

## Exercise 4.1 – Using the Light Microscope

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### LEARNING OUTCOMES

1. List the ways in which a microscope is properly maintained and stored.
  2. Identify and give the function of key parts of a compound light microscope.
  3. Discuss the principles of magnification and resolution; define key terms.
  4. Calculate total magnification.
  5. Use the scanning, low power, and high-power objective lenses to focus the letter “e.”
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### Microscope Care

Even a very powerful microscope cannot deliver high-resolution images if it is not properly cleaned and maintained. Microscopes are rather delicate instruments, and great care must be taken to avoid damaging parts and surfaces.

Each student is assigned a microscope for use during the semester. Be sure to record the number of your microscope and follow the guidelines below when obtaining and storing it.

Care and Storage of the Light Microscope
1. Carry your microscope with two hands, one on the arm and the other under the base.
2. Lift and place the microscope to reposition it on the benchtop; do not drag it.
3. Clean the stage and objective lenses before and after use with lens paper/cleaner.
4. Lower the light intensity before turning off the microscope.
5. Move the stage to its lowest position before storage.
6. Position the 4X objective to point down toward the stage before storage.
7. Return your microscope to the corresponding number compartment in the cabinet.

Basic components of the light microscope are shown in Figure 4.3.

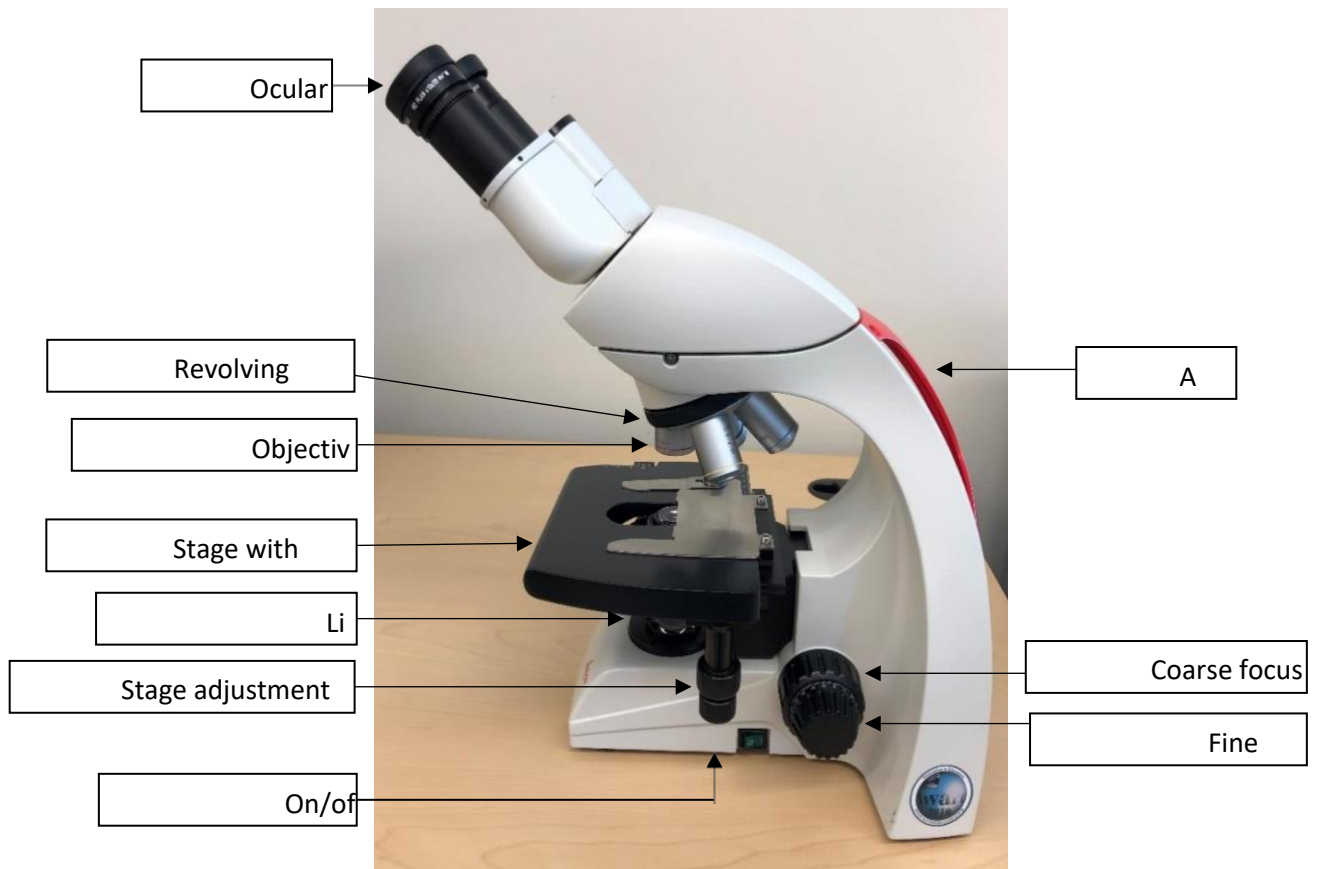


Figure 4.3a: Components of a typical brightfield microscope.

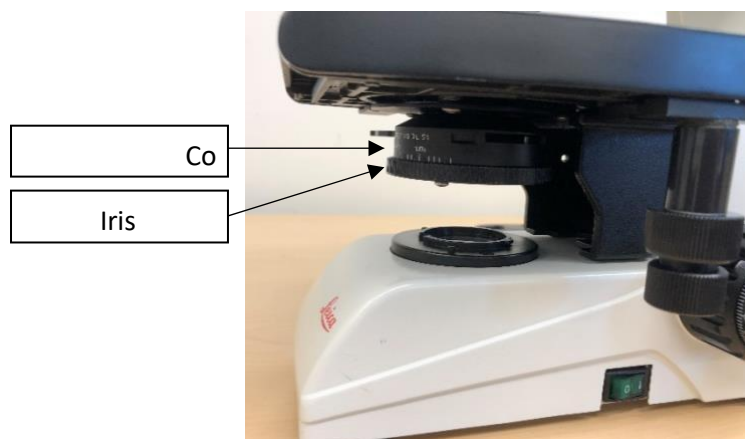


Figure 4.3b: The amount of light traveling through the condenser lens is controlled by turning the adjustment for the iris diaphragm located inside the condenser.

## Summary of Microscope Components

- ❑ **Ocular lens:** Eyepiece that usually magnifies 10X; binocular microscopes have two oculars that are adjustable for interpupillary distance between the eyes. Oculars may have a pointer and/or a ruler for measuring cells called an ocular micrometer.
- ❑ **Revolving nosepiece:** Rotates to allow each objective to align in place with the ocular.
- ❑ **Objective lenses:** Seated in the nosepiece. Each objective lens has a different magnifying power: scanning (4X), low power (10X), high power (40X) and oil immersion (100X).
- ❑ **Coarse focus knob:** Outer large knob that raises and lowers the stage to bring the specimen into initial focus; used with the scanning objective lens.
- ❑ **Fine focus knob:** Smaller inner knob that raises and lowers the stage to bring the specimen into sharp focus; used with low power, high power, and oil immersion lenses.
- ❑ **Mechanical stage:** Horizontal surface on which slide is placed and held by stage clips; the stage is moved left and right by turning the x-y mechanical stage knobs.
- ❑ **Illuminator:** Light source turned on by a switch on the base and controlled by a rheostat located on the side of the base that adjusts the brightness of the light.
- ❑ **Iris diaphragm and condenser:** The iris diaphragm can be adjusted to control the amount of light passing from the illuminator through the bottom of the slide. It is located inside the condenser, which is a lens system that gathers and directs light up from the illuminator.

## Magnification and Resolution

Light microscopes use visible light and a series of lenses to view microscopic specimens. The condenser lens focuses the light as it goes through the specimen and can be adjusted for optimization. The objective lenses magnify the specimen, capturing the transmitted and reflected light to create a real image of the specimen. The ocular lens further magnifies the image and creates a virtual image for viewing. This difference can be observed with a slide of the letter “e” or “p” and noting how the image changes when viewed through the ocular.

What is observed through the microscope is the *field of view*. While an entire organism might be visible in the field of view using the scanning lens, only a small portion of it may be seen under high power. Since microorganisms have a wide range of sizes, the most appropriate objective to use for each varies. For example, while a large protist such as *Amoeba* may be viewed under low power, this would not be suitable for viewing bacteria which are much smaller.

Most modern microscopes are *parfocal* in that they remain in relative focus when changing magnifications. This property eliminates the need for extensive re-focusing when switching between objective lenses.

Related to the concept of field of view are *depth of field* and *working distance*. Depth of field refers to the nearest and furthest planes of a specimen that are in focus at the same time. Depth of field depends on thickness of the specimen and decreases as magnification increases. The working distance, or space between the slide and objective lens, decreases as magnification increases. To avoid damaging the objective lenses or the slide, the coarse focus knob should only be used for initial focus when the working distance is greatest.

## Calculating Total Magnification

Magnification is the process of making an object appear larger than it is. The magnification of each objective is printed on the metal portion of the lens. The *scanning* objective has a magnification of 4X and is used when first bringing an image into focus. The next objective is *low power* which magnifies 10X. The high-power objective, sometimes called the *high dry* objective because it is used without immersion oil, magnifies 40X and is used when fine focusing an image. Finally, the *oil immersion* objective has a magnification of 100X and is used when viewing bacterial cells.

The ocular and objective lenses work together to create a magnified image. Total magnification (TM) is calculated by multiplying the ocular and objective magnifications:

$$\text{Total magnification} = (\text{ocular magnification}) \times (\text{objective magnification})$$

For example, if the ocular is 10X and the 40X objective lens is selected, TM is  $(10X)(40X) = 400X$ . Total magnification using each objective lens for your microscope is given in Table 4.1.

Table 4.1. Total Magnification

Objective Lens Magnification	Ocular Lens Magnification	Total Magnification
Scanning (4X)	10X	40X
Low power (10X)	10X	100X
High power (40X)	10X	400X
Oil immersion (100X)	10X	1000X

Unlike magnification, *resolution* is the ability to distinguish two objects as separate entities. The resolving power for a light microscope is about 0.2 micrometers, meaning any two objects that are closer than two tenths of one micrometer will be seen as a single point.

The following exercise is designed to provide practice using the light microscope to view a slide with the letter “e.” Work through the steps slowly and apply the same principles when viewing stained slides in later exercises.

## Exercise 4.1 – Using the Light Microscope: Viewing the Letter “e”

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### OBJECTIVE

Use the light microscope to practice focusing under scanning, low power, and high power.

### MATERIALS

- ❑ EQUIPMENT: Light microscope, Sta-clear paper, lens paper, lens cleaner  
❑ SLIDE: Letter “e”

### PROCEDURE - *Take your time and work through steps in order.*

1. Obtain a microscope from the cabinet. Remember to carry it with two hands and reposition it on the bench by lifting rather than dragging.
2. Place the microscope directly in front of you on the bench. Sit up straight and push in your chair so that you are comfortable. Do not bend over or kneel on your chair to view slides.
  - Record the number that is found on the back of your microscope: \_\_\_\_\_
3. Verify that the student before you stored the microscope correctly, making sure:
  - \_\_\_\_\_ The stage is clean, has no slides, and is free from oil.
  - \_\_\_\_\_ The scanning (4X) objective lens is pointing down toward the stage.
  - \_\_\_\_\_ The stage is lowered completely.
  - \_\_\_\_\_ The rheostat (light intensity dial on the base) is turned down all the way.
4. Clean the oculars and objective lenses with lens paper and lens cleaner, checking that each objective lens is securely screwed into the revolving nosepiece.
  - Record the magnification printed on the oculars: \_\_\_\_\_X
5. Plug in your microscope and turn it on using the power switch on the base.
6. Raise the light intensity by turning the rheostat to a high number on the base and adjust brightness by closing the iris diaphragm rather than lowering the rheostat.
7. Move the oculars together or apart so that you can use both eyes to view the slide. Note that one ocular will have a pointer and the other will have a micrometer for measuring cells.
  - Record the interpupillary distance between the oculars: \_\_\_\_\_

8. Obtain a slide of the letter “e” from the slide tray and clean it using Sta-Clear paper and lens cleaner.
9. Place the slide on the stage with the label facing up and to the left, securing corners in the stage clips so that it lies flat and pushing the slide back as far as it will go.
  - Record the appearance of the letter as it appears looking at the stage: \_\_\_\_\_
10. Using the stage control knobs, position the slide so that the letter is over the light source.
11. Look through the oculars and keep turning the coarse focus knob until the image comes into focus. This may require significant rotation of the focus knob. If you go too far and miss the image, turn the knob slowly in the opposite direction.
  - Record the appearance of the letter as it appears through the oculars: \_\_\_\_\_
  - Record the total magnification using this objective: \_\_\_\_\_X
  - Circle the appearance of a “p” as it would be viewed through the oculars: p   d   b   q
12. View the slide under low power by rotating the 10X objective in place and turning the fine focus knob until the image is clear. If necessary, adjust the iris diaphragm to lower the light.
13. If directed to do so, raise your hand for the instructor to verify your observation.
  - Record total magnification using this objective: \_\_\_\_\_X
  - Which property maintains focus while changing objectives? \_\_\_\_\_
14. View the slide under high power by rotating the 40X objective in place and turning the fine focus knob until the image is clear. Increase light by opening the diaphragm. If the image is blurry, use lens paper to firmly clean the bottom of the objective.
  - Record total magnification using this objective: \_\_\_\_\_X
  - What happens to the field of view as magnification increases? \_\_\_\_\_
  - Which objective lens is most appropriate for viewing the letter? \_\_\_\_\_
15. Return the slide to the corresponding numbered slot on the tray.
16. When you are done using the microscope, prepare it for storage by ensuring:
  - \_\_\_\_\_The stage is clean, has no slides, and is free from oil.
  - \_\_\_\_\_The scanning (4X) objective lens is pointing down toward the stage.
  - \_\_\_\_\_The stage is lowered completely.
  - \_\_\_\_\_The rheostat (light intensity dial on the base) is turned down all the way.
17. Show your microscope to the instructor before returning it to the cabinet.