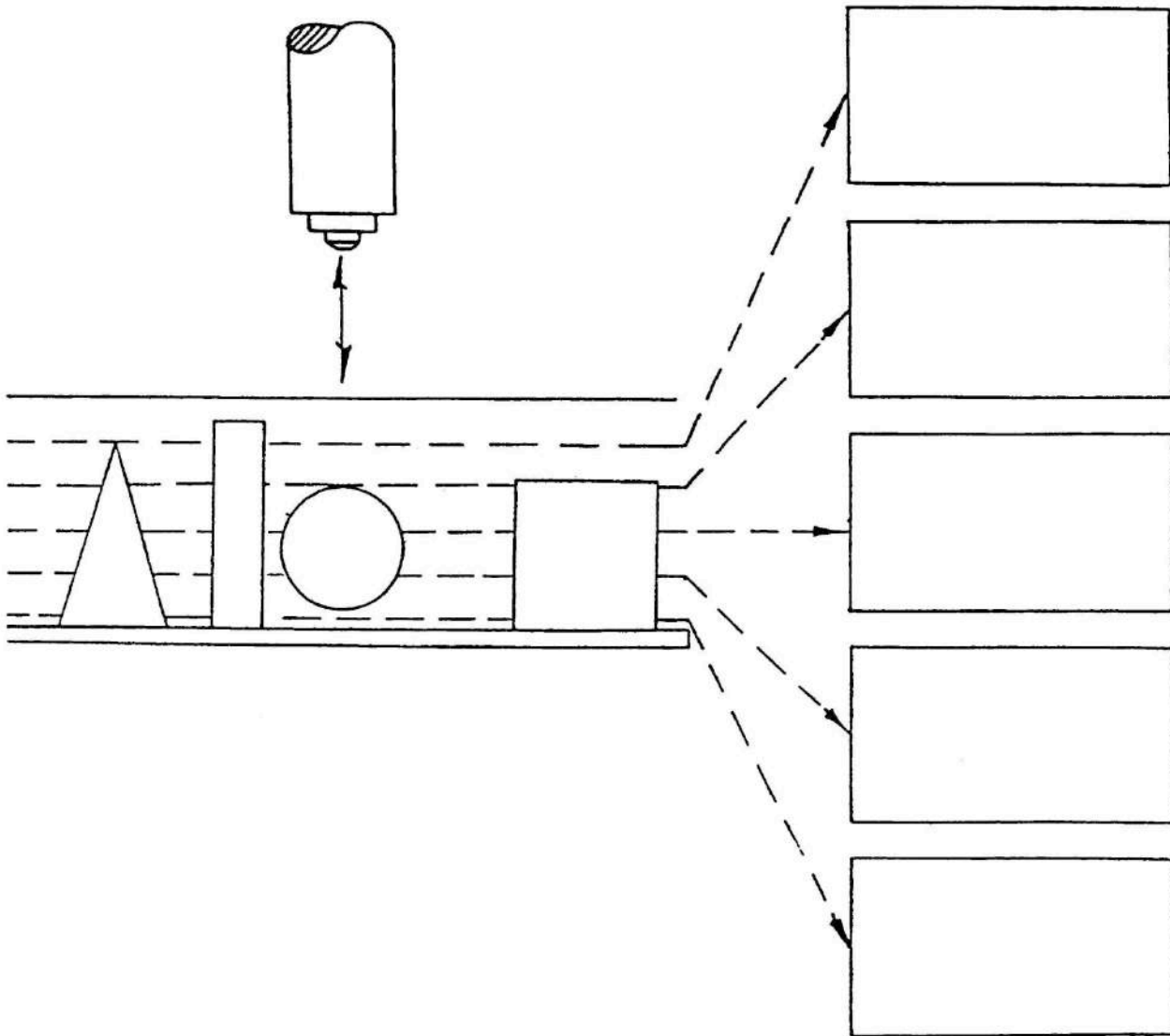
b. length of cell "x" is

$$2000 \mu\text{m} / 3 \text{ cells} = 660 \mu\text{m}/\text{cell}$$



DEPTH OF FOCUS EXERCISE

Below are four objects located between a slide and a coverslip. The actual objects would of course be 3-D. The objects are left to right; a cone, a cylinder, a sphere and a cube. As one focuses down through the various levels a two dimensional representation will be visible. At each level (represented by the dotted lines to the slide diagram) draw what two dimensional shapes would be present and give their proper location on the slide diagram.



ANSWER KEY FOR DEPTH OF FOCUS EXERCISE

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