BIOLOGY NOTESONK

MPI

NAME	_
SCIENCE NUMBER	_
HOLT USER NAME	_
HOLT PASSWORD	_
LAB PARTNER	_
LAB PARTNER EMAII /PHONE/CONTACT	

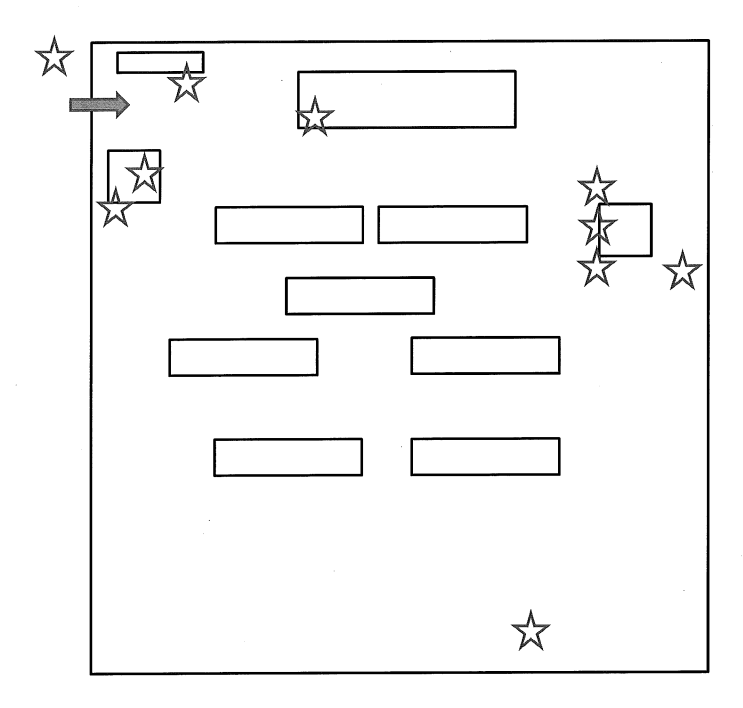
I AM A	LEARNER.

THE THINGS THAT I CAN DO THAT MAKES IT EASIER TO STUDY ARE

*

*

*



Bio Basics

(dissecting)scanning

objective

low power high power field of view micrometer

dependant variable independent variable fine adjustment knob

coarse adjustment knob

scanning electron microscope

Chap 1

Skepticism
Observation
Hypothesis
Experiment
control group

theory SI

Biology cell

homeostasis

universal laws correlation data

bias

-Homeostasis-

Metabolism Responsiveness

Heredity

Chap 3

Atom

Compound

Element Ion

Molecule valence electron

acid

adhesion base

buffer cohesion

рH

solution amino acid

ATP

Carbohydrate

DNA Lipid

RNA

nucleic acid nucleotide protein

activation energy

active site energy enzyme product reactant substrate

Covalent

Ch 7

Cell membrane

-Cytoplasm=

Ribosome Prokaryote Eukaryote Nucleus

Organelle Vesicle

endoplasmic reticulum

Golgi apparatus

Vacuole Chloroplast Mitochondrion Flagellum

Tissue Organ

organ system

colonial organism

Ch 8

Phospholipid lipid bilayer equilibrium

concentration gradient

diffusion

carrier protein

osmosis

sodium-potassium pump

signal

receptor protein
second messenger
Hypertonic solution
Hypotonic solution
Isotonic solution

Concentration gradient

Lab must be typed! Sections must be <u>headed and</u> <u>bold and underlined</u>. Remember to label all graphs tables and charts with a TITLE! Be sure to include your name and science number. 25% off each day late

Title: What is the theme of this lab? Make sure to include a specific scientific theory/process or fact. 20 points- this includes points for names and science number as well as formatting.

Hypothesis: include independent and dependant variables- no questions! 10 points

Procedures/Experimental Design: numbered with full sentences. Be specific! The procedures should allow someone to recreate the experiment in absentia. 20 points

Data: include results, tables, graphs, photos and measurements. All graphs must have axes properly labelled and titled or no credit will be given for the graph. 30 points

Conclusion: use your data AND research to explain why or why not your hypothesis was accurate. This should be the WHY. Explain any possible experimental error here. 20 points

1

HOW TO GET A GOOD LAB GRADE

Here is an example of the points and what is expected. There will be a "cheat" sheet on the homework website for each lab report

NAME(S)

Sci# (both names must be included if a group lab is submitted)

TITLE: This should include the title of the lab and the chapter. Ex. Ch 3 Biomolecules of Food Lab.

20 points (includes name, science number, proper format)

<u>PURPOSE</u>: State what the purpose of this lab is and a hypothesis when applicable. Ex. In this lab, we will be examining the amount and proportions of carbohydrates, fats and protein in a typical fast food meal. 10 points

<u>BACKGROUND</u>: List the relevant information regarding the lab. This should be AT LEAST one paragraph Ex. The four biomolecules are carbohydrates, lipids, proteins and nucleic acids. We will be measuring.....

20 points

DATA: This is where microscope drawings, data tables, graphs and photos are included. EVERYTHING must be labeled and titled. For microscope drawings and pics, the magnification as well as the object must be given. All tables and graphs MUST have a title, key and the axes must be properly labeled. If this information is not included, the data will be given a zero. 30 points

CONCLUSION: you must relate the materials

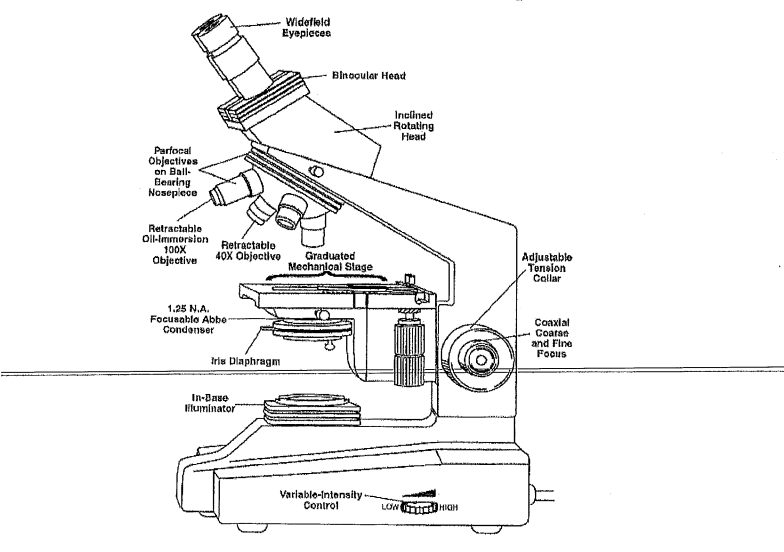
gathered/viewed/observed/collected in lab to the background information as well as any relevant information from class or previous labs. This is also a summary of what the lab showed or was able to exhibit. This will be AT LEAST a paragraph. 20 points

Chapter 0 BIO BASICS

1. Knowing the Microscope

- 1. Review the parts of the compound microscope
- 2. It is important that you have memorized the objectives
 - a. Dissecting = 4X
 - b. Low power= 10X
 - c. High Power = 40X
 - d. Oil (not generally used in this class)=100X
- 3. Know your microscope number, where it belongs, and know how to store it properly. If you do not put your microscope away properly at the end of class, you will receive a detention!

Advanced and Research Microscopes



2. Calculating Magnification

- 1. Look for the number marked with an X on the
 - a. Eyepiece
 - b. Low power objective
 - c. High power objective
- 2. Multiply the number on the eyepiece by the number on the objective.

Ex- eyepiece is 10X and the objective is 40X. the total magnification is 400X

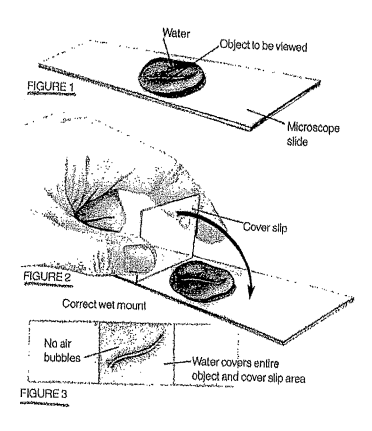
3. Using the Microscope

It is important that you become familiar with using the microscope. If you can not find and focus on an object, you will not be able to complete the labs in the allotted time.

- 1. Take the microscope that corresponds to your science number. If it is not put back correctly (not underdissecting power, not shut off, slide on the stage, cord not wrapped) tell me immediately.
- 2. Carry the microscope to your lab station with one hand under base and the other around the arm grasping the cord so you do not trip.
- 3. Place the microscope on the table and plug it in. Make sure the green light is lit on the outlet. If it isn't, hit the reset button with your finger-not an object.
- 4. Clean the eyepiece and objective with Kimwipes- DO NOT USE PAPER TOWELS- they scratch the lenses. If the microscope is very dirty, you may use alcohol on the Kimwipe. Use a small amount.
- 5. Make sure the diaphragm is open and turn on the microscope.
- 6. Place the slide (coverslip and/or label) facing UP. Make sure the object to be viewed (or the coverslip) is directly over the stage aperture. The light should shine directly through the object or coverslip.
- 7. The microscope should already be under dissecting lens (4X). You should now move the lens to low power (10X). Make sure you hear it click into place or you will see nothing. Without looking through the eyepiece, lower the low power objective til it is almost touching the coverslip.
- 8. Looking through the eyepiece, raise the objective using the rough adjustment knob. When the object comes into focus, stop and switch to fine adjustment. This will allow you to focus clearly. If you have difficulty observing the object, close the diaphragm slightly. This will allow you to see the object with a darker outline.
- 9. Using the stage adapters, move the slide until the object to be viewed is directly in the middle.
- 10. Switch to high power (40X) being careful to not crack the lens on the slide.
- 11. Look through the eyepiece and focus USING THE FINE ADJUSTMENT ONLY!! You will crack the slide and objective if you use the coarse adjustment!
- 12. When finished, return the microscope to dissecting objective (4X), remove the slide, clean anything you spilled on the stage or lenses, shut the power button, coil the cord and return to its parking spot.

4. Making a Wetmount

- 1. Add a drop of water to the center of a clean microscope slide.
- 2. Place the object to be viewed in the drop of water.
- 3. Pick up a coverslip by its edges. Do not touch the surface of the coverslip. Stand the coverslip on its edge next to the drop of water
- 4. Slowly lower the coverslip over the drop of water and the object to be viewed.

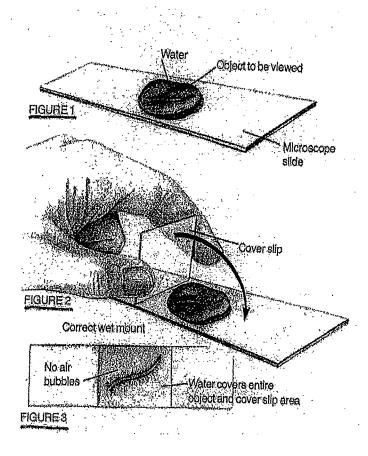


Troubleshooting

- 1. Not enough water: air bubbles will form. Air does not refract light in the same way as water- therefore you will not be able to see an object in or near an air bubble. Air bubbles will appear as dark black dots or lines
- 2. Too much water: water will come out from under coverslip and coverslip will be floating and moving. Take a paper towel and touch the edge of the paper towel to the edge of the coverslip. This is also how you pull stain across a specimen.

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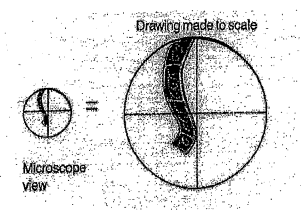
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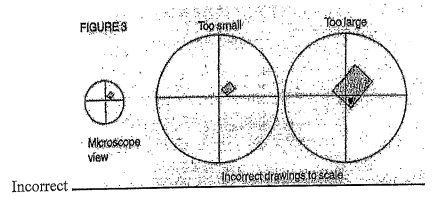
4. Making Scale Drawings

When you draw objects seen through the microscope, the size that you make your drawing is important. Your drawing should be in proportion to the size the object appears to be when viewed through the microscope. This is called drawing to scale. This allows you to compare the sizes of different objects.

- 1. Draw a circle on the paper
- 2. Imagine the circle divided into 4 equal sections
- 3. When looking through the eyepiece, imagine the same 4 equal sections
- 4. Note how much of the object takes up each quadrant. Draw each quarter exactly as it appears in the eyepiece.



Correct



6. Writing a Hypothesis

A hypothesis is a possible or tentative explanation for a question or problem. A properly written hypothesis has a dependent and independent variable.

Dependent Variable- this is what may happen because of the independent variable. In other words it depends on the independent variable.

Independent Variable- this is what is having an effect on the dependent variable

Here are some Examples to help clear this up!

Problem: Does the amount of air in a basket ball determine how high it will bounce?

Hypothesis:

The amount of air in a basketball affects how high it will bounce.
(I.V.) (D.V.)

Problem: Does the temperature affect how active the lizard is?

Hypothesis:

Warmer temperatures increase the activity level of a lizard.
(I.V.) (D.V.)

Problem: IS the speed that a boy walks affected by how baggy his pants are?

Hypothesis:

The speed that boy walks is affected by how baggy his pants are

(D.V.) (I.V.)

Listed below are some already written hypotheses. Underline the independent variable and circle the dependent variable in each one. Basically, the dependent variable is something that can be measured like speed, height, odor, etc. and the independent variable is something that is causing different amounts of the thing being measured.

- 1. The amount of sunlight a plant gets affects how tall the plant will grow.
- 2. Female elks with higher level of hormones will migrate faster
- 3. A teacher's attitude is affected by the number of students in her class

15

- 4. The amount of sleep a students gets before a test affects the score he or she earns on a the test
- 5. A person's sex determines how fast they can learn.
- 6. The amount of hairspray a girl uses affects the number of boys who ask her out.
- 7. Eating broccoli increases the number of correct answers on a math test
- 8. Applying fertilizer affects the number of weeds growing in a yard
- 9. The amount of rainfall affects how many flowers a cactus produces
- 10. A rougher road increases the number of times you fall when rollerblading.
- 11. Telling your mother that she is a good cook increases the hour of your curfew.
- 12. Washing the dishes for your mother increases the amount of money she gives you on the weekend.
- 13. Coaches with more years of experience will have a higher percentage of wins.
- 14. The amount of food that a bird eats is affected by the temperature.
- 15. Eating chocolate affects the number of zits you get.

Here is something a little different to try. Now I am going to give you the problem and you have to write a hypothesis. <u>Underline</u> the independent variable and **circle** the dependent variable for each hypothesis you create.

- 1. Does the number of holes in your pants affect the number of detentions you get?
- 2. Does the color of a person's hair affect the scores they get on tests?
- 3. Does the color of a T-shirt you wear affect the number of people who smile at you?
- 4. Does the type of music you listen to affect your grades?
- 5. What affect does the temperature have on the length of an animal's hair?

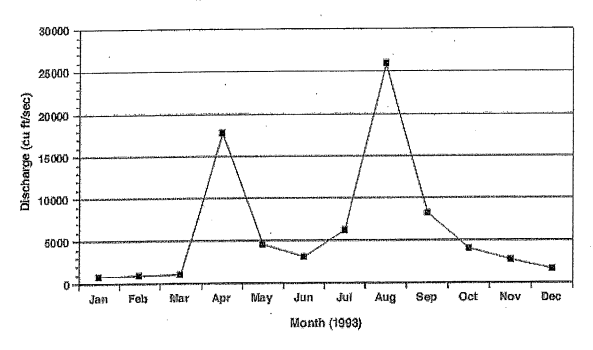
- 6. Does the amount of salt in water affect how fast it will boil?
- 7. Does the way a boy's hair is cut affect how many girl's like him?
- 8. Does music have an affect on the number of eggs a chicken will lay?
- 9. What effect does the price of a pair of jeans have on how good they fit?
- 10. What affect does cockroach poison have on the number of cockroaches in a house?

7. Graphing

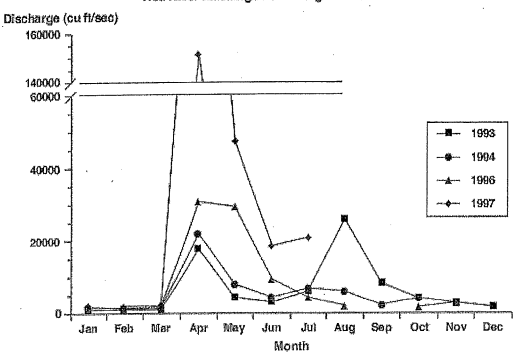
Line Graphs

Line graphs are very useful to plot a value over time. Line graphs are useful when both variables are quantitative (numerical). The line can be studied to find the slope, which can be useful for studying certain properties. The slope is a tool used to mathematically express a trend in the data.

Red River Discharge Rate - Fargo Station

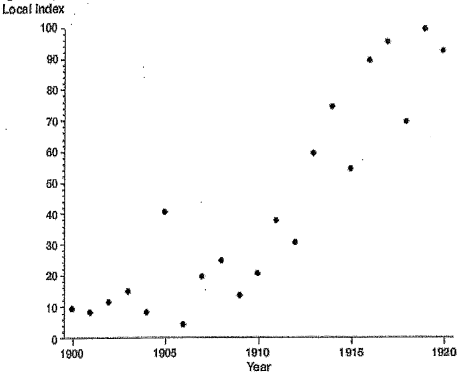


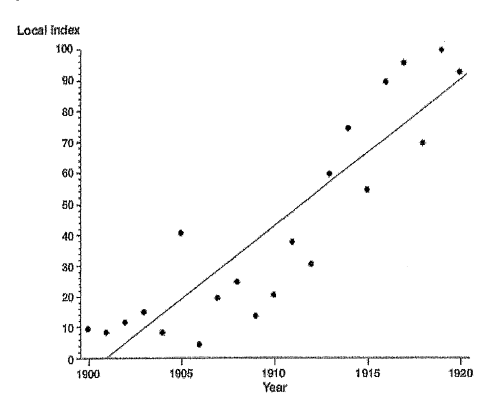
Red River Discharge Rate - Fargo Station



Scatter Plot

With a scatter plot a mark, usually a dot or small circle, represents a single data point. With one mark (point) for every data point a visual distribution of the data can be seen. Depending on how tightly the Points cluster together, you may be able to discern a clear trend in the information

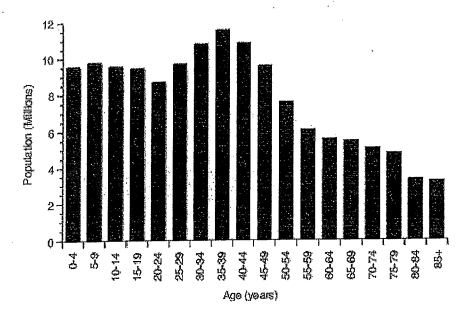




Histogram

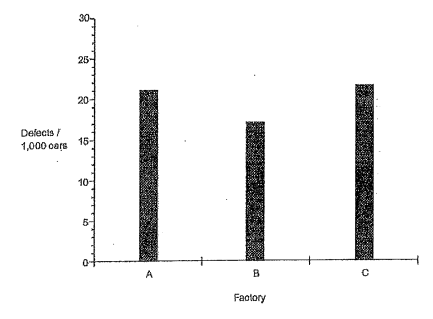
A histogram is a graphic display of frequencies of a value. It is the graphical equivalent of the table of how many of each cateogory fall into the range... The histogram differs from a bar chart in that it is the *area* of the bar that denotes the value, not the height, a crucial distinction when the categories are not of uniform width.

United States Femals Population 1997



Bar Graphs

Bar graphs are a coomon type of graph that are best suited for qualitative information, such as name or group. (there is no unbiform distance between the bars- due to qualitative nature- and a slope can NOT be derived from the information presented.



Name	C	lass	Date

LABORATORY SKILLS

Using Graphing Skills

Pre-Lab Discussion

Recorded data can be plotted on a graph. A graph is a pictorial representation of information recorded in a data table. It is used to show a relationship between two or more different factors. Two common types of graphs are line graphs and bar graphs. In this investigation, you will interpret and construct a bar graph and a line graph.

Problem

How do you correctly interpret and construct a line graph and a bar graph?

Materials

No special materials needed

Procedure

Part A. Interpreting Graphs

1. The type of graph that best shows the relationship between two variables is the line graph. A line graph has one or more lines connecting a series of points. See Figure 1. Along the horizontal axis, or x-axis, you will find the most consistent variable in the experiment. Along the vertical axis, or y-axis, you will find the other variable.

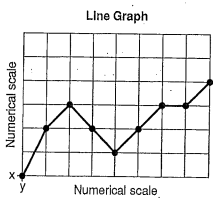


Figure 1

2. Use the line graph in Figure 2 to answer questions 1 through 6 in Observations.

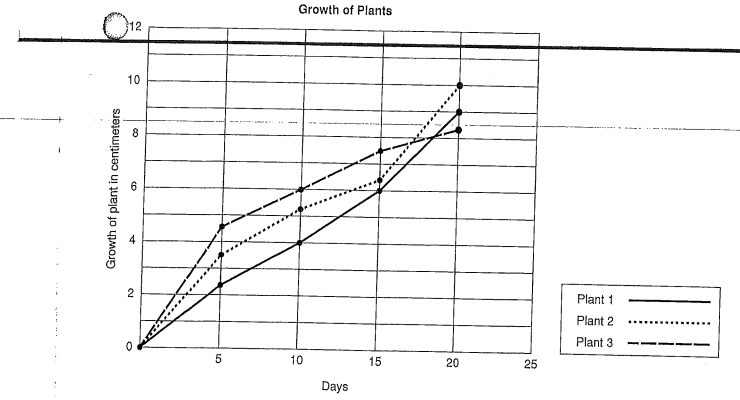


Figure 2

3. A bar graph is another way of showing relationships between variables. A bar graph also contains an x-axis and a y-axis. But instead of points, a bar graph uses a series of columns to display data. See Figure 3. On some bar graphs, the x-axis has labels rather than a numerical scale. This type of bar graph is used only to show comparisons.

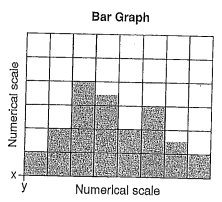
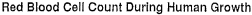


Figure 3





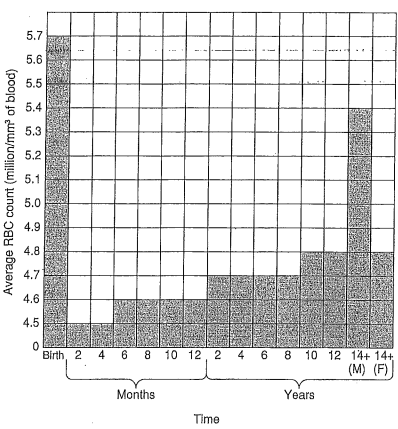


Figure 4

Part B. Constructing Graphs

- 1. When plotting data on a graph, you must decide which variable to place along the x-axis and which variable to place along the y-axis. Label the axes of your graph accordingly. Then you must decide on the scale of each axis; that is, how much each unit along the axis represents. Scales should be chosen to make the graph as large as possible within the limits of the paper and still include the largest item of data. If the scale unit is too large, your graph will be cramped into a small area and will be hard to read and interpret. If the scale unit is too small, the graph will run off the paper. Scale units should also be selected for ease of locating points on the graph. Multiples of 1, 2, 5, or 10 are easiest to work with.
- 2. Use the information recorded in Data Table 1 to construct a line graph on the grid provided in number 12 of Observations. You should label each axis, mark an appropriate scale on each axis, plot the data; connect the points; and give your graph a title.
- 3. Use the information recorded in Data Table 2 to construct a bar graph on the grid provided in number 13 of Observations. You should label each axis, mark an appropriate scale on each axis, plot the data, darken the columns of the graph, and give your graph a title.

Observations

Part A. Interpreting Graphs

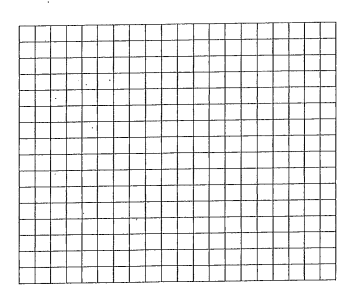
Use the line graph in Figure 2 to answer questions 1 through 6. Which plant grew the tallest? 2. How many plants grew to be at least 6 cm tall? 3. Which plant grew the fastest in the first five days? 4. Which line represents plant 2? 5. After 10 days, how much had plant 3 grown? 6. How long did it take for plant 1 to grow 6 cm? Use the bar graph in Figure 4 to answer questions 7 through 11. 7. At birth, what is the average number of red blood cells per mm³ of blood? 8. What appears to happen to the number of red blood cells between birth and 2 months? 9. What happens to the number of red blood cells between the ages of 6 and 8 years? 10. Between what ages is a human likely to have 4.6 million red blood cells? 11. After 14 years of age, do males or females have a higher red blood cell count?

Part B. Constructing Graphs

Data Table 1 Breathing Rate of the Freshwater Sunfish

	The street restiwater Sums
Temperature (°C)	Breathing Rate (per minute)
10	15
15	25
18	30
20	38
23	60
25	57
27 24	25

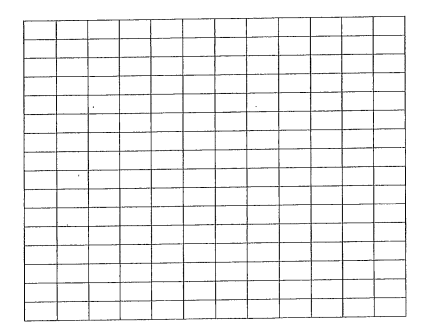
12. Use the grid below to construct a line graph for the information shown in Data Table 1.



Data Table 2 Average Rainfall in Willamette Valley

Month	Jan.	Feb.	Mar.	April	May	June	July	Aug.	Sept.	Oct.	Nov.	Dec.
Rainfall (mL)	15	21	28	24	16	8	2	1	2	3	5	10

13. Use the grid below to construct a bar graph for the information shown in Data Table 2.



	1.	How is a graph similar to a data table?
	2.	How is a line graph different from a bar graph?
	3.	Does a steep curve on a line graph indicate a rapid or a slow rate of change?
Criti	cal	Thinking and Application
		You are conducting an experiment to measure the gain in mass of a young mouse over a tenweek period. In constructing a graph to represent your data, which variable should you place along the x-axis and which variable should you place along the y-axis? Explain your answer.
g deserve of		
<u>()</u>		
	2.	What is an advantage of using multiple lines on a line graph? (See Figure 2.)
	3.	Why is it important to have all parts of a graph clearly labeled and drawn?
_	· 27 -	

Analysis and Conclusions

Name

The sites for this assignment are listed on the "Cells & Microscopes" page of the Kid Zone at http://sciencespot.net/.

Site #1: MOS Scanning Electronic Click the link for "How It Works"	on Microscope and then choose "Slide Show".
1. What does SEM mean?	
	oscopes work?
	microscope use to magnify images?
4. Why are the images black and w	hite?
5. How does the SEM work? Read	the captions and put the steps in order from 1 to 7.
	hits each spot on the sample, secondary electrons are knocked loose from its by a deflector and sent as signals to an amplifier.
The sample is placed i	nside the microscope's vacuum column through an air-tight door.
A set of scanning coils	s moves the focused beam back and forth across the specimen, row by row.
The final image is buil	t up from the number of electrons emitted from each spot on the sample.
	he column before the electron gun emits a beam of electrons, which travels of magnetic lenses designed to focus the electrons to a very fine spot.
The Scanning Electron of microorganisms.	Microscope reveals new levels of detail and complexity in the amazing world
SEM samples are coate	ed with a very thin layer of gold by a machine called a sputter coater.
6. Watch the animation if possible.	Write a paragraph to summarize what you saw.
o, main me management passasses	The state of the s
Chi HA TTA A 1 FEE A N. M.	
Click and drag the specimens on	the left side under the microscope to examine. Then identify the slides by n the right side of the screen. Write the results below.
#1	#6
#2 -	
#3	
₩Λ _	

Done with your worksheet? Visit the other sites listed on the Cells & Microscopes page!

#10 - _____

ACTIVITY #1

"HOW TO MAKE A WET MOUNT SLIDE"

In order to observe cells, you will have to become good at the technique of making a slide. This requires patience and careful handling of equipment. Take your time.

STEP 1

You will need a microscope slide and a coverslip.

STEP 2

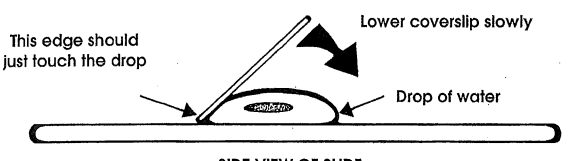
Put a drop of water on the slide.

STEP 3

Put the object into the drop of water. The object must be very thin. You will see the importance of this when you make a wet mount of onion cells.

STEP 4

Place the coverslip over the object by first placing one edge down, and then slowly lowering the other side so that you don't trap air bubbles. Air bubbles will look like discarded tires, and are actually quite interesting in appearance, but they will interfere with your view of the object you really want to see.



SIDE VIEW OF SLIDE



Whenever you make a slide of something during this semester, you should use the wet mount method. It is the very best way to get a clear view of the object, and it prevents the specimen from drying out.

Practice Staining Techniques

The parts (organelles) of a typical cell are mostly transparent. In a technique called staining, color is added to cell parts to help identify and distinguish them.



Procedure

- 1. Use forceps to remove a thin layer of onion skin, and place it in the center of a glass slide. Add a drop of water, and place a coverslip over the specimen.
- 2. Examine the onion skin with a light microscope. Draw what you see.

3. Place a drop of iodine stain along one edge of the coverslip. Touch a piece of paper towel to the opposite edge to draw the water. When the skin is stained, examine it with the microscope.

Analysis —

- 1. Describe how the stain affected the onion skin.
- 2. **Critical Thinking Analyzing Information** What is the advantage of using the paper to draw the stain across the field of view?

Human Epidermal Cells

Introduction

What do your skin cells look like? It is easy to remove some and look at them with a microscope.

Biological Concepts

· Cell structure

· Epidermis

Materiáls

Methylene blue stain, 1% aqueous

Clear tape, 1.0 cm × 1.0 cm

Dissecting needle

Forceps

Microscope

Microscope slide

Slide cover slip

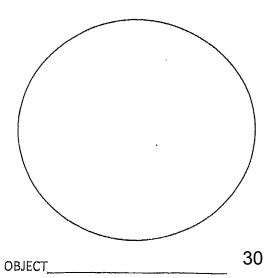
Soap/water

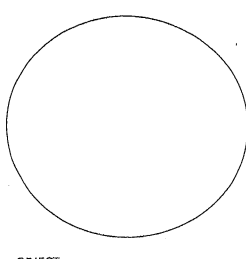
Safety Precautions

Methylene blue is a vital stain—it stains nearly everything, and it is difficult to remove. Prevention is the key when working with vital stains. Wear chemical-resistant gloves and avoid contact with eyes and skin. Wear safety glasses or chemical splash goggles whenever working with chemicals, heat or glassware in the lab.

Procedure

- 1. Wash the underside of a wrist that will be sampled for epidermal cells with soap and water.
- 2. Stick a clean piece of clear tape on the underside of the washed wrist.
- 3. Gently remove the piece of tape from the wrist being careful to avoid getting fingerprints on the tape. A forceps might help to remove the tape and avoid fingerprinting the tape.
- 4. Place the tape, sticky-side up, on a clean microscope slide.
- 5. Stain the top, sticky side of the tape with 2 or 3 drops of 1% methylene blue solution.
- 6. Use a dissecting needle to gently place a cover slip over the sticky tape. Lower the coverslip down onto the tape and then remove the dissecting needle. This should help prevent staining your fingers. Caution: Use methylene blue carefully. It will stain most items including skin, clothing, and table tops.
- 7. Examine the slide under a microscope. Look for cells with low power first, and then switch to high power for details,
- 8. Record your observations of epidermal cells by making drawings. Label your drawings with appropriate magnifications. Use your knowledge of the size of the microscopic field to estimate the size of the cells.





OBJECT____

Measuring with a Microscope

Pre-Lab Discussion

The microscope, developed more than three hundred years ago, is the basic tool of the biologist. The microscope enables biologists to investigate living things and objects that are too small to be seen with the unaided eye. The microscope is able to magnify these tiny specimens by means of lenses located in the eyepiece and objectives. The light microscope is also capable of revealing fine detail. This ability to reveal fine detail is known as resolving power. The type of microscope that you will be using throughout your study of biology is the compound light microscope.

Although it is interesting and informative to observe specimens under the microscope, it is often difficult to know the actual size of the object being observed. Magnification causes us to lose the idea of actual size. You cannot hold up a ruler to a paramecium or a plant cell while it is under the microscope. Therefore size must be measured indirectly—that is, it must be compared with the size of something you already know. The diameter of the microscope field seen through the eyepiece is a convenient standard to use. To measure objects under the microscope, a unit called the micrometer (μ m) is used. One micrometer equals 0.001 millimeter.

In this investigation, you will develop skill in using the compound light microscope. You will also learn how to estimate the sizes of objects under the microscope.

Problem

How is the compound microscope used to make measurements of microscopic specimens?

Materials (per group)

Microscope

Transparent metric ruler

Lens paper

Prepared slides

Safety &

Always handle the microscope with extreme care. You are responsible for its proper care and use. Use caution when handling glass slides as they can break easily and cut you. Note all safety alert symbols next to the steps in the Procedure and review the meanings of each symbol by referring to the symbol guide on page 10.

Procedure

- 1. Take a microscope from the storage area and place it about 10 centimeters from the edge of the laboratory table.
- 2. Carefully clean the eyepiece and objective lenses with lens paper.
- 3. Examine the markings on a metric ruler. Decide which marks indicate millimeter lengths. Place the ruler on the stage so that it covers half of the stage opening, as shown in Figure 1.
- **4.** Prepare your microscope for low-power observation of the ruler.
- 5. Look through the eyepiece. Focus on the edge of the ruler using the coarse adjustment. Adjust the position of the ruler so that the view in the low-power field is similar to Figure 2.
- 6. Place the center of one mark at the left side of the field of view. Make sure that the edge of the ruler is exactly across the center of the field. If the ruler sticks to your fingers, use the eraser end of a pencil to arrange it.

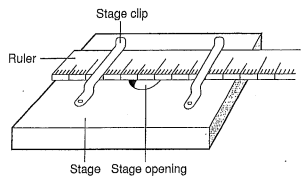


Figure 1

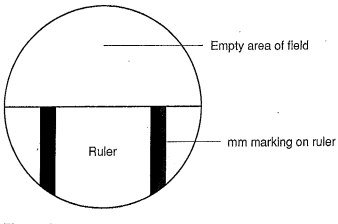


Figure 2

- 7. Note that 1 millimeter is the distance from the middle of one mark to the middle of the next mark. The diameter of the low-power field measures 1 millimeter plus a fraction of another. In Observations, record the measurement of the low-power field diameter in millimeters, expressing the length to the nearest tenth of a millimeter.
- 8. In Observations, record the measurement of the low-power field diameter in micrometers.
- 9. You cannot measure the diameter of the high-power field using the process you have just completed. Viewing a ruler under high power presents problems with light and focusing. Also, the high-power field diameter is less than 1 millimeter. But you can obtain the high-power field diameter indirectly. You know the low-power field diameter and the magnifying power of both objectives. Since the magnification of the objectives is inversely proportional to the field size, you can use this formula:

high-power field diameter = low-power field diameter x low-power magnification high-power magnification

In Observations, record the high-power field diameter in micrometers. Show your calculations.

10. Now that you know the diameter of your field size under both low and high power, you can estimate the sizes of the objects you view under the microscope by comparing them with the diameter of the field of vision. For example, if a tiny organism takes up approximately one-half of a field of view that is 1000 micrometers in diameter, then its size is about one-half of 1000 micrometers, or 500 micrometers.

11.	Obtain prepared slides of various organisms and practice estimating their lengths. Write the
	name of the organism or part you examine and its estimated size in micrometers in the Data
	Table.

12.	When you have finished	examining	the organisms	in step	11, return	your i	microscope	to t	the
	storage area.								

\cap	hs	0	۲ı	12	+i	^	n	c
	100	11	Г١	11	11	()	11	

1.	Measurement of the low-power field diameter =
2.	Measurement of the low-power field diameter = micrometers.
3.	Low-power magnification =

5. Use the formula shown in step 9 of Procedure to calculate the high-power field diameter. Show your calculations.

Data Table

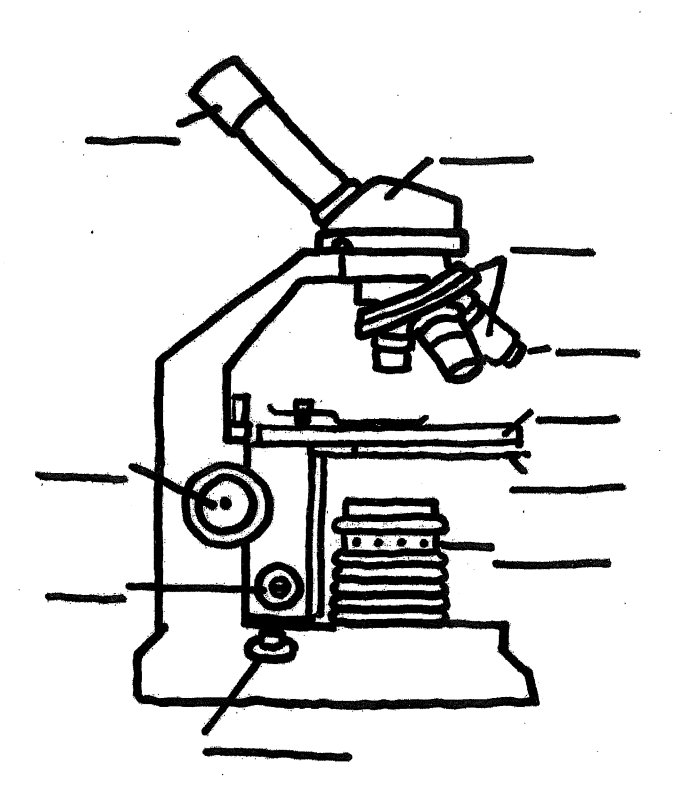
Name of Object	Measurement of Object (µm)

4. High-power magnification = _____

Analysis and Conclusions

2. How many micrometers are in 1 meter?		How many micrometers are in 1 millimeter?
· · · · · · · · · · · · · · · · · · ·	2.	How many micrometers are in 1 meter?
power magnification?	3.	What happens to the field of view when you change from low-power magnification to high-
		power magnification?

·
nes is the diameter of a field decreased when you change from low-power to high- lication?
Application
y 500 of a certain type of bacteria can fit across your low-power field of vision. pproximate size of 1 bacterium?
y 7 of a certain type of protist can fit across your high-power field of vision. What mate size of 1 protist?
be has a low-power magnification of 100X, a high-power magnification of 600X, wer field diameter of 1800 micrometers, what is the high-power field diameter in
t across a low-power field of view whose field diameter is 3000 micrometers, proximate size of each object?



BIO BASICS REVIEW
1. What is the dependant variable?
2. What is the independent variable?
Which is the dependant and independent variable in each of the following? Underline the independent, circle the dependant.
3. The number of holes in a student's jeans affects the number of detentions the student gets.
4. The amount of rain during the spring season determines the number of mosquitoes
5. A teacher's attitude is affected by the number of students in the class
6. The number of flowers on a cactus plant is related to the amount of rainfall.
Determine the variable that goes on each axis
7. The amount of fabric softener used in the laundry affects the grades that the student gets.
8. The type of sneakers a student wears determines the number of friends the person has.

9. The number	r of words a pers	on can text per mi	inute deter	mines tl	he numb	er of pe	ople that lil	ce them.	
10. The tempe	erature influences	the length of a do	og's hair.						
11. Make the	following graphs	•							•
Temp (° C)	# ofters					- T	1		
6	121								
45	150						 -		,
66	61	# otters			 		<u> </u>		
43	118								
51	100								
14	62								
				emp					
12. At 100x m	nagnification, wh	at does the object	ive have w	ritten o	n it?				

	What is the total magnification of the low power objective?
14.	What does the dissecting objective have written on it?
.5.	What objective is the high power?
б.	What steps do you need to take to put away the microscope?
7.	What direction does the object being viewed move when you move the stage to the right?
8.	How do you clean the lenses?
9.	What power objective must you use oil to see?
Э.	When making a wet mount slide, how do you get rid of air bubbles?
1.	If you put 23 drops of water to fill up a ml, how many drops would you need for 5 ml?
2.	If you put 105 drops of water to fill up a teaspoon, how many drops does each ml contain?
3.	Where do you read the fluid in a graduated cylinder?

24. Label the diagram of the microscope

Common SI	Common SI units				
Prefix	none	kilo-	centi-	milli-	
Factor	1 (base unit)	1,000	0.01	0.001	
Units used to describe volume	1 liter (L)	1 kiloliter (kL) = 1,000 L	1 centiliter (cL)= 0.01L	1 milliliter (mL) = 0.001 L	
Units used to describe length	1 meter (m)	1 kilometer (km) = 1,000 m	1 centimeter (cm) = 0.01 m	1 millimeter (mm) = 0.001 m	
Units used to describe mass	1 gram (g)	1 kilogram (kg) = 1,000 g	1 centigram (cg) = 0.01 g	1 milligram (mg) = 0.001 g	

Pre-Lab Questions

Where should you read the volume in a graduated cylinder?
What will you use to add water to the cylinder?
· .
What object will you find the volume of? What is this method called?
What is volumes will you be taking from the beakers of red, blue and yellow?
What are your predictions for this lab? Reading what colors you will be mixing, can you tell what the final colors will be, in order?.
·
•

•
Part A: Count your drops!
Take a guess - How many drops of water will it take to equal 1 milliliter? drops
Follow the directions to find the number of drops in 1 milliliter of water, then answer the questions. Yo will need a small graduated cylinder (25 ml), a beaker of water, and an eyedropper for this section. (1) Fill a small graduated cylinder with 10 ml of water. (2) Count the number of drops it takes to raise the water to 11 ml. Record the number in the chart. (3) Leave the water in the graduated cylinder and count the number of drops it takes to raise the water to 12 ml. Record the number in the chart. (4) Leave the water in the graduated cylinder and count the number of drops it takes to raise the water to 13 ml. Record the number in the chart. (5) Calculate your average and round to the nearest tenth.
#of/drops/to/1/mil // #lof drops to 12 mil // #lof drops to 13 mil // Average
Based on your average, how close were you to your guess?
Based on your average, how many drops would it take to make 1 liter?
Part B: Water Displacement
Follow the directions to find the volume of three marbles using water displacement. (1) Add 20 ml of water to a 100 ml graduated cylinder. Record this amount in the chart. (2) Add three marbles to the cylinder and measure the volume. Record this amount in the chart. (3) Find the difference between the two measurements and record in the chart. The difference between the two measurements will be the volume of the three marbles.
Volume of water Difference in volume Volume of 3 marbles before adding marbles marbles
Part C: Volume by Formula Use the formula to find the volume of the box. Measure to the nearest centimeter (no decimals) before calculating your answer.
Volume = length x width x heightx = -

Part D: Color Challenge

- 1. Obtain the following items from your teacher:
 - 3 beakers with colored water- 25 ml of each color (red, blue, and yellow)
 - 1 graduated cylinder (25 ml 50 ml)
 - 1 eyedropper
 - 6 test tubes labeled A, B, C, D, E, and F
- 2. Perform each step outlined below using accurate measurements.
 - (1) Measure 17 ml of RED water from the beaker and pour into test tube A.
 - (2) Measure 21 ml of YELLOW water from the beaker and pour into test tube C.
 - (3) Measure 22 ml of BLUE water from the beaker and pour into test tube E.
 - (4) Measure 5 ml of water from test tube A and pour it into test tube B.
 - (5) Measure 6 ml of water from test tube C and pour it into test tube D.
 - (6) Measure 8 ml of water from test tube E and pour it into test tube F.
 - (7) Measure 5 ml of water from test tube C and pour it into test tube B.
 - (8) Measure 2 ml of water from test tube A and pour it into test tube F.
 - (9) Measure 4 ml of water from test tube E and pour it into test tube D.
- 3. Complete the chart.

Test Tube	Color	Final/Amount (ml)
A		
В		
С		
D		
E		
F		

Pre-Lab Questions

1	What liquids will you be mixing?
2.	What is the formula of density?
3.	What will you be taking the temperature of?
4.	What must you do to the material you are measuring the temperature of?
5.	What are the SI units for a. Temperature b. Mass c. Density d. Volume
6.	What are your predictions for this lab? Be specific.

SI Units

OBJECTIVES

- Express measurements in SI units.
- · Read a thermometer.
- Measure liquid volume by using a graduated cylinder.
- Measure mass by using a balance.
- Determine the density (mass-to-volume ratio) of two liquids.

MATERIALS

- graduated cylinder, 100 mL
- cups, plastic, (2)
- thermometers, Celsius, alcohol-filled (2)
- ring stand or lamp support
- stopwatch or clock
- corn oil, 25 mL
- cup, clear plastic

- sand, light-colored, 75 mL
- sand, dark-colored, 75 mL
- gloves, heat-resistant
 - light source
 - balance
 - water, 25 mL
 - graph paper



Procedure

MEASURE SAND TEMPERATURE

- 1. Use the data table on the next page to record your results.
- 2. Put on safety goggles, gloves, and a lab apron. Using a graduated cylinder, measure 75 mL of light-colored sand. Pour the sand into one of the small plastic cups. Do the same thing with the dark-colored sand and another plastic cup.
- 3. Make sure the sand is level. You can do this by placing the cup on your desk and sliding it back and forth. Insert one thermometer into each cup.
- 4. Using a ring stand or lamp support, position the lamp approximately 9 cm from the top of the sand, as shown in the picture on page 20. Make sure that the lamp is evenly positioned between the two cups.

5. Before turning on the lamp, record the initial temperature of each cup of sand in the data table.

Sand Temperature

	Temperature	(degrees C)
Time (min)	Dark-colored sand	Light-colored sand
Start		
1		
2	·	
3		
4		
5		
6		
7		
8		
9		
10		

6. CAUTION: Wear heat-resistant gloves when handling the lamp. The lamp will get very hot and may burn you. Start the stopwatch when you turn on the lamp. The lamp will get hot and warm the sand. Check the temperature of the sand in each container every minute for 10 minutes. In your data table, write down the temperature of the sand after each minute.

COMPARE THE DENSITY OF OIL AND WATER

- 7. Use the data table on the next page to record the results from this section.
- 8. Label one clean plastic cup "Oil." Label a second cup "Water." Using a balance, measure the mass of each plastic cup. Record the weight in your data table.

	Density of Two L	iquids
a. Mass of en	npty oil cup	g
b. Mass of er	npty water cup	g
c. Mass of cu	ap and oil	g
d. Mass of cu	ip and water	g
e. Volume of	Coil	25 mL
f. Volume of	water	25 mL
	Calculating Actual	Mass
Oil	Item c – Item a =	g
Water	Item d – Item b =	g
g. Density of oil		g/mL
h. Density of water		g/mL

- 9. Put on an apron. Using a clean graduated cylinder, measure 25 mL of corn oil, and pour it into the plastic cup labeled "Oil." Using a balance, measure the mass of the plastic cup containing the corn oil. Record the mass in your data table.
- 10. Repeat step 9 with water instead of oil. Use the plastic cup labeled "Water."
- 11. To find the mass of the oil, subtract the mass of the empty cup from the mass of the cup and the oil together.
- 12. To find the density of the oil, divide the mass of the oil by the volume of the oil, as shown in the operation below.

Density of oil =
$$\frac{mass\ of\ oil}{volume\ of\ oil}$$
 = $\frac{g}{mL}$

- 13. Repeat steps 11 and 12 to find the mass and density of water.
- 14. Combine the oil and water in the clear cup. What happens?

lab.

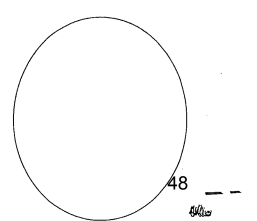
	·
15.	Clean up your materials. Put everything back where you found it. Wash your hands or use the GERM-X by door before leaving the

NAME	SCI#	POINTS:
SI UNITS LAB SHEET	·	
1. Use graph paper to graph the Remember to use the correct correctly! Staple behind this s	variables (use notes) and la	abel the graph
2. Based on your graph, how are	e color and heat absorption	related?
3. How might the color of the clear (Hint: Think of typical summer	othes that you wear affect er clothes.)	you on a sunny day?
4. In the second part of the lab, y	ou combined the oil and v	vater. How are your
observations related to the der	nsities of the liquids?	•
5. What could you infer about th floating in water?	e value for the density of i	ce if you observe it
6. How would your calculated de volume measurement on the gr	ensity values be affected if raduated cylinder?	you misread the
7. Pumice is a volcanic rock that you prove this density if you d (Hint: The density of water is	id not have a balance to w	0 g/cm ³ . How would eigh the pumice?

Pre-Lab Questions

What magnification is the dissecting lens? The low power? High power?
What will you be looking at under the microscope? Why is one object only looked at under low power?
List the steps for focusing under low power.
What is very important when focusing under high power?
What are your predictions for this lab? How many microns will the field of view be under low power?

Draw what you believe an human skin cell will look like under high power



Scale Drawings

OBJECTIVES

• Use the microscope to view prepared slides under low and high power

MATERIALS

Slides of tissues

Procedure:

- 1. You will take one of the prepared slides
- 2. Using your knowledge of the microscope, you will make 2 scale drawings, one under low power and one under high power
- 3. Use a clear ruler under the low poer objective only to get a scale, which you MUST include in your drawing under low power.

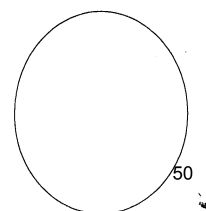
Questions

1. Complete 2 scale drawings below. Be sure to include a circle and labels as they are not provided for you in this lab.

Pre-Lab Questions

What m	agnification is the dissecting lens? The low power? High power?
What wi	ill you be making a wet mount of? What should you be careful of?
List the s	steps for focusing under low power.
What is v	very important when focusing under high power?
in the fiel	your predictions for this lab? How do you think the object will appear to ld of view when you move it right? Up? What do you think will be most to accomplish?.

Draw what you believe an onion cell will look like under high power



Using a Microscope

In almost every type of biological research, the microscope plays a fundamental role. Biologists use it to study the fine structures of cells and tissues, things that are too small to be seen with the unaided eye. The microscope used most often is the *light microscope*, which uses light to form an enlarged image of a specimen. A commonly used type of microscope is the *compound light microscope*. Compound light microscopes are used to view tiny living organisms as well as preserved cells mounted on glass (a *microscope slide*) and covered with a *coverslip*. This type of slide is prepared with water or some other liquid, such as a stain, and is called a *wet mount*.

Under the compound light microscope, most objects and microorganisms are observed in a drop of water. If you think of that drop of water as a pond and the objects and microorganisms as fish in the pond, you will begin to see why it is important to be able to focus at different depths. *Depth-of-field* focusing is always done under high power with the fine adjustment.

In this lab, you will practice using a compound light microscope. You will learn how to make a wet-mount slide and will observe several cell structures.

OBJECTIVES

- Show the proper use and care of a compound light microscope.
- Use the compound light microscope at low power and at high power.
- Prepare a wet-mount slide to examine under the microscope.
- Compare the movements of several images seen through a compound light microscope.

MATERIALS

- safety goggles, lab apron, protective gloves
- compound light microscope
- coverslip (5)
- dissecting needle or pencil
- forceps
- glass microscope slide (5)
- Elodea leaves
- lens paper

- medicine dropper
- methylene blue
- onion section
- paper towel
- prepared slide
- small plants, such as moss
- threads
- tweezers
- water



Procedure

PART 1: THE COMPOUND LIGHT MICROSCOPE

1. Complete Table 1 as you do Part 1.

FIGURE 1 THE COMPOUND LIGHT MICROSCOPE

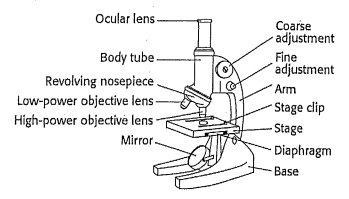


TABLE 1 THE PARTS OF A COMPOUND LIGHT MICROSCOPE

Microscope part	Function
Ocular lens	
(magnification:)	
Body tube	
Arm	
Stage	
Coarse adjustment	
Fine adjustment	
Lamp or mirror	
Revolving nosepiece	
Low-power objective lens	
(magnification:)	
High-power objective lens	
(magnification:)	
Diaphragm	
Base	

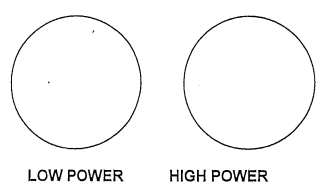
- 2. Carry a microscope to your lab table by holding the microscope arm with one hand and supporting the base with the other hand. CAUTION: A microscope is expensive and fragile. It is important to use it correctly to avoid damaging it and avoid breaking slides or destroying specimens. When you use a microscope, be sure it rests securely on your lab table away from the edge.
- 3. Locate each microscope part listed in **Table 1** and shown in **Figure 1**. Observe the magnification power (a number followed by an ×) of the ocular lenses and the low- and high-power objective lenses. Record these numbers in **Table 1**.
- 4. If your microscope has a built-in lamp, plug it in and turn it on to reflect light through the hole in the center of the stage.
- 5. Raise the objectives (or lower the stage) as far as possible by turning the coarse-adjustment knob. Secure a prepared slide to the stage, using the stage clips. Turn the low-power objective into position over the stage. While observing the stage from eye level, use the coarse-adjustment knob to position the objective as close to the slide as it will go without touching the slide.
- 6. Look through the ocular. Always keep both eyes open as you look into the eyepiece. Keeping both eyes open avoids eye strain. If the lens is dirty, ask your teacher to demonstrate the correct way to clean it. CAUTION: Never use anything other than lens paper to clean the lenses of the microscope. Focus with the coarse-adjustment knob only. CAUTION: Never focus by moving the objectives downward. You may run the objective into the slide and break the slide or damage the objective.
- 7. Complete focusing by slowly turning the fine-adjustment knob back and forth. When the object you are viewing is in focus and exactly in the middle of your field of vision, switch to high power. Use the fine-adjustment knob to refocus. CAUTION: Never use the coarse-adjustment knob at high power.

PART 2: MAKING A WET MOUNT

- 8. Use tweezers to strip a thin, transparent section of skin from the inner layer of a piece of onion.
- 9. Place the section of skin in the center of a clean, dry slide. With a medicine dropper, apply a drop of methylene blue stain to the skin. CAUTION:

 Glassware is fragile. Notify your teacher immediately of any broken glass.
- 10. Hold a coverslip at a 45° angle to the slide at the edge of the drop of methylene blue. Lower the coverslip slowly to avoid forming air bubbles. Under the microscope, air bubbles look round and have dark edges.

- 11. Place your wet mount onion cell slide on the microscope stage. Using the low-power objective, center and focus the microscope on the cells that make up the skin. Then switch to high power.
 - Make a drawing of what you see.



- What happens to the image of the cells as you go from low power to high power?
- 12. As you look through the eyepiece, slowly adjust the diaphragm to obtain the appropriate light for viewing.
 - What happens as you adjust the diaphragm?
- 13. As you look into the microscope, use your stage adaptor to move the slide to the right and then to the left.
 - What happens to the image as you move the slide to the right?
 - What happens to the image as you move the slide to the left?
- 14. Observe one cell carefully for several minutes under high power.
 - Locate a vacuole. How many are there?
 - What other cell structures do you see?
- 15. Obtain a sample of protists. Make a wet mount, and observe it under high power. Observe one cell carefully for several minutes.

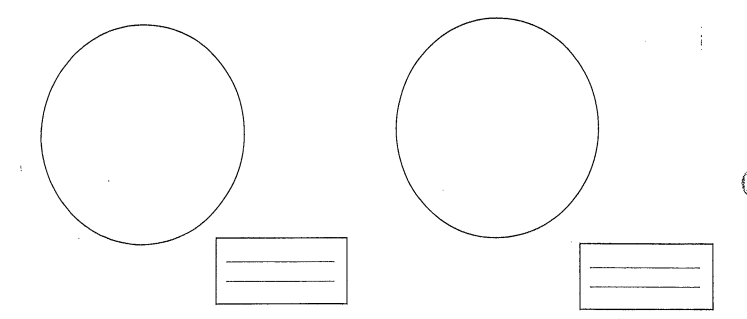
• Identify the structures you see.
What movement do you see?
PART 3: DEPTH-OF-FIELD FOCUSING
16. Make a wet mount slide of two threads by crossing the threads in the center of a clean glass microscope slide. Use a medicine dropper to add a drop of water. Add a coverslip to the slide.
17. Place your wet mount on the stage of the microscope. Under low power, adjust the slide on the microscope stage so that the point where the threads cross is in the center of your field of vision. Bring both threads into focus.
 18. Switch to high power. Using the fine adjustment, can you see both threads in focus at the same time? Why or why not? What can you infer about the depth-of-field and the objective used for viewing?
19. Slowly turn the fine-adjustment knob back and forth, and practice focusing on different parts of the two threads.
20. Dispose of your materials according to the instructions from your teacher. Clean up your work area, and wash your hands before leaving the lab.

	uestions for Microscope Lab What does the magnification number on the ocular lens mean?
2	Calculate the total magnification of your compound light microscope at low power and at high power. (Multiply the ocular (eyepiece) magnification by th objective magnification.) Show the calculation.
3.	Is the largest field of view seen under high power or low power?
4.	Why is it necessary to be able to focus at different depths?
5.	When making a wet mount, why must you always use a coverslip?
6. yo	When the slide is moved to the right, the object through the eyepiece appears to move in which direction? What occurs when the slide is moved away from 1?
	If a microorganism was moving from right to left across your field of view under a compound light microscope, which way would you move the slide to keep the microorganism in view? Why?
	Which thread from the prepared slide was on the bottom? What color was on

9. In addition to compound light microscopes, there are more powerful ones that scientists use. Research (use book **56**internet) to determine the differences in

what they use to image an object and the amount of magnification for the following microscopes:

- a. scanning electron microscope
- b. transmission electron microscope
- c. scanning tunneling electron microscope
- 10. Draw the onion cell correctly below under low AND high power. Make sure to use proper labeling and drawing techniques as discussed in class!!



Ch 1 Crossword/Flashcards-complete the crossword, then make a

flashcard (term on one side, definition on the back) for all 10 terms

	1	2													
			3	I			4	1							
5					6]	•					
											 			 	 T1
							7		<u></u>						
-	-														
8															
9															

Across

- 1. in biology, the smallest unit that can perform all life processes
- 5. the maintenance of a constant internal state in a changing environment
- 7. the process of obtaining information by using the senses; the information obtained by using the senses
- 8. Le Système International d'Unités, or the International System of Units, which is the measurement system that is accepted worldwide
- 9. a habit of mind in which a person questions the validity of accepted ideas

Down

- 2. a procedure that is carried out under controlled conditions to discover, demonstrate, or test a fact, theory, or general truth
- 3. group in an experiment, a group that serves as a standard of comparison with another group to which control group is identical except for one factor
- 4. the scientific study of living organisms and their interactions with the environment
- 5. a testable idea or explanation that leads to scientific investigation
- 6. a system of ideas that explains many related observations and is supported by a large body of evidence acquired through scientific investigation 58

CHAPTER 1 VOCAB

Skepticism	objectively	:		
Observation			,	
Hypothesis				
Experiment				
control group				
theory	· ·	****		
SI				
Biology	T			
cell		***************************************		
homeostasis				
universal laws				
correlation data				
bias				
TA AT 1 X 1 11 11 11 11 11 11 11 11 11 11 11				
Ones			• .	
	59 L			

CHAPTER 1 BIOLOGY AND YOU

I.	SCIENTIFIC THOUGHT involves making observations, using evidence to draw
conc.	lusions, being skeptical about ideas, and being open to change when new discoveries
are n	nade.
Π.	UNIVERSAL LAWS-Science is governed by truths that are valid everywhere in
the u	niverse. These truths are called
Ш.	SCIENCE AND ETHICS -Scientific experimentation and discovery can have
serio	us ethical implications. Because of this, scientific investigations require ethical
behar	vior are a system of moral principles and values.
Scien	ntists performing investigations must report only accurate data, must allow peers to
revie	w their work, and must behave ethically with the people involved in their
inves	etigations.
IV.	WHY DO YOU NEED SCIENCE? An understanding of science can help you
take 1	better care of your health, be a wiser consumer, and become a better-informed
citize	en.
	A. The same critical thinking process that scientists use is a tool that you can use in your
	everyday life- ex deciding which route to work has the least amount of traffic
	B. You can use what you learn to increase the quality of your physical life- ex what is the best acne medication, which vitamins help you live better, etc
V.	C. New technologies are around the corner. Understanding biology and science will help you make informed decisions- ex new drugs for obesity, removing the need for sleep. nanotechnology SCIENTIFIC EXPERIMENTS-Scientists conduct controlled experiments or
	rm studies in order to test a
Ç	A. An experiment is a procedure that is carried out under controlled conditions to test
	a hypothesis.
	B. There are often cases in which experiments are not possible or not ethical. In these
	cases, researchers perform studies or use correlation data (statistics gathered from
	subjects that show a relationship)
	C. Scientists verify their by conducting their experiments
	many times and by checking to see if other scientists have found similar results.
	D. Every person has his or her own point of view. A point of view is called a
	E. Scientists try to prevent bias from affecting their work, but bias can still influence
	an experiment. Sources of funding, personal involvement in a product, and other conflicts
	of interest can affect an experiment.

~	F.			text and think critically about
	scient	ific theories. Ex- diet mirac	les	· •
Y	G.		·	othesis is that a hypothesis is a
1	-			onditions and a theory is a general
	explar	nation for a broad range of	data that is consistently	proven correct by new studies.
		•		•
VI	THE	STUDY OF LIFE-Biolo	ogy is the scientific st	udy of living organisms and
the	ir interac	tions with the environme	nt. Some of the branc	hes of biology are
	•	biochemistry,	•	microbiology,
	•	ecology,	•	botany,
		cell biology,	•	zoology,
		genetics,	•	physiology
		evolutionary theory,		1 7 20
		' crosscounty through	•	
Y 77Y	DD ODEI			
VII.	PROPE	RTIES OF LIFE-The six	c properties of life ar	e:
1.		i. All living things are made	of one or more cells.	
	•	ii. A cell is the smallest unit c		
2.		i All living arganisms must		existentment in order to function manuals.
÷ •		ii. The maintenance of a stabl	e internal environment in sp	nvironment in order to function properly. pite of changes in the external environment
•		is called homeostasis.		
3.		i. Living organisms carry out		ns in order to obtain energy.
		ii. The sum of all the chemica	l reactions carried out in an	organism is called metabolism.
		iii. Almost all of the energy us	ed by living things original	ly comes from the sun.
			•	
4.				
•		i. In addition to maintaining a external environment.	a stable internal environmen	nt, living organisms respond to their
		ii. Can you think of a way tha		ur environment today • Reproduction
		 Most living things can repr their own kind from one ge 		process by which organisms make more of
5.				
,		i. When an organism reprodu	ices, it passes on its own tra	aits to its offspring in a process called
		heredity. ii. Inherited characteristics ch	ange over generations. This	s process is called evolution
6.		III, IIIIoiitod olidiaatotiotios oli		s process is carred evolution.
1		i. All living organisms grow.		
		ii. As organisms grow, many	change. This process is call	ed development. nent refers to change in a single individual
		during that individual's life		mont rotors to change in a strigle marylanar

CHAPTER 1	SEC 1	Due Date	
Why is skepticis	sm important in science?		
•			
	i		·
Describe four w	ays to practice scientific thought.		
			
	•		
.What is a unive	rsal law?		
	I I avvo		
. Identify two uni	/ersai laws.		
			. <u> </u>
Give three exa	mples of ethical scientific behavior.		
	•	•	•
	·		
			• •
Think about sor	me decisions you make every day. Give to	WO	
examples of ho	w you can use scientific thought to help y	ou make good decisior/	าร.
	•		
	•		



CHAP	TER 1 SEC 2 Due Date	
1.	How do most scientific investigations begin?	
2.	What is the difference between a dependent variable and an independent variable?	1.
3.	How is a theory different from a hypothesis?	
4.	<u>Underline</u> the independent variable and <u>circle</u> the dependent variable in the follows. Telling your mother that she is a good cook increases the hour of your curfew.	ing.
5.	Washing the dishes for your mother increases the amount of money she gives you or	n
6.	Coaches with more years of experience will have a higher percentage of wins.	
7.	The amount of food that a bird eats is affected by the temperature.	
8.	Eating chocolate affects the number of zits you get.	

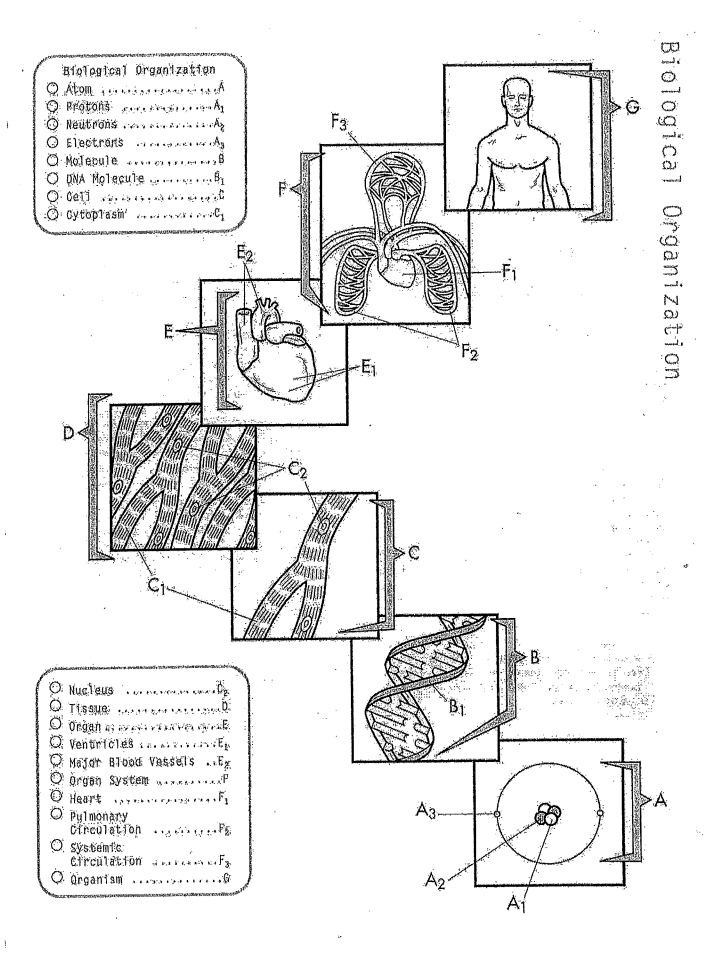
CHAPTER 1 REVIEW QUESTIONS

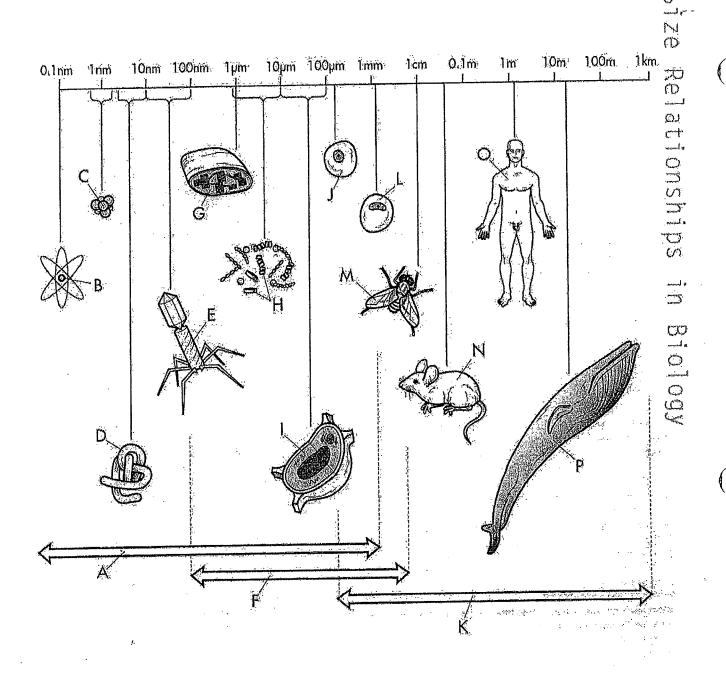
- 1. What is bias? Why do scientists not use this to support ideas?
- 2. What is skepticism? Why would making a new discovery be a result of skepticism?
- 3. What is a universal law? What branches of science do universal laws mostly apply to?
- 4. Which are some examples of unethical behavior in scientific investigations?
- 5. How can an understanding of science help you live a better life?
- 6. Noticing that your heart rate AND respiration increase when you exercise is an example of what? What part of the scientific process is noticing changes? What about writing down your heart rate and respiration?
 - 7. What is a hypothesis?.
 - 8. Most typically, what is the order in which the steps of scientific investigations are applied?
 - 9. If experiments are not possible or ethical, scientists can do what?
 - 10. How are scientific hypotheses tested?
 - 11. What is the definition of an experiment?
 - 12. What are the independent and dependant variable? (definitions)
 - 13. What is a general explanation for a broad range of data called?
 - 14. Is a scientific theory always correct? When can they be revised (updated or changed)?
 - 15. How do scientists build a theory? What do they use to support the theories?
 - 16. What number is the metric system based on? (what are the powers)?
 - 17. How many kilometers is one meter equal to?
 - 18. A specialized tool used to magnify organisms so that they can be observed is a
 - 19. What is sterile technique? Give an example of how sterile techniques prevents contamination
 - 20. Know your safety procedures.
 - 21. What is biology the study of?
 - 22. What are the six properties of life?
 - 23. As a characteristic of all living things, homeostasis relates most directly to what biological themes?

65

- 24. All living things maintain a balance within their cells and with the environment through the process called what?
- 25. What is the process by which organisms make more of their own kind?
- 26. Children tend to resemble their parents due to what trait of living things?
- 27. When sunflowers turn their flowers to follow the sun, or you wake up from your alarm clock going off, this is an example of which property of life?

ß





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	Stre Retationships in Biology	
O Electron Microscope Range O Atom O Small Molecula:	O Light Microscope Range O Chioropiast O Bacteria O Plant/Animal CellI	O Unaided Eye Bange K O Frog Egg Cell O Insect O Rodent O Human
O Virus E	O Human Egg Cell	Whale

Using the Scientific Process

Scientific Process

- Collecting observations
- Forming hypothesis
- Making predictions
- Verifying predictions
- Performing control experiments
- Forming a theory

To show how each stage of a scientific investigation leads logically to the next, perform the following exercise and identify each stage.

- (1) Collecting observations
 - (a) Measure (for one-minute) your resting respiratory & pulse rate
 - (b) Record the number of times you breathe during one minute period
 - (c) Record the number of times your heart beats during one minute period

Resting Respiratory Rate	Resting Pulse Rate	
(2) Formulate a hypothesis	S:	
Have will awayoing offact	vane easting easniesta	ry & pulse (heart) rate?

(3) Making Predictions:

Note the number of breaths & pulses you think you will take in the same period of time after you have jogged in place for one minute

Predicted Respiratory	Predicted Pulse
Rate	Rate

(4)	Verifying	predictions:
-----	-----------	--------------

Exercise by jogging in place for one minute and then immediately record your respiratory & pulse rate after exercise.

Respiratory Rate After Exercise	Pulse Rate After Exercise

(3) Ferrorming control experiments
How do you know that exercise was the factor that affected your respiratory rate and pulse rate.
(6) Formulate a theory
Suggest a connection between exercise and your respiratory & pulse rates.

Pre-Lab Questions

1.	How will you measure head circumference?
2.	How will you measure running speed?
3.	Who's information will you be recording?
4.	How will you measure height?
5.	What are the SI units for a. Shoe size
6.	b. time c. height d. weight What are your predictions for this lab? Be specific.

LAB Graphing

OBJECTIVES

- to use observation to record data
- to use data to create a graph that best demonstrates a correlation

MATERIALS

- graph paper
- string
- ruler

Procedure:

- 1. You will use a string to measure height and head circumference. Mark the string with a marker or pen instead of cutting it and then measure it using a yardstick.
- 2.To determine how high someone can jump, measure where the tip of their hand is on a wall (use masking tape), then have them place a different piece of tape on the wall as high as they can put it while jumping. The distance between the 2 pieces of tape is the height they jumped.
- 3.Use 3 attempts for jumping, breath holding, and running. Take the best of the three.
- 4. Use the materials provided to record the variables of the following:
 - a. Does someone's height determine how high they can jump?
 - b. The effect a person's age has on their weight. (Use kilograms)
 - c. Does a student's head circumference effect the length of time a person can hold their breath? Use class data.
 - d. Does a person's shop effect the number of jumping jacks they can complete in 1 minute?
 - e. Does a person's shoe size influence how fast a person can run 100 meters?
 - f. The favorite types of ice \overline{d} ram in the class. (yes this is a graph)

- 5. Unless otherwise specified, you will use the people in your lab table. This means there will be a minimum of four people.
- 6. If there are not enough people in your group, I will combine groups.
- 7. You must use SI units except for shoe size, which is American shoe size. We will run the 100 meters as a group and you will be required to remember your own time.
- 8. Create a different graph for each situation. They are not all line graphs. Use what you know about graphs to chose the best type of graph for the information
- 9. The x and y axes (meaning the independent and dependant variables) must be correct, the labels must be correct, an appropriate title must be chosen, a ruler must be used to draw the graphs and of course graph paper must be used. Your name must be on each page and they must be stapled. Each person must hand in their own set of graphs.

Chapter 3 Word Search/ Flashcards Homework- find all 30 words in the letters below whose definitions follow the puzzle. Then create flashcards for the terms

(30). Remember to put the definition on the opposite side as the term

W	N	Q	0	N	S	P	0	U	D	E	В	E	В	M	E	T	N	E	X
X	C	0	H	E	s	I	0	N	N	I	L	U	0	A	V	V	U	N	N
P	K	F	R	P	X	H	I	Z	G	U	P	\mathbf{T}	F	В	S	${f T}$	C	E	I
T	E	V	I	T	C	A	Y	P	C	S	A	I	E	F	I	E	L	R	E
A	0	D	H	0	C	M	D	E	V	A	E	M	L	S	E	В	E	G	T
C	Q	D	I	N	E	E	L	J	C	L	T	C	U	I	J	R	0	Y	0
L	A	X	X	I	P	0	L	\mathbf{T}	I	C	\mathbf{T}	A	N	\mathbf{T}	M	C	T	J	R
R	C	R	Y	M	M	F	I	E	I	A	N	D	M	E	Z	L	I	L	P
P	L	A	В	A	В	V	Z	E	Z	M	R	J	A	W	L	E	D	F	Δ
S	Z	X	S	0	A	A	L	L	В	J	N	V	Y	U	F	A	E	C	D
P	C	Y	N	T	H	C	D	U	Q	\mathbf{Z}	J	L	H	J	S	I	V	L	Z
R	T	M	I	В	U	Y	Z	H	R	I	K	Q	L	U	0	S	W	S	F
D	W	0	Z	N	A	A	D	E	E	K	В	X	G	D	L	I	U	L	W
Y	N	Q	F	A	E	T	A	R	T	S	В	U	S	N	U	Q	Q	D	P
Y	0	В	Y	S	F	C	Q	A	A	M	I	C	G	U	T	H	J	L	N
W	V	C	U	M	T	X	E	Y	I	T	U	0	M	0	I	M	Q	K	N
Y	Q	K	R	A	J	A	J	L	F	J	E	H	N	P	0	G	H	T	F
F	I	C	N	A	C	I	D	G	J	I	В	T	Z	M	N	A	I	N	V
T	F	T	I	V	R	D	W	P	H	Q	K	F	A	0	Z	0	N	E	T
T	N	E	M	E	L	E	P	R	0	D	U	C	T	C	N	D	L	R	M

	the smallest unit of an element that maintains the chemical
	properties of that element
	a substance that can not be broken down into simpler substances by
	chemical means- ex Carbon, hydrogen
	an electron that is found in the outermost shell of an atom and
	determines the atom's chemical properties
	a substance that is made of atoms of two or more elements joined by
·	chemical bonds
	a group of atoms that are held together by chemical forces

	an atom or molecule that has gained or lost one or more electrons and has a net positive or negative charge
)	the force that holds molecules of a single material together the attractive force between 2 different substances that touch each other
	a homogeneous mixture with 2 or more substances uniformly dispersed
	any substance that increases hydronium (hydrogen ions H+)
	any substance that increases hydroxide ions (OH-) a scale that shows acidity or alkalinity (basicness). A logarithmic scale.
	a substance that acts as an acid and base and stabilizes a solution's
	pH a class of molecules that has carbon, hydrogen and oxygen. Includes sugars, starches and fiber
	long hydrocarbon chains that includes fats, waxes and steroids
,	long chains of amino acids. Main component of everything in cells
	a class of molecules that contain a carboxyl group and amino group
Northern American Control of Cont	long chains of nucleotides. Includes DNA and RNA contains a sugar, nitrogenous base and phosphate group. In chains, makes DNA and RNA
	deoxyribonucleic acid, determines hereditary information ribonucleic acid, also carries genetic information, but uses info to make proteins
	adenosine trisphosphate, the energy currency of ALL cells
	capacity to do work
	a substance that is part of a chemical reaction
	a substance that is formed from a chemical reaction the minimum amount of energy that is needed to begin a chemical reaction a molecule, usually made of protein, that helps chemical reactions happen in cells
	— the reactant (beginning substance) catalyzed by an enzyme
	the site on an enzyme where the reaction takes place

Chapter 3 Chemistry of Life

	Il matter is made up of atoms en down by chemical means.	s. An atom is the small	est unit of matter that
A. Matte	er is anything that has	and t	akes up space.
uncharge	nucleus of an atom is made up d neutrons. Negatively charg ne nucleus in a large region ca	ed electrons have very	little mass and move
	lement is a substance made up For example, each atom of th	~	
	ns of an element may have dif		trons. These atoms
	AL BONDS -Chemical bond ome stable when they have ei		
A. Electi	rons in the outermost level, o	r shell, are called	·
	s tend to combine with each hell. When atoms combine, a	_	
C. When compound	a atoms of different elements d is a substance made of the b	combine, aonded atoms of two or	forms. A r more elements.
D. Types	s of bonds-		
1	- sharing	valence electrons form	s a covalent bond.
	a) Acovalent bonds.	is a group of atoms l	neld together by
	b) A water molecule, H ₂ C), forms when an oxyg	en atom forms
	covalent bonds with two h		
ator lost	-Atoms can a ning electrons, resulting in a mor group of atoms that has a electrons. The attractive for ic bond.	positive or negative ch an electric charge beca	arge. An ion is an use it has gained or

III. POLARITY – some bonds may have charges that are not distributed equally. Molecules with partial charges on opposite ends are said to be
A. In some covalent bonds, the shared electrons are attracted more strongly to one atom than to the other. As a result, one end of the molecule has a partial negative charge, while the opposite end has a partial positive charge.
B. The partially charged ends of polar molecules attract opposite charges. Because of this behavior, polar molecules can dissolve other polar molecules and ionic compounds.
C. Nonpolar substances, such as, grease, and, do not dissolve well in water.
D. When bonded to an oxygen, nitrogen, or fluorine atom, a hydrogen atom has a partial charge nearly as great as a proton's charge. It attracts the negative pole of other nearby molecules. This attraction, called a, is stronger than attractions between other molecules, but not as strong as covalent bonds.
IV. PROPERTIES OF WATER -Most of the unique properties of water result because water molecules form hydrogen bonds with each other.
A. When water freezes, the crystal structure formed due to hydrogen bonding makes ice than liquid water.
B. Water can absorb a large amount of heat without changing temperature. This property can help organisms maintain a constant internal temperature.
C. The attraction of particles of the same substance, such as water, is called Cohesion keeps water from evaporating easily; thus, water is a liquid at ordinary temperatures.
D. Water molecules also stick to other polar molecules. This attraction between particles of different substances is called
V. SOLUTIONS - A solution is a mixture in which ions or molecules of one or more substances are evenly distributed in another substance.
A. Many substances are transported throughout living things as solutions of water. Dissolved substances can move more easily within and between cells.
B. Some water molecules break apart to form(H+) and(OH-) ions. In pure water, hydronium and hydroxide ions are present in equal numbers.

vi.	ACIDS/BASES- Acids and bases are compounds that change the	to darance of these
	A. Acids are compounds that form extra when dissolved in water.	(H+) ions
	B. Bases are compounds that form extra when dissolved in water.	(OH-) ions
	C. When acids and bases are mixed, the extra hydronium and hyreact to form water.	ydroxide ions
VII. j	\mathbf{pH} is a measure of how acidic or basic a solution is.	
	A. Each one-point increase in pH represents ahydronium ion concentration. (logorathmic scale)	decrease in
	B. Pure water has a pH of Acidic solutions have a pH7.	7, and
	C. The pH of solutions in living things must be stable. For a stamaintained, the solutions in living things contain buffers.	ble pH to be
·	D. A is a substance that reacts to preven a solution.	nt pH changes in
VIII. called	BUILDING BLOCKS OF CELLS – biomolecules contain lorganic). They include carbohydrates, proteins, lipids and nuclei	•
	A Carbohydrates are molecules A sugar contains carbon, hydrogen, and oxygen in a ratio of	
	1. Carbohydrates are a major source of energy	
,	2. Chitin and cellulose are complex carbohydrates that pro	vide support.
	a) is found in the shell the cell walls of mushrooms.	s of insects and
	b) is found in the cell	walls of plants.
	B Lipids are another class of which includes fats, phospholipids, steroids, and waxes.	of biomolecules,
	1. Lipids consist of chains of carbon atoms bonded to each hydrogen atoms. This structure makes lipids	
	2. The main purpose of is to store energy. I energy even more efficiently the carbohydrates.	ats can store

	3. The cell's boundary(cell membrane) is made of
	The structure of cell membranes depends on how this molecule interacts with water.
(CProteins are chains of amino
8	Proteins may be involved in structure, support, movement, communication,
	ransportation, and carrying out chemical reactions.
	1. A protein is a molecule made up of amino acids, building blocks that link to form proteins.
	a) Every amino acid has an group and a group. Units of amino acids can form links called peptide bonds.
	b) The group gives an amino acid its unique properties different amino acids are found in proteins.
	2. For each type of protein, there are different levels of structure
	a) amino acids are arranged in a specific order, the protein's primary structure. 1
	b) The interactions of the various side groups may form coils and folds, the protein's secondary structure. 2
	c) The overall shape of a single chain of amino acids is the protein's tertiary structure. 3
	d) The quaternary structure is the overall shape that results from combining the chains to form proteins. 4
) u 	- A nucleic acid is a long chain of nucleotide nits. A nucleotide is a molecule made up of three parts: a, a, and a group.
	1. Nucleotides of deoxyribonucleic acid, or, contain the sugar deoxyribose. DNA molecules act as "instructions" for the processes of an organism's life
	2. Nucleotides of ribonucleic acid, or, contain the sugar ribose. RNA also interacts with DNA to help decode the information.
	3. Adenosine triphosphate, or, is a nucleotide that has three phosphate groups and supplies energy to cells. Energy is released in the reaction that breaks off the third phosphate group.

		change occurs when only the form or shape of the
n	natter (changes.
	. A_ ıbstan	change occurs when a substance changes into a different ce.
C8	alled t	Matter is neither created nor destroyed in any change. This observation is ne Every change in matter a change in energy.
		rgy may change from one form to another, but the total amount of energy t change. This observation is called the
		CAL REACTIONS -Chemical reactions can only occur when the gy is available and the correct atoms are aligned.
		nging a substance requires a chemical reaction. During this process, etween atoms are broken, and new ones are formed.
В	. A_	is a substance that is changed in a chemical reaction.
C.	A_	is a new substance that is formed.
en ch	ergy o emica	mical reactions can only occur under the right conditions. The activation of a reaction is the kinetic energy required to start a l reaction. Even if enough energy is available, the product still may not ne correct atoms must be brought together in the proper orientation.
		ICAL REACTIONS - By assisting in necessary biochemical reactions, organisms maintain homeostasis.
		ving things, chemical reactions occur between large, complex cules. Many of these reactions require large activation energies.
В.	An_	is a molecule that increases the speed of reactions.
		Enzymes hold molecules close together and in the correct orientation. An zyme lowers the activation energy of a reaction.
		Each enzyme has an, the region where e reaction takes place.
		The shape of the active site determines which reactants, or substrates, ill bind to it. Each different enzyme acts only on specific substrates.

4. Most enzymes need a certain range of _____ and

CHAPTER 3	SEC 1	DUE DATE .
i. How are atoms and e	elements related?	
2. Fill in the blank space	es in the table below.	
·		
•		
Type of particle	Location within an atom	Charge
	outside the nucleus	
Proton		
	in the nucleus	O (noutral)
	in the nucleus	0 (neutral)
. Why do atoms form o	chemical bonds?	
	,	
. How is a covalent bo	nd different from an ionic bond?	
•		
-		,
	, t t t t t t t t t t t t t t t t t t t	
Market and the second s		
s. What is a hydrogen b	ond?	
	,	
		·

				•	
·					
. Why does sodi	um have a po	ositive charge	when it is in s	olution?.	
			-		

Bellninger:DayMIT W:Th:F:Date	ess Ouestion	
Answer:	The second secon	
The state of the s		
The Control of the Co		

		•	
CHAPTER 3	SEC2	DUE DATE	<u>.</u>
1. A student emp		ass. The student observes that small droplets o	
water,remain stud	ck to the glass. Which tw	o properties of water explain the student's	
observation?			
· · · · · · · · · · · · · · · · · · ·			
2. Oceans and oth	ner bodies of water warm	n up more slowly than air or land.	
Describe how the	hydrogen bonds betwee	en water molecules cause this effect.	
		· · · · · · · · · · · · · · · · · · ·	
		water, some of the CO ₂ molecules react with v	vate
This forms carbor	nate ions and hydronium	ions. Will a solution of CO2 in water be acidic,	
basic, or neutral?	Explain your answer.		
•		n living things contain buffers?	
	, ,		
	•		

Bellringeri DayMiTeW, Tihlif, Date: <u>Austropassion</u> <u>Sandara (Caranter Caranter Cara</u>

HAPTER 3	SEC 3	DUE DATE	·
. What are biomole	cules?		
. Fill in the spaces i	in the table below.		·
Type of blomolec	cule: What are the third this type	he building blocks of	main function of f biomolecule?
arbohydrate	在第二十二十二十八十二十八十二十二十二十二十二十二十二十二十二十二十二十二十二十二		
	atoms	arbon and hydrogen	
rotein			1
		·	,
	Nucleotides		·
What is the differe	nce between a nucleic	acid and a nucleotide?	
		<u>:</u>	
	,		
			,
· · · · · · · · · · · · · · · · · · ·		,	
Inger:DayM:TW:Th.F.D	ate <u>s</u> Questio		
inger: DayM:T-W-Th F. Di ing p	ate Question		
	ate Question		

- 1. Where do living things get the energy they need?
- 2. How is a physical change different from a chemical change?
- 3. Give two conditions that must be met for a chemical reaction to occur.
- 4. Identify the products and the reactants in the chemical reaction shown below. Write only the chemical formulas for the products and reactants.

$$CO_2 + H_2O \rightarrow C_6H_{12}O_6 + O_2$$

- 5. Why are enzymes important to living things?
- 6. What is the relationship between an active site and a substrate?
- 7. Why may an enzyme not work properly if temperature or pH changes?

CHAPTER 3 REVIEW

- 1) Atoms are composed of what?
- 2) What are ionic bonds? How do they form?
- 3) What is an element?
- 4) What is a molecule?
- 5) What is a covalent bond? How does it form?
- 6) Why don't oil and water mix?
- 7) What types of bonds share electrons? Donate electrons?
- 8) What element is contained in all biomolecules? What are the types of biomolecules?
- 9) What is a polar molecule? Why is water a polar molecule? What does the polarity do?
- 10) What is a non-polar molecule? What charges does a non-polar molecule have? What is a polar molecule? What charges do a polar molecule have?
- 11) What so nonpolar molecules look like? Are the ends charged? How do non-polar molecules behave?
- 12) What is an electron? Where is an electron cloud found? Where do electrons stay? What is the valence shell?
- 13) What is the smallest particle of matter that retains the properties of the element? What is the difference between an atom, element and molecule?
- 14) What are polysaccharides, sugars, chitin and cellulose?
- 15) What are the unique properties of water?

- 16) If the electrons in the valence shell are shared, what type of bond is it? What does this do to the stability of the molecule?
- 17) What are hydronium ions? What do excess hydronium ions do to the pH of the substance?

What are hydroxide ions? What do excess hydroxide ions do to the pH of the substance?

- 18) What types of molecules are classified as carbohydrates?
- 19) What types of molecules are classified as lipids?
- 20) What type of molecule is this?

21) What type of molecule is this?

- 22) How are lipids and carbohydrates similar? What do they have in common in structure?
- 23) What are the four levels of protein organization? What are the characteristics of each level?
- 24) What is cohesion? What is adhesion? What properties of water do each of these influence?
- 25) What are the attractions between water molecules called? How do they form? Why are they important?

26) Give 3 examples of lipids 27) Why is each amino acid unique? What portion of the structure is responsible for this uniqueness? 28) What is the substrate of an enzyme? What is the reactant of an enzyme? What is the active site of an enzyme? 29) What is matter composed of? 30) What are the two types of nucleic acids? 31) A pH less than seven means that the substance is what? 32) How do you form an electron bond? Are the electrons shared or donated? 33) What are the parts of a DNA molecule? 34) What is the link between a carboxyl group of an amino acid and the amino group of another amino acid called? 35) How do enzymes make reactions proceed? What do they do to the activation energy? 36) Where are long chains of amino acids found? 37) What molecule does an enzyme act on? 38) If a substance has a pH greater than 7 mean the substance is what?

40) What is the force that allows water to climb up a glass tube called? What is the type of

39) How does ATP store energy?

bonding that is responsible for it called?

- 41) What are the charges of a
 a. neutron
 b. proton
 c. electron
 42) Where can each subatomic particle be found?

Pre-Lab Questions

What i	is pH? What foes the p represent? What does the H represent?
What v	vill you be using to test solid surfaces?
What p	H is most acidic? What pH is most basic?
What is Why?	a buffer? Name a common buffer? What type of water will you be using?
	e your predictions for this lab? What in your shop will be acidic (3 items). ill be basic (3 items). What do you believe will be neutral?
-	
-	

pH of Shop Materials Lab

Materials

- pH test strips
- pH color change guide

- Q tips
- Distilled water

Procedures:

- 1. You will be exploring your shops to determine the pH of various materials commonly used in your area of interest. You will be working in groups of two. Gloves and eye protection must be worn during this lab!
- 2. You will determine the pH of substances by using the pH test papers. These papers are designed to test liquids, but you can test pH of solids, although the results are not as accurate. The papers are embedded with a chemical that causes a color change in the presence of hydrogen and hydronium ions.
- 3. The liquids you test should sampled using the following method. Immerse the cotton portion of the Qtip into the liquid to be tested. The qtip should then be wiped on the test paper. Do not immerse the test strip in any liquids. There are chemicals on the paper that could contaminate the liquids you are testing. Also, avoid smearing any liquid on your gloves or body as this may irritate your skin or contaminate the pH tests of other materials.
- 4. After placing test liquid on pH test paper, wait 8-10 seconds and then compare the color of the test strip where the liquid was placed to the pH color change guide. The pH of the liquid is the number next to the color that most closely matches the test strip. If the color seems to be between the guide colors, estimate between the pH values.
- 5. Record the name of the substance that you tested, the chemical name (if known) and the pH value that you determined using the test strips in the data table below.
- 6. DO NOT wait longer than 12 seconds to read your test tape because some materials oxidize and will give you inaccurate pH results.
- 7. DO NOT try to remember the values- the strips will change with time, and may even return to the original color when dry. Record your readings as soon as you complete the reading.
- 8. The pH test strips can not accurately determine the pH of strongly colored dyes. If you have a material that is colored, wipe the liquid on the test tape and then with the other side of the Qtip (dipped in distilled water) remove any excess liquid. This prevents staining of the tape and allows more accurate pH measurement.
- 9. If you chose to sample a solid object (only 25% of the total number of objects tested may be solid), saturate a clean Qtip in distilled water, wipe the Qtip on the solid object for 30 seconds, then rub the Qtip on the test tape and read as described above. Remember to record your results immediately!
- 10. Throw all used Qtips and pH test strips away in a proper trash receptacle immediately following testing.

	Substance Tested	Chemical Name (if chemical name is not known, state the function of the substance-ex Fantastik=cleaner)	Room number	pН
1				
2				
3				
4				1
5				
6	·			
7				
8				
9		04		
10		91		

MAM	AME	_SCI#	POINTS:	
Que	uestions:			
ACID	clude with this lab sheet a list of the materials your CIDIC to MOST BASIC. Include the substance test TED!!!			
1.	1. Why is it important to use <i>distilled</i> water to moist	en Qtip when	sampling materials?	
2.	2. What is the pH range of an alkaline solution? Nam	ne three alkali	ne substances.	
3.	3. What is the pH range of an acidic substance? Nam	e three acidic	substances.	,
4.	4. What chemical properties does pH actually represe	ent?		. !
5.	5. An increase in the pH from 8 to 9 indicates that the amount?	number of h	ydroxide ions has increased	by what
6.	6. A decrease in the pH from 3 to 2 represents an inci	ease or decre	ase in hydrogen ions? By ho	w much?
7.	7. Why is it important to know the acidity/ alkalinity	of the substan	ces you are working with?	
8.	8. Does the pH range for acids seem to go against into knew nothing about acids and bases, but were told would you say acids were on? Does this make it ea common substances? Why?	that the pH ra	nge was between 1 and 14,	for pH)? If you what end

Measuring Food Energy

Pre-Lab Discussion

All living things need energy to carry out metabolic activities. Animals—unlike many plants, protists, and bacteria—do not have the means to get energy directly from sunlight or simple inorganic chemicals. The energy requirements of animals must be met by taking in food.

The energy content of food can be determined by burning a sample of food in a device called a calorimeter. Heat energy released by combustion is absorbed by a container of water. Any rise in water temperature is measured and then used to determine the value of the heat energy released by the burning food sample. Heat energy is expressed in units called calories. One calorie is the amount of heat needed to raise the temperature of 1 gram of water by 1 degree Celsius. This unit, however, is too small for evaluating food energy. A Calorie, which is equal to 1000 calories, is used to measure food energy.

In this investigation, you will construct a simple calorimeter and use it to measure the amount of heat energy contained in certain foods.

Problem

How is the energy in food measured?

Materials (per group)

Ring stand

Four food samples Heat-resistant gloves

Test tube clamp Test tube

Triple-beam balance 100-mL graduated cylinder

Paper clip Cork stopper

Matches

Fireproof pad

Thermometer

Metric ruler

Safety & 🙉 🕮 🝩

Put on a laboratory apron if one is available. Put on safety goggles. Handle all glassware carefully. Use extreme care when working with heated equipment or materials to avoid burns. Note all safety alert symbols next to the steps in the Procedure and review the meanings of each symbol by referring to the symbol guide on page 10.

Procedure

- 72
- 1. To assemble a calorimeter, set up a ring stand, test tube clamp, test tube, and fireproof pad as shown in Figure 1.
- 2. To make a food platform for the calorimeter, bend the outer end of a paper clip straight down so that it is at a right angle to the rest of the clip. Insert the free end of the clip into the middle of the narrow end of the cork stopper. See Figure 2.
- 3. Place the food platform on the fireproof pad. Adjust the height of the test tube so that the space between the food platform and the bottom of the test tube is 2 cm.
- 4. Use a graduated cylinder to measure exactly 15 mL of water into the test tube. Record the mass of the water in the appropriate place in the Data Table. Note: Remember that 1 mL of water has a mass of 1 g.
- 5. Measure the temperature of the water in the test tube. Record this number in the appropriate place in the Data Table. Note: Be sure to remove the thermometer from the test tube after you record the temperature.
- 6. Select a food sample and find its mass using the triple-beam balance. Record the mass in the appropriate place in the Data Table. Also record the name of the food sample used in the appropriate place in the Data Table.
- 7. Place the food sample on the paper clip platform. Ignite the food sample with a match, and quickly place the platform under the test tube. CAUTION: Wear safety goggles when doing this part of the investigation. Be careful when using matches. Allowthe food to burn completely. Reignite the sample if necessary.

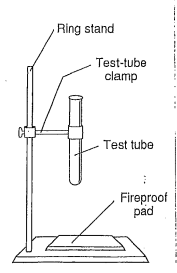


Figure 1

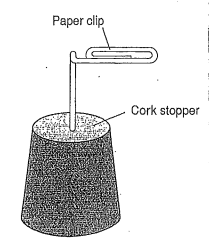


Figure 2

- 8. After the sample has burned completely, measure the temperature of the water in the test tube. CAUTION: Do not touch the test tube; it may be hot. Record the temperature of the water.
- 9. Find the mass of the remainder of the burned food sample, Record the mass,
- 10. Determine the change in mass of the food sample. Record the result.
- 11. Determine the change in the temperature of the water in the test tube. Record the result.
- 12. Repeat steps 3 through 11 using three other food samples. Note: Remember to empty the water out of the test tube and to use cool water for each sample.
- 13. Use the formula below to find the energy value, or Calories, per food sample. Record the results in the appropriate place in the Data Table. Note: The specific heat of water is 1 Calorie per kilogram degree Celsius.

Calories Change in Mass Specific
$$\frac{1 \text{ kg}}{\text{per}}$$
 = water x of x heat x $\frac{1 \text{ lood g}}{1000 \text{ g}}$

14. Use the formula below to find the Calories per gram of food sample. Record the results in the appropriate place in the Data Table.

Calories per gram = Calories per food sample/Change in mass of food sample

Observations

Data Table

Data Table						
	Food Sample					
Variable						
Mass of food sample before burning (g)						
Mass of food sample after burning (g)						
Change in mass of food sample (g)						
Mass of water (g) (1 mL = 1 g)						
Temperature of water before heating (°C)						
Temperature of water after heating (°C)						
Change in water temperature (°C)						
Calories per food sample						
Calories per gram						

Analysis and	Conclusions	
--------------	-------------	--

· '10.

1.	What is the difference between a calorie and a Calorie?						
2.	Why must the food sample be ignited before placing the platform under the test tube?	,					
3.	Why must the thermometer be removed from the test tube when the food sample is burnin	g?					

- 14. - 14.	
¹³⁾⁷⁷	
5.	Fats yield more food energy than proteins or carbohydrates. Which of your food samples mos
	likely contained the greatest amount of fat?
itical	Thinking and Application
1.	Swimming for one hour burns up 600 Calories. For each food sample you tested, calculate how
	many grams of food you would have to eat to get this energy.
2.	Fad diets, which have become popular in the past two decades, involve the consumption of large amounts of a limited variety of foods. Explain why some fad diets may be an unhealthful way to lose weight.
3.	Although fiber is not officially classified as a nutrient, it is an important component of the American diet today. What is the role of fiber in the human body?
	Contrast the snacks for a person who is trying to lose weight with those for a person who is
	growing very rapidly,

Going When

Using the procedure from this investigation, determine the Caloric value of various diet foods and their counterparts. Is there a difference in their Caloric values?

- 5. In the Data Table, write the name of the type of cell that you examined. Describe the general shape of the cell in the space provided. Estimate the length of the cell and record this figure. Refer to Laboratory Investigation 4 if you need to review how to estimate the size of objects under the microscope. Put a check mark next to the cell structures you are able to observe under low power.
- 6. Switch to the high-power objective lens. CAUTION: When turning to the high-power objective lens, you should always look at the objective from the side of your microscope so that the objective lens does not hit or damage the slide. Look for cell structures unobservable under low power. Put a check mark next to these structures in the Data Table. Based on your observations, decide if the cell is prokaryotic or eukaryotic and record this in the Data Table.
- 7. In the appropriate place in Observations, draw and label what you see using the high-power objective lens. Record the magnification of the microscope.
- 8. Repeat steps 1 through 7 using other prepared slides provided by your teacher.
- 9. Repeat steps 1 through 7 using an unidentified prepared slide provided by your teacher.
- 10. When you have finished examining all of the prepared slides, return the microscope to the storage area.

Observations Data Table

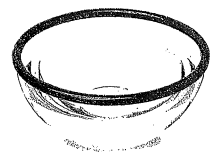
					Cell	Struc	tures			
Cell Type	Shape	Size (µm)	Cell wall	Cell wall Cell membrane Nucleus Nuclear envelope	Nuclear envelope	Cytoplasm	Vacuoles	Plastids	Prokaryotic or Eukaryotic	
,										
Unknown		1								

4				
Prepared Slide 1 Hig	h-power objective		Prepared Slide	2 High-power objective
Magnification			Magnification _	
		•		
		97		

Break water's surface tension

THE SURFACE OF WATER pulls in all directions. This is called surface tension. Try breaking water's

surface tension and watch what happens!



Fill up a small bowl with cold I water until it is about three-quarters full.



Try floating a paper clip on top of the water's surface. Can you see the surface stretching under the weight of the paper clip?

Find in your kit:

e 2 boat shapes

» paper clip

Float the

small drop of dishwashing liquid

Itwo boats on

the surface of the

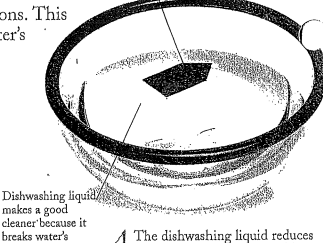
water. Gently add a

behind the boats and

watch what happens.

Find around your home:

- o dishwashing liquid
- bowl of water



How far do your boats travel?

makes a good cleaner because it breaks water's surface tension.

> Float the two boats next to each other on the water's surface.

More experiments

Place a different amount of dishwashing liquid behind each boat and race them. Which boat travels farthest?

The water's surface tension. The

surface tension at the front of the

boats pulls them forward.

What other substances can you use to break water's surface tension?



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Float a clay boat



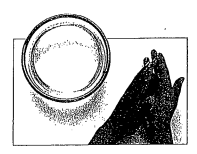
If BOATS ARE MADE FROM heavy materials, why don't they sink? By simply reshaping a ball of clay, you can learn how boats float on water.

Why does the water

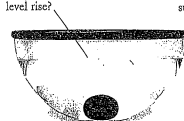
Find in your kit:

modeling clay

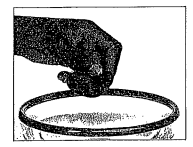
Find around your home:
small bowl of water



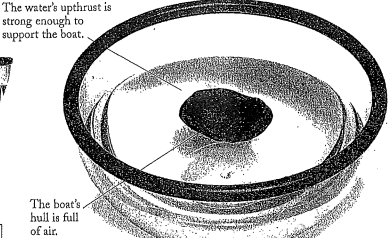
Pour some water into a small bowl until it is about three-quarters full. Roll the modeling clay into a ball.



Drop the clay ball gently into the water. The clay ball sinks because it is more dense than the water.



Now carefully place the clay boat in the bowl of water. Do you think that the boat will float?



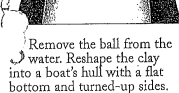
The boat shape
displaces, or pushes
away, more water than
the clay ball did. This
increases the water's
upthrust and makes the
boat float on the surface.
Try putting a cargo of
small marbles in the boat.
Does the boat float
or sink?

More experiments

Test whether your clay boat can hold different types of cargo from your kit. Then record the results in your Scientist's Notebook.



6 7



WATER MOLECULES.

ELECTRON DIAGRAM*

PROTON_P+

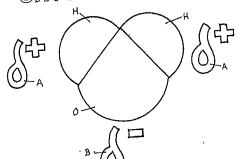
NEUTRON.

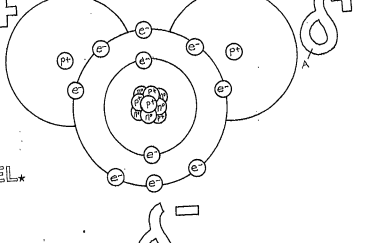
ELECTRON.

POSITIVE CHARGEA NEGATIVE CHARGEB

SPACE-FILLING MODEL*
HYDROGENH

OXYGEN.

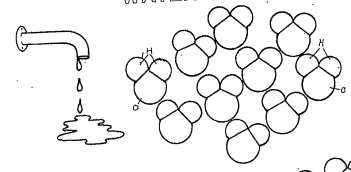




EMPIRICAL FORMULAH20

H20.

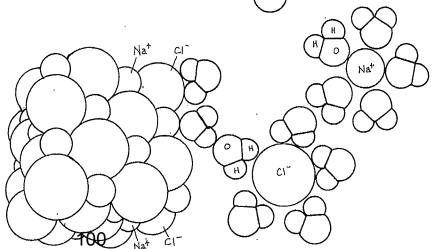
HYDROGEN BONDING OF WATER MOLECULES.

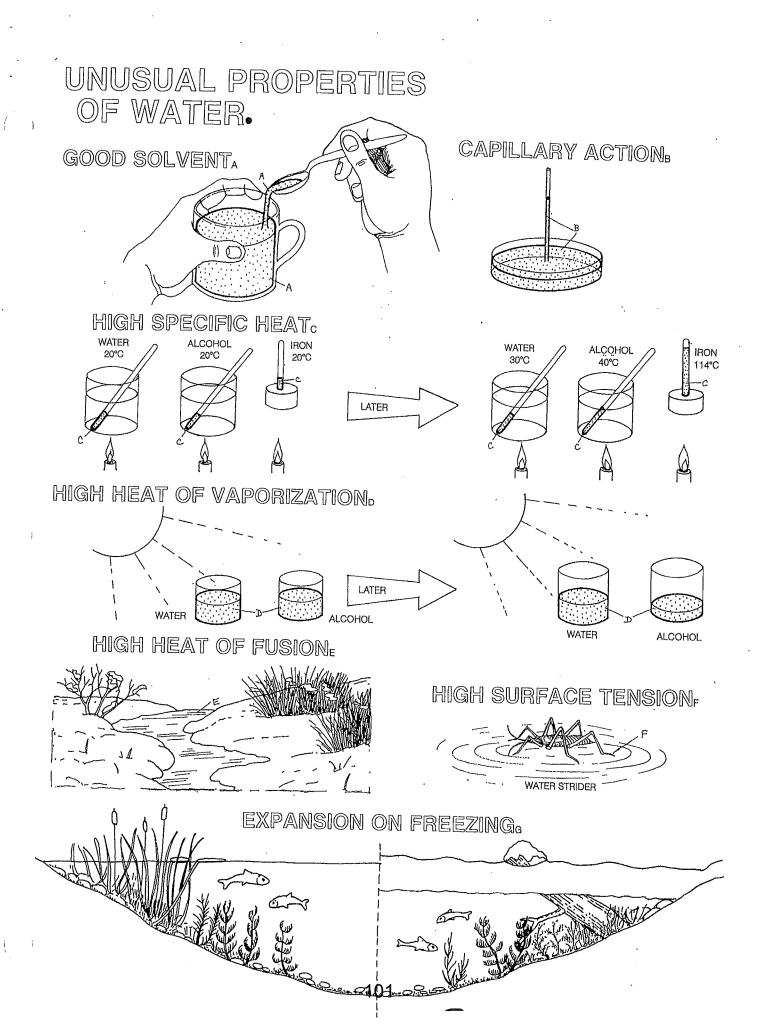


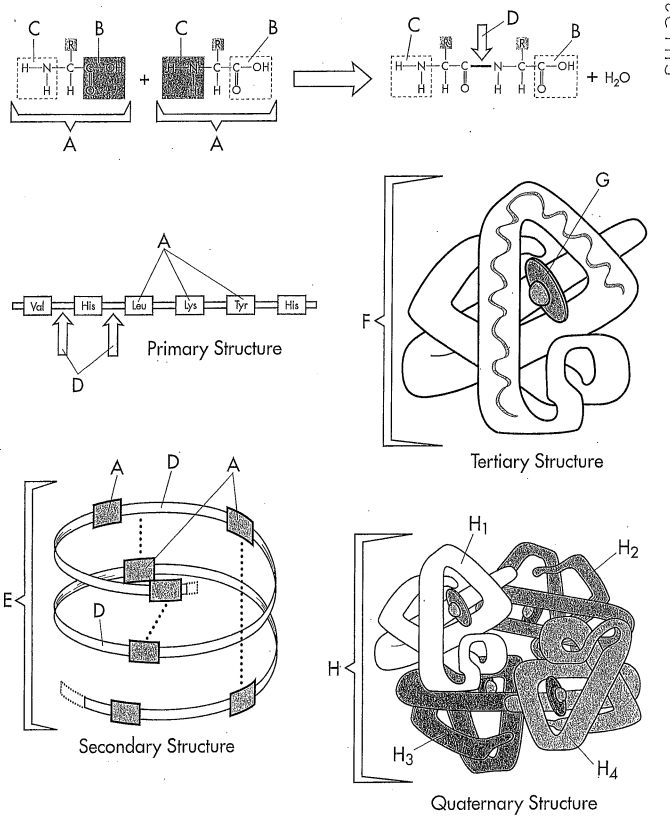
DISSOLVING OF AN IONIC COMPOUND*

SODIUM IONNA+

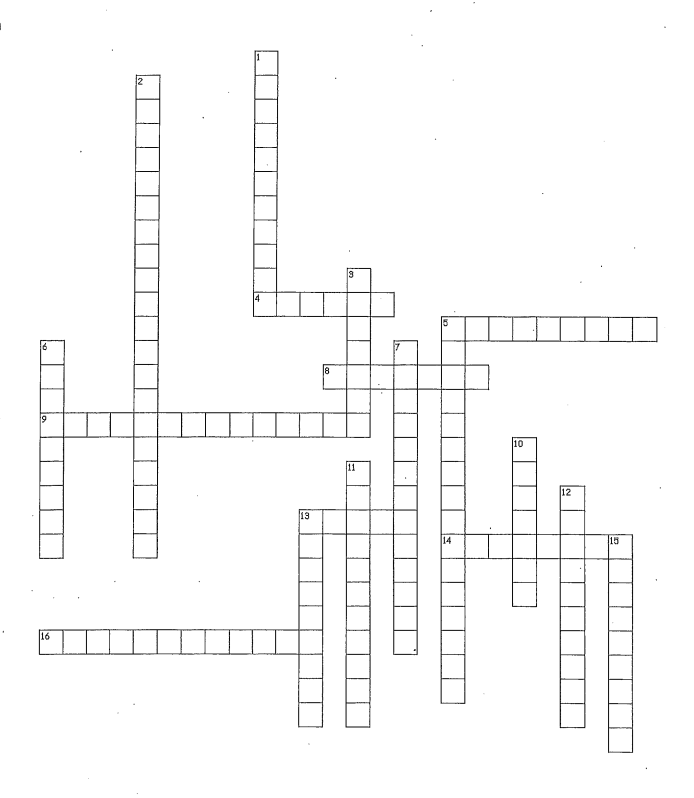
CHLORIDE IONa-







Ch 7 Crossword and Flashcards—complete the crossword, then make a flashcard for each term 18 cards



Across

- 4. a group of similar cells that perform a common function
- 5. the region of the cell within the membrane
- 8. a small cavity or sac that contains materials in a eukaryotic cell
- 9. a cell organelle that helps make and package materials to be transported out of the cell
- 13. a collection of tissues that carry out a specialized function of the body
- 14. a cell organelle where protein synthesis occurs
- 16. a phospholipid layer that covers a cell's surface and acts as a barrier between the inside of a cell and the cell's environment

Down

- 1. an organelle found in plants and algae cells where photosynthesis occurs
- 2. a system of membranes that is found in a cell's cytoplasm and that assists in the production, processing, and transport of proteins and in the production of lipids
- 3. in a eukaryotic cell, a membrane-bound organelle that contains the cell's DNA
- 5. a collection of genetically identical cells that are permanently associated but in which little or no integration of cell activities occurs
- 6. a long, hairlike structure that grows out of a cell and enables the cell to move
- 7. in eukaryotic cells, the cell organelle that is surrounded by two membranes and that is the site of cellular respiration
- 10. a fluid-filled vesicle found in the cytoplasm of plant cells or protists
- 11. a group of organs that work together to perform body functions
- 12. a single-celled organism that does not have a nucleus or membrane-bound organelles
- 13. one of the small bodies that are found in the cytoplasm of a cell and that are specialized to perform a specific function
- 15. an organism made up of cells that have a nucleus and membrane-bound organelles

CHAPTER 7 VOCAB

Cell membrane	<u></u>
Cytoplasm	
Ribosome	
Prokaryote	
Eukaryote	
Nucleus	
Organelle	
Vesicle	
endoplasmic reticulum	
Golgi apparatus	
Vacuole	
Chloroplast	
Mitochondrion	
Flagellum	
Tissue	
Organ	
organ system	
colonial organism	
,	

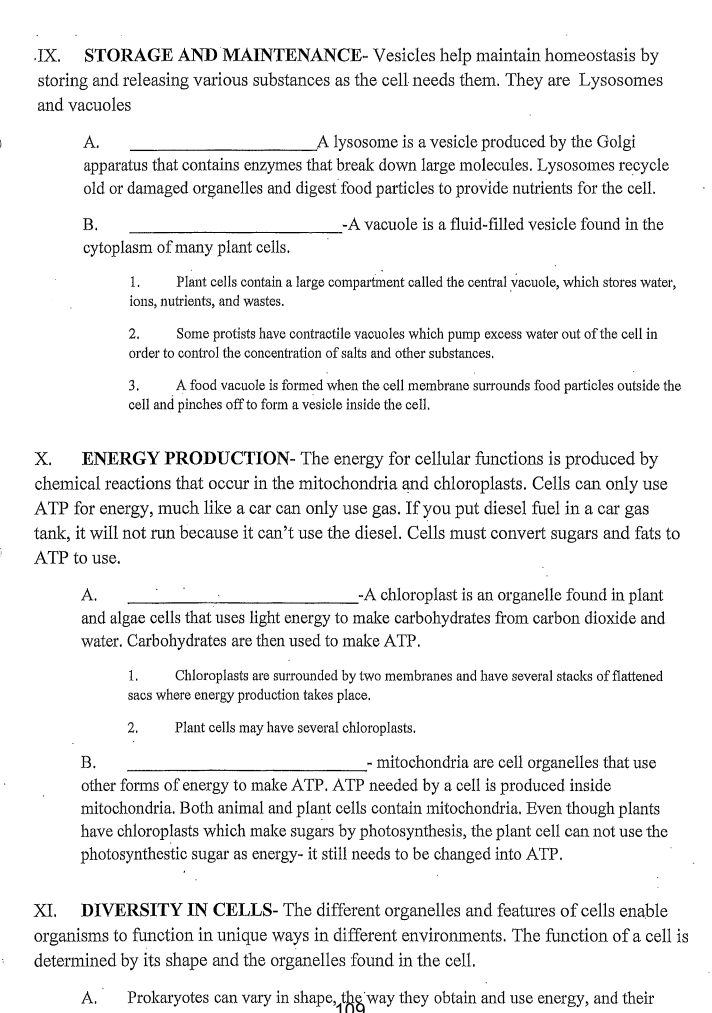
Chapter 7 Cell Structure

I.		DISCOVERY OF CELLS- Microscope observations of organisms led to
the d	iscove	ry of the basic characteristic common to all living things.
	A.	Robert Hooke used a microscope to discover cells in
	B. organ	Anton van Leeuwenhoek used a more powerful microscope to see single-celled isms in pond water.
Π.	CEL	L THEORY-The cell theory states:
	Α.	All living things are made up of one or more
	В.	Cells are the basic units of structure and function in organisms.
	C.	All cells arise fromcells.
III. by a c		L FUNCTION- A cell's shape reflects the cell's function. Cell size is limited
	A.	All substances that enter or leave a cell must cross the surface of the cell.
	B. its sur	A cell's ability to move substances across its surface can be estimated by finding rface area-to-volume ratio.
	C.	Cells with surface area-to-volume ratios can ange substances more efficiently.
	D. to-vo. cells.	When comparing cells of the same shape, small cells have greater surface area- lume ratios than large cells. Small cells function efficiently than large
	out m	L FEATURES- Because of their complex organization, eukaryotic cells can ore specialized functions than prokaryotic cells can. All cells share common eatures, including a cell membrane, cytoplasm, ribosomes, and DNA.
	A.	The cell membrane is the outer layer that covers a cell's surface and acts as a between the outside environment and the inside of the cell.
	В.	The cytoplasm is the region of the cell within the cell membrane. The includes the fluid inside the cell called the cytosol.
	C.	A is a cellular structure (but does not have a brane) that makes proteins.
	TITOTII	Miller Miller Manager Properties

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	D. regula		of a cell provides instructions for null large activities, and enables cells to reproduce.	naking proteins,
V.	DIFI	EREI	NCES BETWEEN TYPES OF CELLS	
	A. single		aryotic cell.	inism made of a
		1. geneti	Prokaryotic cells do not have a nucleus or other internal conic material of a prokaryotic cell is a single loop of DNA.	npartments. The
		2.	Prokaryotes are more and existed f	irst
	B. more	eukar	- A eukaryote is an organism m ryotic cells. All multicellular organisms are made of e	- ·
		1. the nu	The DNA of a eukaryotic cell is found in an internal comparticleus.	ment of the cell called
		2. structi	All eukaryotic cells have membrane-bound organelles. An or ture found in the cytoplasm that carries out specific activities in	-
_	its shaj	pe, and	MEWORK OF THE CELL-The cytoskeleton help d organize its parts. Eukaryotic cells have an intricate ed the cytoskeleton which provides the interior fram	e network of
	A.	There	e are three different kinds of cytoskeleton fibers:	
		1.	Microfilaments	
		2.	Microtubules	
		3.	Intermediate fibers.	
activi	structi ties. D	ons fo	NG CELLULAR ACTIVITY- DNA is the "brain" or making all proteins. The proteins then go on to consilike a general, the proteins are the soldiers. The sole but without the general making the decisions, there	mplete ALL diers are actually
	A. of the		contains instructions for making proteins which control	most of the activity
	В.	The D	ONA of eukaryotic cells is stored in the	,
		ıs. Nu	uble membrane called the nuclear uclear pores located on the nuclear envelope act as chang o move in and out of the nucleus. 107	

cyto	s are r plasm	The is a structure within the nucleus where ribosome re made. These ribosome parts are transported out of the nucleus into the asm where they are assembled to form a complete ribosome. Ribosomes are the less that make the proteins.								
E	Ribosomes that are suspended in the cytosol are called ribos									
F.	Fre	that remain inside the cell.								
G.	Rib	posomes that are attached to the endoplasmic reticulum are called ribosomes. Bound ribosomes make proteins that								
are e	xport	ed from the cell.								
H. the c		osomes can switch between being bound or free, depending on what proteins eds to make.								
		N PROCESSING- The endoplasmic reticulum and Golgi apparatus are protein processing								
A.		- The endoplasmic reticulum								
and t	he Go	olgi apparatus are organelles that prepare proteins for extracellular export.								
		Proteins that are sent outside the cell are packaged in vesicles. Vesicles are small, abrane envelopes that enclose the proteins and keep them separate from the rest of the plasm.								
	2. other	The endoplasmic reticulum, or ER, is a system of membranes that moves proteins and r substances through the cell and make the vesicles.								
	3.	The endoplasmic reticulum is divided into two portions: rough ER and smooth ER.								
	4.	The ribosomes on the rough ER make proteins that are packaged into vesicles.								
	5.	Enzymes of the smooth ER make lipids and break down toxic substances.								
		- The Golgi apparatus is a set of nembrane-bound sacs. The Golgi apparatus helps modify, sort, and package ets for distribution.								
	1. mem	The ribosomes located on the rough ER make proteins which then cross into the branes of the ER.								
•	2.	The ER membrane then pinches off and forms a vesicle around the proteins.								
	3. enzy	Vesicles move from the rough ER to the Golgi apparatus, where they are modified by mes and repackaged in new vesicles then are sent out of the cell or stored.								



ability to move.

- 1. Many prokaryotes have a flagellum, a long, hair-like structure that grows out of the cell and enables the cell to move through its environment.
- 2. Prokaryotes may also have pili, short outgrowths that allow the cell to attach to surfaces or other cells.
- B. Eukaryotic cells can vary in shape, external features and internal features. Eukaryotic cells usually have a "specialty"- a specific job they must do for the health of the whole organism. Remember eukaryotes are multicellular.
 - 1. Your skin cells and brain cells do not have the same job and so do not look or function the same.
 - 2. Animal and plant cells are two types of eukaryotic cells. Both have many of the same organelles, but plant cells also have chloroplasts, a large central vacuole, and a cell wall.

		ELS OF ORGANIZATION- Plants and animals have many highly cells that are arranged into tissues, organs, and organ systems.
	A. a com	A is a distinct group of similar cells that perform amon function.
	B. work	An is a collection of tissues that together to form a structure which performs a specific function.
	C. group	An is composed of a of organs that work together to perform major body functions.
XIII.	BOD	Y TYPES- organisms can be unicellular or multicellular.
	A. groups	organisms can thrive independently or live together in
	B. activit	Cells that are permanently associated but do not work together or integrate cell ies are called organisms.
	function cell, we a process	True multicellularity occurs only in eukaryotes. In a multicellular body, cells are ependent – they can NOT live alone. Distinct types of cells have specialized ons to help the organism survive. Most multicellular organisms begin as a single thich divides to form more cells. These cells then grow and become specialized in ess called differentiation. Once differentiation occurs, the specialized cells can not in their own.

CH		2	cr	n	7
1 H	ш	\mathbf{r}		к	•

1. Indicate whether each structure or feature below is found in a prokaryotic cell, a eukaryotic cell, or both.

Cell structure or feature	Prokaryotic cell	Eukaryotic cell
Nucleus	no	yes
Cell membrane		
Cytoplasm		
DNA		
Ribosomes		
Membrane bound organelles		

2. What are the three parts of the cell theory?

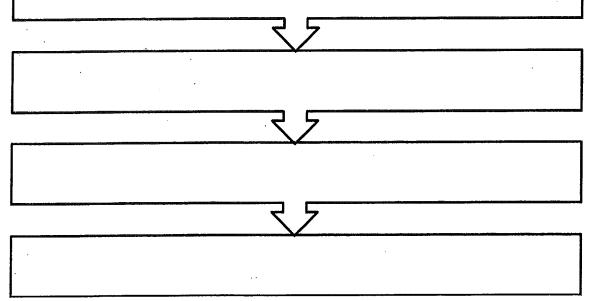
- 3. Could a cell be the size of an elephant? Explain your answer.
- 4. How does the location of DNA differ in prokaryotic and eukaryotic cells?

Bellringer:Day MIT:W/Th F.Date遗

Otlestion

1. Complete the process chart to describe how proteins are made andmoved out of the cell.

Ribosomes use the instructions carried by RNA to build proteins.



2. How does DNA direct the cell's activities, such as making proteins, if DNA stays inside the nucleus?

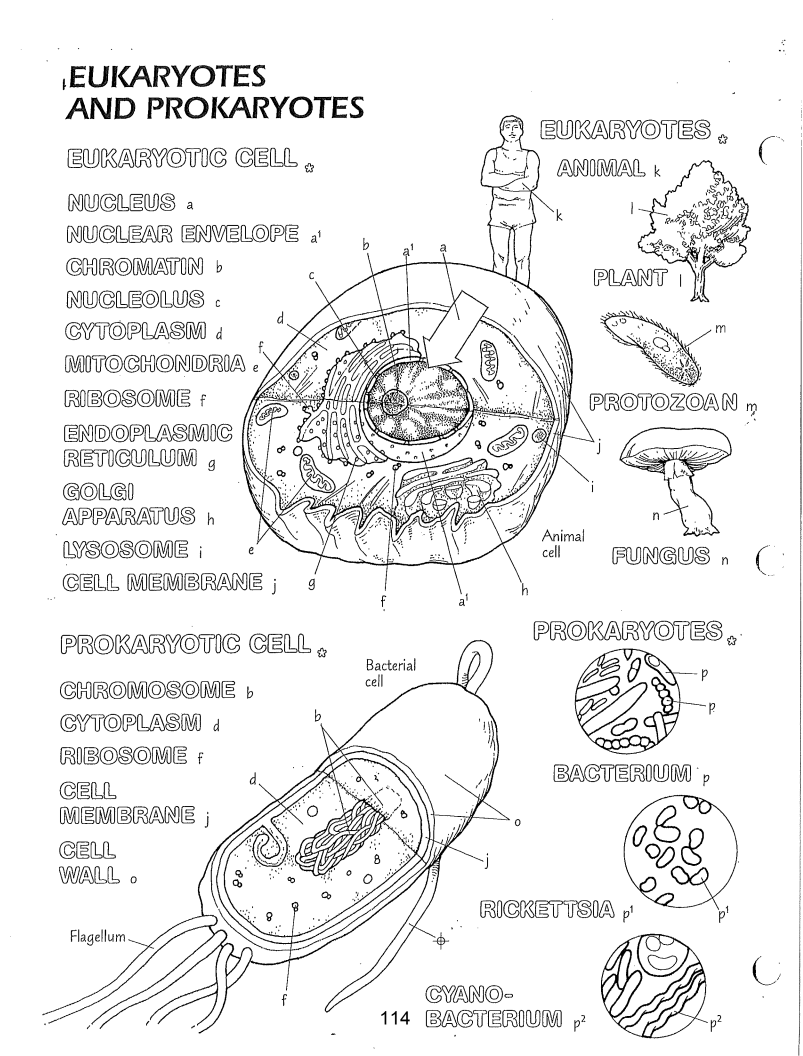
3. Why do plant cells need both chloroplasts and mitochondria?

Bellringer Day Mrt. W This Date

Oilestion

∖nswer≝

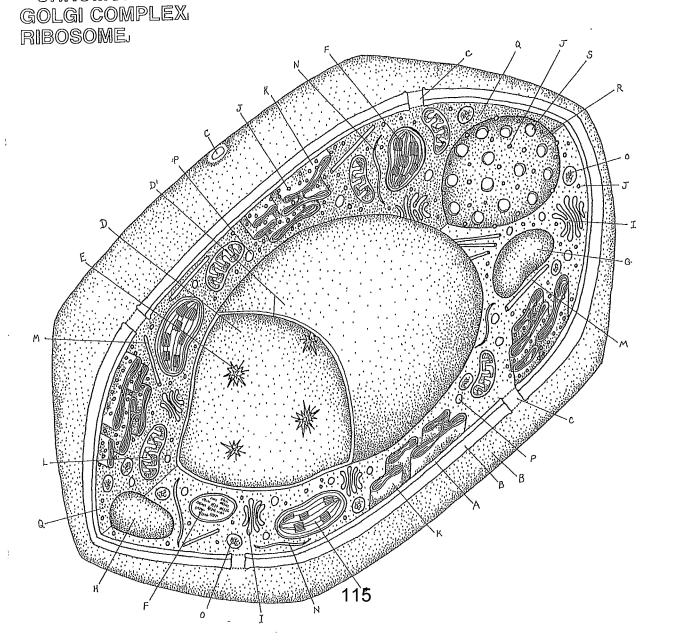
CHAPTER 7	SEC 3	DUE DATE .
1. Why are spe	cialized cells found only in m	nulticellular organisms?
2. Identify four	ways that prokaryotes can d	iffer from one another.
	, , , , , , , , , , , , , , , , , , , ,	
3. Why are cold	onial organisms not truly mul	ticellular?
		·
4. How would p	ili be important to colonial ba	acteria?
300.		
5. What are the	four levels of organization o	f complex multicellular organisms?
LAM. Att.		
ellringer Day Mit V	V.Th F.Date Charles	
nsweralis is a second		
		tage to the second seco



PLANT CELL.

CELL MEMBRANEA
CELL WALL
PLASMODESMA
VACUOLE
TONOPLAST
CRYSTAL
PLASTIDS
CHLOROPLAST
LEUCOPLAST
CHROMOPLAST

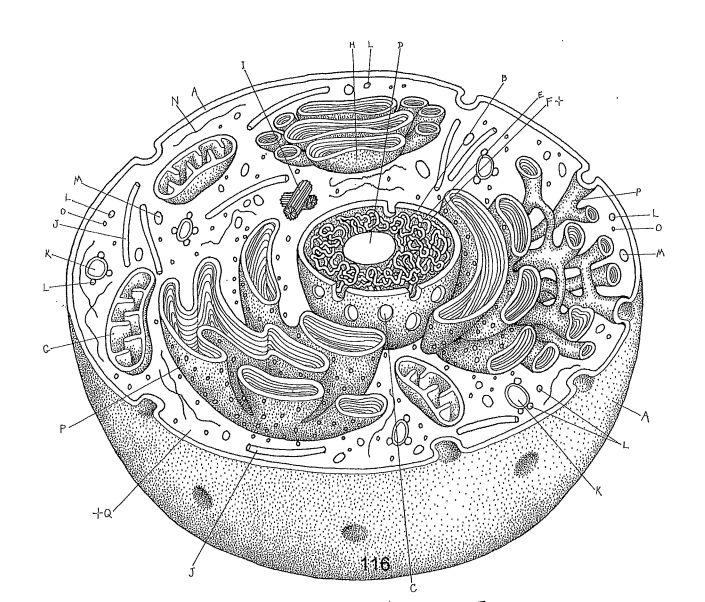
ENDOPLASMIC RETICULUM
MITOCHONDRION
MICROTUBULE
MICROFILAMENT
LYSOSOME
MICROBODY
HYALOPLASM
NUCLEUS
NUCLEAR ENVELOPE
NUCLEAR PORE



ANIMAL CELL.

CELL MEMBRANEA
NUCLEUS*
NUCLEAR ENVELOPEB
NUCLEAR POREC
NUCLEOLUSD
CHROMATINE
NUCLEAR SAPE+
CYTOPLASM*
MITOCHONDRIONG
GOLGI COMPLEXH

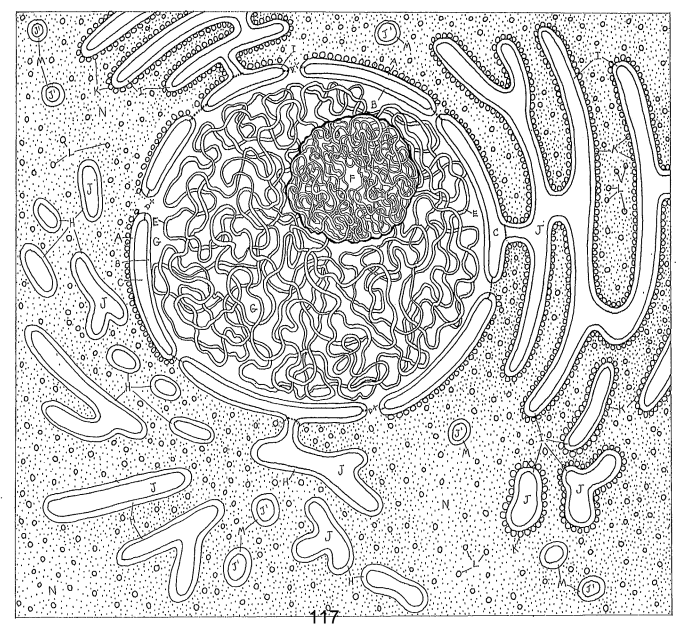
CENTRIOLE,
MICROTUBULE,
VACUOLE,
LYSOSOME,
MICROBODY,
MICROFILAMENT,
RIBOSOME,
ENDOPLASMIC RETICULUM,
HYALOPLASMO;



NUCLEUS AND ENDOPLASMIC RETICULUM.

NUCLEAR ENVELOPE*
OUTER MEMBRANE*
INNER MEMBRANE*
PERINUCLEAR SPACE*
NUCLEAR PORE*
CHROMATINE
NUCLEOLUS*
NUCLEAR SAP*
SMOOTH ENDOPLASMIC
RETICULUM*

ROUGH ENDOPLASMIC
RETICULUMI
CISTERNAJ
ATTACHED RIBOSOMER
FREE RIBOSOMEL
VESICLEM
CONTENTSJ
HYALOPLASMIN



MITOCHONDRIONA

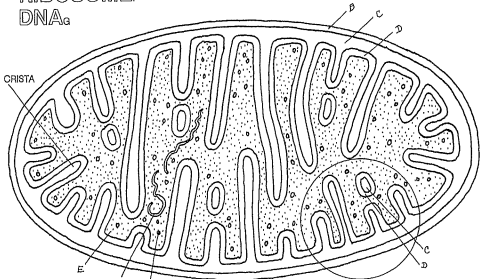
OUTER MEMBRANE:

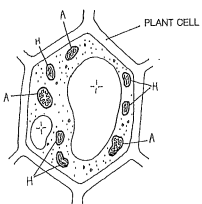
INTERMEMBRANE SPACE.

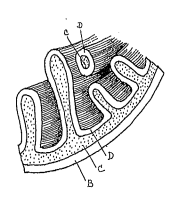
INNER MEMBRANE

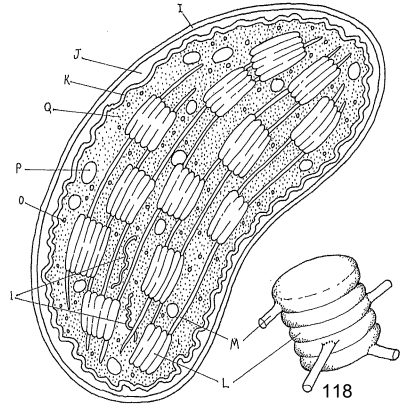
MATRIXE

RIBOSOME,









CHLOROPLASTH OUTER MEMBRANE INTERMEMBRANE SPACE, INNER MEMBRANEK GRANUM*

THYLAKOID: STROMAL LAMELLAM DNAM RIBOSOME: STARCH GRAIN: STROMA:

GOLGI COMPLEX, LYSOSOMES, MICROBODIES.

GOLGI COMPLEX.

SACCULE/VESICLE.

MEMBRANE.

COMPARTMENT.

COMPARTMENT.

GOLGI COMPLEX IN ACTION.

AMINO ACID MOLECULES.

CELL MEMBRANE.

RIBOSOME.

ROUGH ER MEMBRANE.

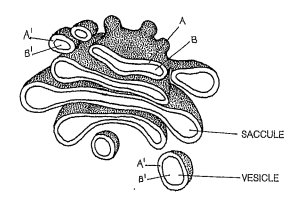
CISTERNA.

POLYPEPTIDE CHAINS.

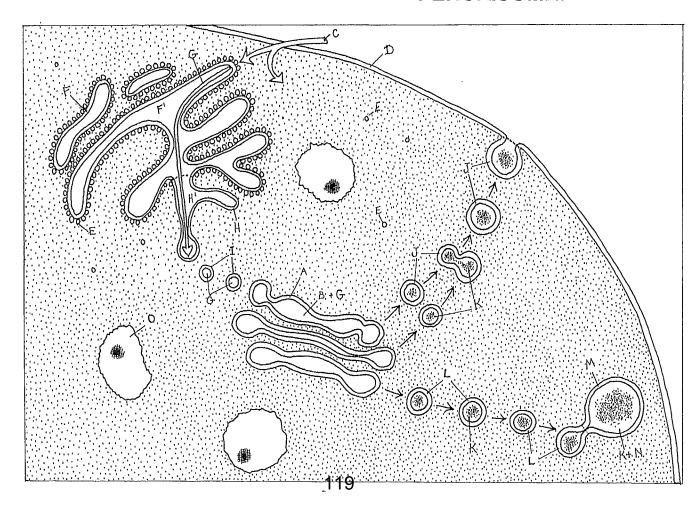
SMOOTH ER MEMBRANE.

CISTERNA.

TRANSITION VESICLE.



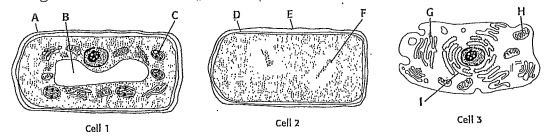
SECRETION VESICLE,
PROTEIN COMPLEX
LYSOSOME
FOOD VACUOLE
FOOD
MICROBODY
PEROXISOME



Science Skills

INTERPRETING GRAPHICS

Biology students were working on a class project. They prepared copies of transmission electron micrographs of a bacterium, a plant cell, and an animal cell for display in their classroom. Unfortunately, the pictures were not labeled and got mixed up. Help these students correctly identify the cells and cell structures. Use the figures below to answer questions 1–5.



In the space provided, write the names of each cell's labeled structures (A–I). Using this information, write the identity of each cell—bacterium, plant cell, or animal cell.

1.	Cell 1 identity
	A
	B
	C
	Cell 2 identity
	D
	Е
	F
	Cell 3 identity
	G
	Н.
	I

Characteristics of Prokaryotic and Eukaryotic Cells

Pre-Lab Discussion

Cells are the basic units of structure and function of all living things. There are two major divisions into which all cells fall—prokaryotic and eukaryotic.

Prokaryotic cells are cells that lack a nucleus and membrane-bound organelles. Bacteria and related microorganisms are prokaryotes. Eukaryotic cells are cells that contain a nucleus and membrane-bound organelles. Organisms such as animals, plants, fungi, and protists are all eukaryotes.

In this investigation, you will observe several prepared slides to examine the differences between prokaryotic and eukaryotic cells. You will also use these differences to classify an unknown specimen.

Problem

What are the differences between prokaryotic and eukaryotic cells?

Materials (per group)

Microscope
Lens paper
Prepared slides of prokaryotic
and eukaryotic cells

Safety &

Always handle the microscope with extreme care. You are responsible for its proper care and use. Use caution when handling glass slides as they can break easily and cut you. Note all safety alert symbols next to the steps in the Procedure and review the meanings of each symbol by referring to the symbol guide on page 10.

Procedure

- 1. Take a microscope from the storage area and place it about 10 centimeters from the edge of the laboratory table.
- 2. Carefully clean the eyepiece and objective lens with lens paper.
- 3. Place your first prepared slide on the microscope stage so that it is centered over the stage opening. Hold the slide in position with the stage clips.
 - 4. Using the low-power objective lens, locate the cell(s) under the microscope. Turn the coarse adjustment knob until the cell comes into focus.

Prepared Slide 3 High-power objective Magnification		Prepared Slide 4 High-power objective Magnification	
	Prepared Slide 5	High-power objective	
	Magnification		
	Unknow	/n	
Analysis and Conclusions			
	ıs, do all cells have t	the same shape? Support your answer.	
O. D J	an de all collé baye t	he same size? Support your answer.	
2. Based on your observation		nie same size: bupport your unswer.	
	•		
3. What cell structures are co	ommon to all cells?		

۰C۰

,		
	4	. What cell structures are found only in eukaryotic cells?
	5	Are the nuclei always found in the same place within different types of cells? Support your
		answer.
Crit	ical	Thinking and Application
0,10		
	1.	Skin cells seem to fit together like pieces of a jigsaw puzzle. How is this arrangement of cells
		helpful to an organism?
1	2.	Why do cells have different shapes and sizes?
	3.	What cell structure might you be able to compare to the main (principal's) office in your
		school? Explain your answer

Going Further

- 1. Observe characteristics of living cells by making wet-mount slides of plant and animal tissues or protist cultures. Construct a data table to record the shapes and sizes of the cells and the structures they contain.
- 2. Think about the cell structures that you were unable to see with a compound light microscope. Use resources from your library to locate electron micrographs of these structures.
- 3. Research the use of some of the stains used in the preparation of wet-mount slides. Some of the stains that might be included in your report are methylene blue, neutral red, acetocarmine, Congo red, Janus green B, and Sudan III. What cell structures do each of these stains make more visible?



PART C - Observation Of Organelles In Living Tissue

We have already mentioned that mitochondria are responsible for breaking down food in the presence of oxygen. The enzymes that the mitochondria use to do this are called *dehydrogenases*, which remove hydrogen atoms from food molecules. We can "fool" the mitochondria by giving it other molecules with hydrogen, and the mitochondrion's enzymes will react the same way. The kit contains a dye called *Janus Green B*. The molecules of this dye are greenish-blue in color, and will be transported to the mitochondria as if they were food. If hydrogen atoms are removed from the dye molecules, though, the dye looses its color. By observing the loss of color in the mitochondria, we can observe mitochondria at work.

Obtain a second piece of celery, the same size as the first, a razor blade, a microscope slide and a cover slip, a pair of forceps, and a microscope.

Cut a piece of celery in the same manner as in Part A, isolating a piece of thin tissue from in between two celery threads. Add a drop of water to the slice, and drop a cover slip on top of it. Place the slide under the microscope as before and focus it, first on low power and then on high power.

Obtain a bottle of Janus Green B stain. Tear off a piece of paper towel, and have it ready next to your lab station. Place a drop of the Janus Green B stain on the slide immediately next to the cover slip. Place the piece of paper towel along the opposite edge of the coverslip. The paper towel will absorb the water underneath the coverslip, and the drop of stain on the other side will be drawn into its place. Observe the cells as they are stained. The stain will congregate in the mitochondrial cells, and will turn them bluish green. Observe the slide for several minutes, alternating with your partner, until you notice a change in the color of the mitochondria. Record and explain your observations:

Discussion Questions

Why do you think that it is important that enzymes be restricted to inside organelles?

What advantage do you think a cell has because of its organization into organelles?
Could the model for an organelle be used as a model for a cell? What are the differences between cells and organelles?
The organelles of a cell have been compared to the parts of the body: each part of the body performs a specific function. In what ways do you think this analogy is correct?
When you observed the living tissue of the celery, you may have noticed that the organelles were flowing around inside the cell. This process is called cytoplasmic streaming. Why do you think that this might be important to the cell?

What Cell Parts Can You See with the Microscope?

Living things are made of cells. All cells have parts that do certain jobs. Cells have an outer covering called the cell membrane. Cell membranes give cells their shapes and control what enters and leaves the cells. The clear, jellylike material inside the cell is the cytoplasm. The nucleus is the control center of the cell. Plant cells have a thick outer covering called the cell wall. It is on the outside of the cell membrane.

Cell parts can be studied by making wet mount slides. A wet mount slide is a temporary slide. It is not made to last a long time. You can make wet mount slides of living and once-living materials to study cell parts.

GOALS

In this exercise, you will:

a. make wet mount slides for examination under the microscope.

b. study four cell parts—the cell wall, cytoplasm, nucleus, and cell membrane.

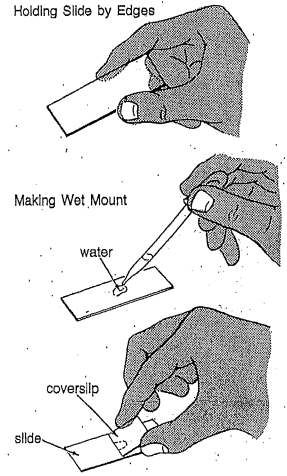
MATERIALS

glass slide coverslip light microscope water dropper forceps stain cork shaving bamboo shaving onion skin prepared slide of frog blood

PROCEDURE

- 1. Follow the steps below to make a wet mount slide.
 - a. Get a clean microscope slide and coverslip. Handle the slide and coverslip by the edges to keep them clean.
 - b. Use a dropper to put a drop of water in the center of the slide.
 - c. With forceps, place the object to be examined in the drop of water.
 - d. Hold the coverslip at an angle.

 Gently lower it onto the drop of water.

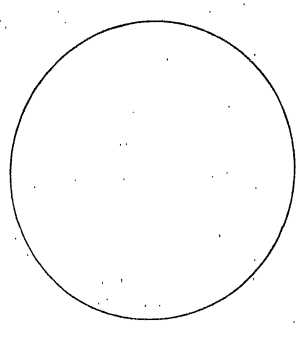


2. Prepare a wet mount of the cork shaving. Follow the steps just given on the last

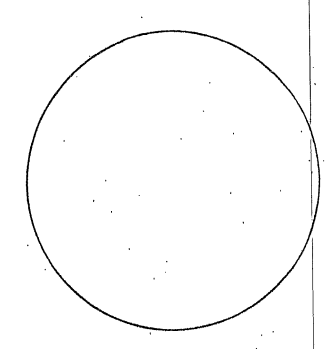
3. Examine the slide of cork under low power of the microscope. Switch to high power. Examine the cork cells under high power. Draw cork cells that you see in

the circle below. Label the cell wall.

4. Prepare a wet mount of a bamboo stem shaving. Examine the bamboo under low and then high power of your microscope. Draw the bamboo cells you see. Label the cell wall.

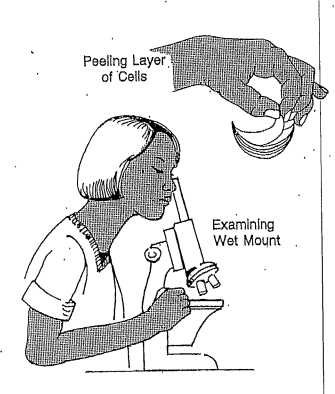


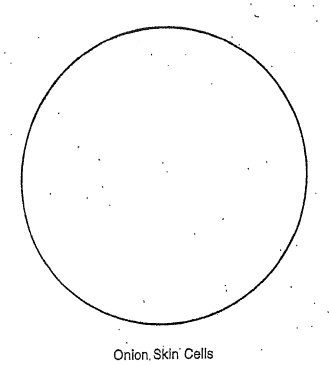
Cork Cells



Bamboo Cells

- 5. Peel the thin layer of cells from the inside of an onion as shown here. Make a wet mount of the onion skin cells. Add one drop of stain in place of water.
- 6. Examine the onion slide under low and high power of your microscope.
- 7. Find the cell wall, nucleus, and cytoplasm. Draw onion cells that you see in the circle on page 17. Label the parts.
- 8. Examine a prepared slide of frog blood with low and then high power. In the circle on page 17, draw frog blood cells that you see. Label the nucleus, cytoplasm, and cell membrane.
- .9. Complete the table on page 17.





Frog Blood Cells

Parts of Cells

Cell type	Cell wall present? (yes or no)	Nucleus present? (yes or no)	Cytoplasm present? (yes or no)	Shape of cell?	Cell living or dead?
Cork		•	•		
Bamboo					1
Onlon					
Frog blood					

QUESTIONS

- 1. What is the name of the small units that make up cork?
- 2. Describe how the small units of cork look.
- 3. Are the cork cells filled with living material or are they empty?

1	Are bamboo cells living or dead?	
4. .	How are cork cells and bamboo cells alike?	_
5.		_
	77 1166 1 C - 11	_
6.	How are onion cells different from the cork cells?	_
7.	Compare the onion skin cells and the frog blood cells.	
		_
8.	What cell parts that you observed are found only in plant cells?	
	TO A TOTAL OF THE STATE OF THE	
AP 1.	PLICATIONS Why do cells have different shapes?	
•		
2.	Skin cells seem to fit together or overlap. How is this cell arrangement helpful to	O
	the organism?	
	If blood cells were box-shaped, like onion cells, why would they be unable to d	0
3.	, , , , , , , , , , , , , , , , , , ,	Ī
	their job as well?	
Fil	OCABULARY I in the blanks with the proper word or words.	
1.	. Cells have an outer covering called the	L
2.	. The jellylike material inside the cell is the	-
	. The control center of a cell is the	L
,	Plant calls have a thick outer covering called the	
5.	. When you place an object in a drop of water on a slide and put a coversity on	
	it, a slide is made.	

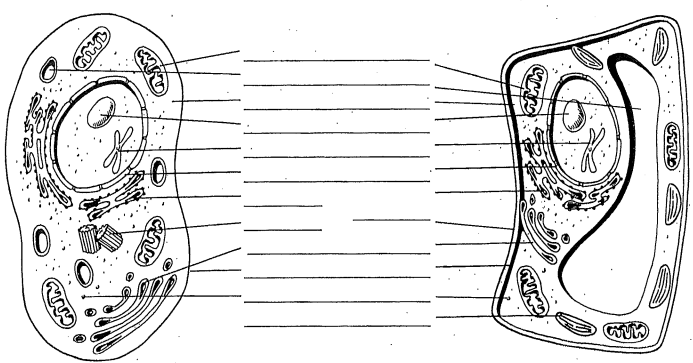
Pre-Lab Questions

Is a plan	nt a prokaryote or eukaryote? What is a prokaryote?
-	
What is	the cell cycle? Is mitosis a part of the cell cycle?
Why do -	es a cell undergo mitosis?
- Can you 	see DNA in all cells under the microscope? Why or why mot.
	e your predictions for this lab? Be specific- what do you think is the
ongest r 	part of mitosis.
_	

CELL PARTS AND THEIR JOBS

In your textbook, read about cell parts and their jobs in Section 2:2.

1. Label the parts of these two cells in the spaces provided.



- Cell B Cell A
- 2. Read the descriptions of cell parts below and write in the name of the cell part. Use the color indicated to shade the pictures above.
 - a. Use red for the part that gives the cell shape and holds the cytoplasm.
 - b. Use green for parts that make food.
 - c. Use brown for the thick outer covering that protects and supports the cell.
 - d. Use blue for the part that stores substances.

 - e. Use black for parts that release energy from food.
 - f. Use purple for parts that carry hereditary information.
 - g. Use pink for the cell part that helps with cell reproduction.
 - h. Use orange for the parts that package and store chemicals.
- 3. List two cell parts found only in a plant cell.
- 4. Where in a cell do most chemical reactions take place?

NAME
~ABPARTNER

SCI#_

Cell Parts Model

No space is wasted inside a cell. Packed into the cell are all parts essential to its survival.

Procedure

- 1. Fill a sealable plastic sandwich bag halfway with tap water. Add several drops of blue food dye. Before you seal the bag, push out any remaining air.
- 2. Roll this water-filled bag into a cylindrical shape. Use two long strips of tape to secure this shape.
- 3. Fill two small plastic jewelry bags with water. Before sealing the bags, add several drops of green food coloring to each bag.
- 4. Place the water-filled sandwich bag and the two small jewelry bags into a gallon-size plastic bag.
- 5. Fill this outer bag two-thirds full with water. Push out any remaining air, and seal the bag.



Analysis

Sta	ate what each plastic bag in this model represented.
,	
De	escribe how the "central vacuole" affects the contents of your cell model.
fro	ritical Thinking Predicting Outcomes Explain how removing water om the model's central bag might affect the tension and shape of the outer astic bag.
_	

Summary Questions

- 1. Discuss the Cell Law and explain why it's discovery was so important.
- 2. How big is a human cheek cell compared to the dot over the letter "i"? (State your answer as a percent of the diameter.)
- 3. How do you know whether a cell is eukaryotic?
- 4. How long in mm is a typical onion cell?
- 5. How many layers of cells are in the Elodea leaf?
- 6. Which layer of the Elodea leaf has thick-walled cells?
- 7. Where are color pigments located in the cells of plants? Illustrate your answers.

8. What is the stomata, where are they found, and how are they important?

Gram Stain

Differential stains, which are more complex than simple ones, are used to divide bacteria into groups. Bacteria stain differentially because they differ in cell wall composition. The Gram stain separates almost all bacteria into two large groups: the Gram-positive bacteria, which stain blue (Fig. 6), and the Gram-negative bacteria, which stain pink (Fig. 7). This classification is basic to bacteriological identification.

- 1. Prepare the smear, air-dry, and heat-fix by following Steps 1 through 8 in the "Simple Stains" staining instructions above.
- 2. Flood with Hucker ammonium oxalate crystal violet for 60 seconds.
- 3. Rinse with tap water.
- 4. Flood with Gram's iodine solution for 60 seconds.
- 5. Rinse with tap water.
- **6.** Decolorize with 95% ethanol. Allow the ethanol to drip across the slide until the runoff is almost clear.
- 7. Rinse with tap water.
- 8. Flood with safranin for 60 seconds.
- 9. Rinse with tap water.
- 10. Blot carefully.
- 11. Observe with an oil immersion lens.

Morphological observations and the Gram stain are the first steps in identifying an unknown bacterium. Differential media are then used for definite identification.

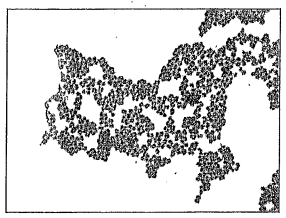


Figure 6. Gram-positive bacteria.

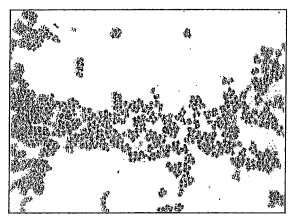
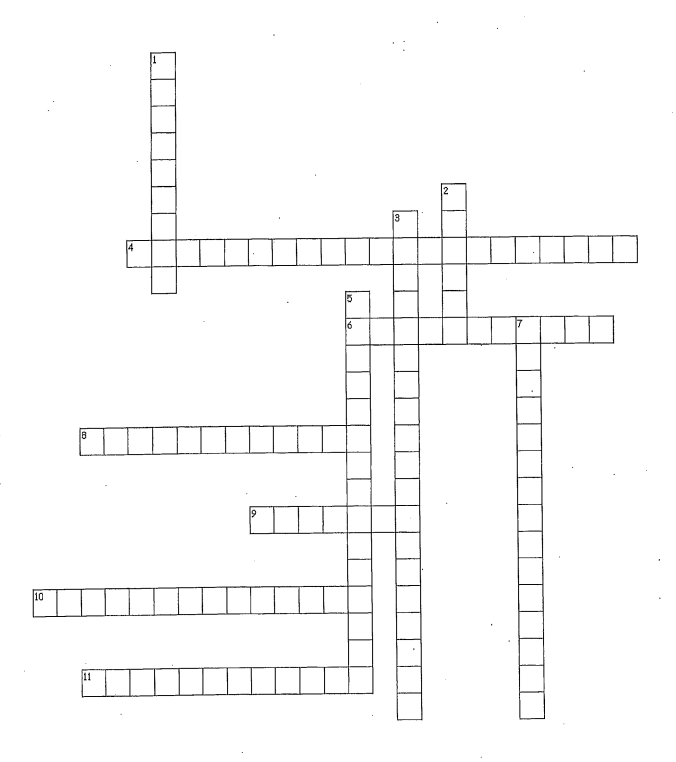


Figure 7. Gram-negative bacteria.

Chapter 8 Flashcards and Crossword- complete the crossword and then complete a flash card for each of the terms (11)



Across

- 4. a difference in the concentration of a substance across a distance
- 6, a state that exists when the concentration of a substance is the same throughout a space
- 8. a lipid that contains phosphorus and that is a structural component in cell membranes
- 9. the diffusion of water or another solvent from a more dilute solution (of a solute) to a more concentrated solution (of the solute) through a membrane that is permeable to the solvent
- 10. a protein that transports substances across a cell membrane
- 11. the basic structure of a biological membrane, composed of two layers of phospholipids

Down

- 1. the movement of particles from regions of higher density to regions of lower density
- 2. anything that serves to direct, guide, or warn
- 3. a carrier protein that uses ATP to actively transport sodium ions out of a cell and potassium ions into the cell
- 5. a molecule that is generated when a specific substance attaches to a receptor on the outside of a cell membrane, which produces a change in cellular function
- 7. a protein that binds specific signal molecules, which causes the cell to respond

Chapter 8 Cells and Their Environment

I	- Homeostasis is the maintenance of stable
interna	al conditions in a changing environment. One way that a cell maintains
	ostasis is by controlling the movement of substances across the cell membrane.
The ce	ell membrane is a gatekeeper. The cell membrane also provides structural support
to the	cytoplasm, recognizes foreign material, and communicates with other cells, all of
which	contribute to maintaining homeostasis.
П.	The cell membrane is made of phospholipids. A phospholipid
is a sp	ecialized lipid made of a phosphate "head" and two fatty acid "tails."
A. pass.	The phospholipids form a barrier through which only small, nonpolar substances can Ions and most polar molecules are repelled by the nonpolar interior of the lipid bilayer
В.	The phosphate head is and is attracted to water.
C.	The fatty acid tails are and are repelled by water.
D. layer	Because there is water inside and outside the cell, the phospholipids form a double called the lipid
	1. The nonpolar tails, repelled by water, make up the interior of the lipid bilayer.
	2. The polar heads are attracted to the water, so they point toward the surfaces of the lipid bilayer.
III.	MEMBRANE PROTEINS- Proteins in the cell membrane include cell-
surfac	e markers, receptor proteins, enzymes, and transport proteins.
	<u>Cell-surface markers</u> -act like a name tag. A unique chain of sugars acts as a marker entify each type of cell. These sugars (carbohydrates) are attached to the cell surface by sins called glycoproteins. Glycoproteins help cells work together.
B.	-enable a cell to sense its surroundings by binding to in substances outside the cell. When this happens, it causes changes inside the cell.
	-Many substances that the cell needs cannot through the lipid bilayer. Transport proteins aid the movement of these substances into out of the cell.
D.	allow reactions to take place- can break
a larg	ger molecule into 2 smaller molecules

TRANSPORT ACROSS THE MEMBRANE- There are 2 types of transport-IV. passive and active PASSIVE TRANSPORT-In passive transport, substances cross the cell membrane down their concentration gradient. No energy is required for this. Passive transport includes Small, nonpolar molecules can pass directly 1. through the lipid bilayer. This type of movement is called simple diffusion. Oxygen moves down its concentration gradient into the cell. Carbon dioxide diffuses out of the cell. Also, natural steroid hormones, which are nonpolar and fat soluble, can also diffuse across the lipid bilayer. - Many ions and polar molecules that are 2. important for cell function do not diffuse easily through the nonpolar lipid bilayer. During facilitated diffusion, transport proteins help these substances diffuse through the cell membrane. Two types of transport proteins are channel proteins, channel proteins, sometimes called pores, serve as tunnels through the a) lipid bilayer. Each channel allows the diffusion of specific substances that have the right size and charge. Ions, sugars, and amino acids can diffuse through the cell membrane through channel proteins carrier proteins- A carrier protein binds to a specific substance on one side of the cell b) membrane. This binding causes the protein to change shape. As the protein's shape changes, the substance is moved across the membrane and is released on the other side. Carrier proteins transport substances that fit within their binding site. -Water can diffuse across a selectively 3. permeable membrane in a process called osmosis. Osmosis allows cells to maintain water balance as their environment changes. Remember that in osmosis, ONLY the water molecules are free to move. If the solution is hypertonic, or has a higher solute concentration than the cytoplasm does, water moves out of the cell. The cell loses water and shrinks. If the solution is isotonic, or has the same solute concentration that the cytoplasm does, b) water diffuses into and out of the cell at equal rates. The cell stays the same size. If the solution is hypotonic, or has a lower solute concentration than the cytoplasm does, c) water moves into the cell. The cell gains water and expands in size. If left unchecked, the swelling caused by a hypotonic solution could cause a cell to burst. The rigid cell walls of plants and fungi prevent the cells of these organisms from d) expanding too much. In fact, many plants are healthiest in a hypotonic environment. Some unicellular eukaryotes have contractile vacuoles, which collect excess water inside the cell and force the water out of the cell.

Animal cells have neither cell walls nor contractile vacuoles. Many animal cells can

avoid swelling caused by osmosis by actively removing solutes from the cytoplasm

cells	concen	IVE TRANSPORT- Active transport requires energy to move substances against tration gradients. In order to move substances against their concentration gradients se energy. Most often, the energy needed for active transport is supplied directly of ATP.
	1. "pump	- In active transport, the carrier proteins do require energy to substances against their concentration gradient.
		a) The sodium-potassium pump is a carrier protein that actively transports three sodium io out of the cell and two potassium ions into the cell. This pump is one of the most important carrier proteins in animal cells. It prevents sodium ions from building up in the cell, resulting in osmosis into the cell making it burst.
		b) The concentration gradients of sodium ions and potassium ions also help transport other substances, such as glucose, across the cell membrane.
	_	- Many substances, such as proteins and polysaccharides, are too o be transported by carrier proteins. Instead, they cross the cell membrane in vesicles, which are rane-bound sacs made by pinching off of the membrane.
		a) The movement of a large substance <u>into</u> a cell by means of a vesicle. Vesicles that form by endocytosis may fuse with lysosomes or other organelles.
		b) The movement of material <u>out</u> of a cell by mean of a vesicle. These vesicles are usually from the Golgi
V. chemi		DING SIGNALS- Cells communicate and coordinate activity by sending nals that carry information to other cells.
_	et cells	naling cell produces a signal, often a molecule, that is detected by the target cell. have specific proteins that recognize and respond to the signal. These proteins are ne cell membrane (except in steroids)
В.	Neigh	aboring cells can communicate through direct contact between their membranes.
C.	Long	-distance signals are carried by hormones and nerve cells. are distributed widely in the bloodstream
VI.		CEIVING SIGNALS- cells have receptor proteins. A
~	-	ein binds only to signals that match the specific shape of its binding site (the front door will not open your neighbors front door) The outer part of the

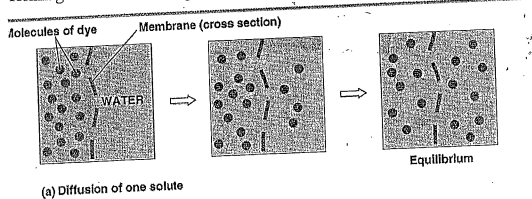
receptor protein is folded into a unique shape, called the binding site. Only the "right" shape can fit into the receptor protein while the "wrong" shape have no effect on that

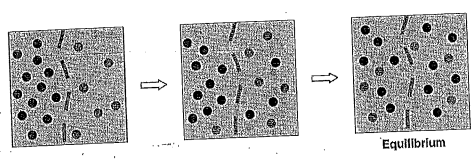
particular receptor protein. Once it binds 38 signal molecule, the receptor protein changes

its shape in the membrane. This change in shape relays information into the cytoplasm of the target cell.

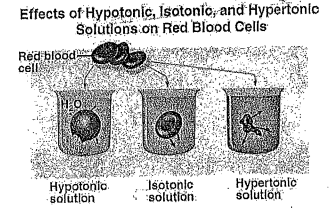
VII. RESPONDING TO SIGNALS-The cell may respond to a signal by

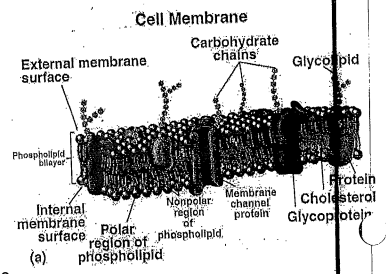
- A. changing its membrane permeability
- B. activating enzymes
- C. forming a second messenger.





(b) Diffusion of two solutes





CHAPTER 8	SEC 1	DUE DATE .	
		brane that help a cell maintain homeostasis?	
	,	·	
2. Label the two many between the two la	ain parts of the structure yers of the lipid bilayer?	e below. Which of these parts faces the area? Which faces out? Why?	
·			
3. Why are ions and	d polar molecules unabl	le to pass easily though the lipid bilayer?	
4. What are two fun	ctions of cell-surface m	narkers?	
5. Suppose a cell w	ere exposed to a drug	that caused transport proteins in the cell membra	ane
to stop working. WI	nat would happen to the	e cell?	
			e de la companya de l

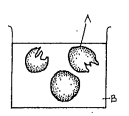
CHAPTER 8	SEC 2	DUE DATE
. Why does diffusion o	of water happen w	hen there are dissolved particles on one side of a membrane but not on
he other?		
2. Complete the follow	ing table	
TYPE OF SOLUTIO	N	DESCRIPTION
HYPERTONIC		
HYPOTONIC		
		The concentrations of solutes and water in the solution are equal to those in the cell cytoplasm.
		Water diffuses into and out of the cell at equal rates.
3. If a cell were unable	to make ATP, how	v would the cell's transport processes be affected?
	•	
eliringer Day M T W T	n F Date	*III.BX (Question) *** *** *** *** *** *** *** *** *** *
inswer <u>)</u>		

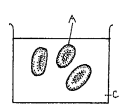
CHAPTER 8	SEC 3	DUE DATE	
1. What are two	ways cells can communica	te over long distances? What is c	ne way cells can
communicate wi	ith cells that are nearby?		
2. What is the fu	ınction of receptor proteins?		
		·	
3. What happens	s when a receptor protein b	inds to a signal molecule?	
	,	v	
4 What are three	o ways a cell may respond	when a signal molecule binds to	
	e ways a cell may respond	when a signal molecule billus to	a receptor
protein?			
	· · · · · · · · · · · · · · · · · · ·		
			•
	· · · · · · · · · · · · · · · · · · ·		·
5. Why is it impo	ortant that each receptor pro	otein binds to only one signal mol	ecule?
5. Why is it impo	ortant that each receptor pro	otein binds to only one signal mol	ecule?
5. Why is it impo	ortant that each receptor pro	otein binds to only one signal mol	ecule?
			ecule?
Ilringer: Day MiT-W		otein binds to only one signal mol	ecule?
PREHINGS V. O.R. V. W. PAYZYSS SCHLASH OF FIF			ecule?
liringer: Day MiT-Wi			ecule?
liringer: Day MiT-Wi			ecule?

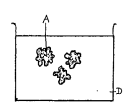
()

OSMOSIS.

ERYTHROCYTE PURE WATER 0.85% SALT SOLUTION. 2% SALT SOLUTION.





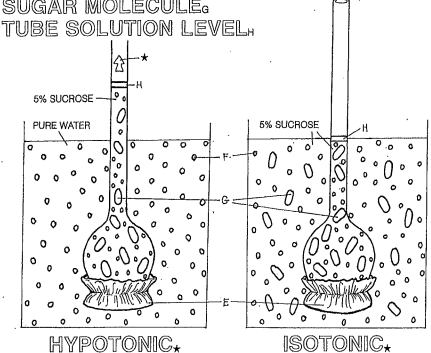


OSMOMETER*

SELECTIVELY PERMEABLE

MEMBRANE

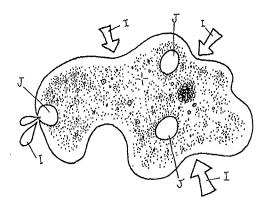
WATER MOLECULE, SUGAR MOLECULE.

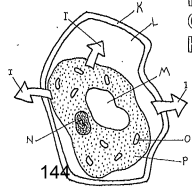


★ 10% SUCROSE 5% SUCROSE

HYPERTONIC*

AMOEBA* WATER CONTRACTILE VACUOLE, WILTING PLANT CELL* CELL WALL AIR SPACEL SHRUNKEN VACUOLE NUCLEUS CHLOROPLASTO HYALOPLASM

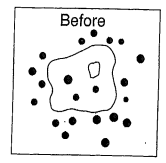


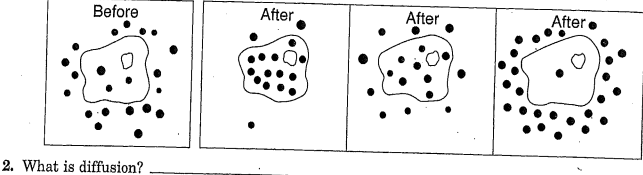


SPECIAL CELL PROCESSES

In your textbook, read about diffusion and osmosis

1. The first picture below, labeled Before, shows a cell surrounded by oxygen molecules before diffusion takes place. Each of the small black dots represents an oxygen molecule. Which of the three pictures labeled After shows where these oxygen molecules would be found after diffusion takes place? Circle your answer.





- 3. How do molecules get through the cell membrane?
- 4. What is osmosis?
- 5. Which way would the water molecules move in the following situations?
 - a. cucumber slice is placed in salt water _____
 - b. salt is poured on a snail _____
 - c. vegetables are sprinkled with water _____
 - d. potato slice is placed in pure water _____
- 6. Circle the letter in front of the sentence that best explains the process of osmosis.
 - a. Osmosis is the movement of water into or out of a cell from where it is in large amounts to where it is in small amounts.
 - b. Osmosis is the movement of water into or out of a cell from where it is in small amounts to where it is in large amounts.
 - c. Osmosis is the movement of salt into or out of a cell from where it is in large amounts to where it is in small amounts.

Pre-Lab Questions

Why must you use 2 beakers?
What is osmosis? If there is no water, can osmosis occur? What is the difference
between osmosis and diffusion.
What is the egg white made of? What is corn syrup made of?
Why must you soak the egg in and why are you doing this?
What is a HYPOTONIC solution?
What is a HYPERTONIC solution?
What is a ISOTONIC solution?
What are your predictions for this lab? Be specific.

Observing Osmosis in Eggs

Some chemicals can pass through a cell membrane, but others cannot. Furthermore, not all chemicals can pass through a cell membrane with equal ease. The cell membrane determines which chemicals can diffuse into or out of a cell.

As chemicals pass into and out of a cell, they move from areas of high concentration to areas of low concentration. Cells in *hypertonic* solutions have solute concentrations lower than the solution that bathes them. This concentration difference causes water to move out of the cell into the solution. Cells in *hypotonic* solutions have solute concentrations greater than the solution that bathes them. This concentration difference causes water to move from the solution into the cell. The movement of water into and out of a cell through the cell membrane is called *osmosis*.

In this lab, you will use a model of a living cell to predict the results of an experiment that involves the movement of water through a membrane.

OBJECTIVES

- Explain changes that occur in a cell as a result of diffusion.
- Distinguish between hypertonic and hypotonic solutions.

MATERIALS

- safety goggles, lab apron, protective gloves
- balance
- beakers, 250 mL (2)
- beakers, 600 mL (2)
- corn syrup
- distilled water

- eggs (2)
- paper towels (2)
- tablespoon or tongs
- vinegar, 400 mL
- wax pencil



Procedure

DAY 1: SOAKING EGGS IN VINEGAR

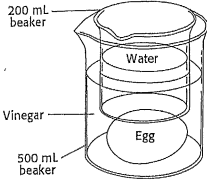
- 1. Label one 600 mL beaker "Egg 1: water" and the other 600 mL beaker "Egg 2: syrup." Also label the beakers with the initials of each member of your group.
- 2. Measure the mass of each of two eggs to the nearest 0.1 g, and record your measurements in the second column of Table 1. CAUTION: Uncooked eggs may contain harmful bacteria. Do not touch your face after you have handled raw eggs. Clean up any material from broken eggs immediately. Wash your hands with soap and water after handling the eggs.
- 3. Put on safety goggles and a lab apron. Pour 200 mL of vinegar into each labeled beaker. Using a tablespoon or tongs, place an egg into each beaker. Always return each egg to the same beaker.

TABLE 1 EGGS IN VINEGAR

Egg	Mass of fresh egg with shell	Observations after 24 h	Mass after 24 h in vinegar
1			
2			

4. Place a 250 mL beaker containing 100 mL of water on each egg to keep it submerged, as shown in Figure 1. Add more vinegar if the egg is not covered by the vinegar already in the beaker. If some vinegar spills over when the 250 mL beaker is placed on the egg, carry the beaker carefully to the sink and pour out some vinegar. Store the beakers for 24 hours in the area specified by your teacher.

FIGURE 1 EGG IN VINEGAR



5. Clean up your work area and wash your hands before leaving the lab.

DAY 2: SOAKING EGGS IN TWO LIQUIDS

- 6. After 24 hours, observe the eggs. Record your observations in Table 1.
- 7. Put on safety goggles and a lab apron. Label two separate sheets of paper towel "Egg 1" and "Egg 2." Pour the vinegar from the beakers into the sink. Using a tablespoon or tongs, remove the eggs and rinse them with water. Place each egg on the appropriately labeled paper towel. Measure the mass of each egg, and record the measurement in the last column of **Table 1**.
- 8. Return Egg 1 to its beaker, and add water until the egg is covered. Return Egg 2 to its beaker, and add corn syrup until the egg is covered. Store the beakers for 24 hours in the same place as before.
- 9. Clean up your work area and wash your hands before leaving the lab.

DAY 3: MEASURING CHANGES IN THE EGGS

- 10. Predict how the mass of each egg has changed after 24 hours in each liquid. (Hint: An egg is surrounded by a membrane. Inside the membrane, the egg white consists mainly of water and dissolved protein. The yolk consists mainly of fat and water. Corn syrup is sugar dissolved in water. The protein, fat, and sugar are solutes.) Record your predictions in Table 2.
 - What will have occurred if your egg gains or loses mass?
- 11. Observe your eggs. Record your observations in **Table 2**. Measure and record the final masses of the two eggs.

TABLE 2 EGGS SOAKED IN TWO LIQUIDS

Egg	Liquid	Predicted change after 24 h	Observations after 24 h	Final mass of egg
1				
2				

- 12. Dispose of your materials according to your teacher's instructions.
- 13. Clean up your work area, and wash your hands before leaving the lab.

Analysis

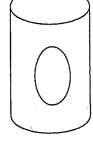
What	caused the change in appear	ance in Egg 1 after it soaked in water?
·		

3.	What caused the mass of the egg to increase after soaking in the vinegar solution?
4.	What material seems to have moved through the membrane of Egg 2 after it soaked in the corn syrup? In what direction did the material move?
5.	How did your results in step 11 compare with your prediction?
6.	Which egg was in a hypertonic solution? Explain what you used for evidence.
7.	Which egg was in a hypotonic solution? Explain what you used as evidence.

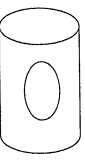
8. What do you think would happen to a red blood cell placed in a test tube of distilled water? Explain using principles of osmosis why you believe this would occur.

9. Draw a diagram of where water molecules move from a cell in a)hypertonic solution b) hypotonic solution c) isotonic solution.

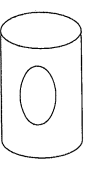
A) hypertonic



B) hypotonic



C) isotonic



Pre-Lab Questions

How many cell models will you be making? What is the difference in them?
What will you use to measure volume of the models?
what is the SI for volume? What is the formula to find the volume of an object?
What is the surface area? What is the surface area of a cube? What is the surface area of a sphere?
What are your predictions for this lab? Which model will have the greatest volume? How does this apply to a cell? Is this an important concept and why.

Modeling Cells: Surface Area to Volume

Are there limits to how large a cell can grow? Everything that enters and exits a cell passes through the cell membrane. As the size of a cell increases, its surface area increases, but so does its volume. Consider how people enter a crowded event at a large stadium. Everyone funnels through a few gates. In a larger stadium, it takes people longer to move in and out. Similarly, in a larger cell, it takes materials longer to reach their destination inside the cell. This means that it is more difficult for a large cell to have its needs met through the cell membrane. In this lab, you will examine surface area-to-volume ratios on a small scale, using model cells. You will use the collected data to draw conclusions about why this ratio might limit the size of a cell.

OBJECTIVES

- Prepare and compare various cell models.
- Calculate surface area and surface area-to-volume ratios.
- Use your data to form conclusions about size limitations on cells.

MATERIALS

- calculator (optional)
- cell model patterns (3)
- funnel
- graduated cylinder, large
- metric ruler

- paper, heavy
- · safety goggles
- sand
- scissors
- tape



Procedure

- 1. Put on your safety goggles. Trace and cut out three cell models. Your teacher will provide you with the patterns or dimensions for each model. Fold the models to form three-dimensional shapes, as in **Figure 1**. Use tape to keep each model together.
- 2. Use the ruler to measure the length, width, and height dimensions of each model. Record the dimensions in Table 1.
- 3. Calculate the total surface area for each model. To do this, find the area of each side (length 3 width), then multiply that number by 6. Enter the data in Table 1.

FIGURE 1
CELL MODEL

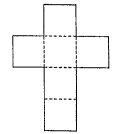




TABLE 1 MODEL CELL CALCULATIONS

Cell	Dimensions (cm)	Surface area (cm²)	Volume (cm³)	Surface area-to- volume ratio
A				
В				
С				

- 4. Use the funnel to fill each model with sand. Use the ruler to level off the sand.
- 5. Find the volume of sand in each model, and enter the data in **Table 1**. You can do this by using either of two methods.
 - a. Measure the amount of sand in each model by pouring the sand through a funnel into a graduated cylinder.
 - b. Calculate the volume, using the following formula: volume = length × width × height
- 6. Calculate the surface area-to-volume ratio for each model. Use the following formula:

surface area ÷ volume = surface area-to-volume ratio

Record the values in Table 1.



Clean up your materials and wash your hands.

NA	MESCI#POINTS:
1.	Why do you need to multiply by 6 in step 3?
2.	Which cell model has the largest surface area?
	The largest volume?
	The largest surface area-to-volume ratio?
	Which of these measurements is the most important for hypothesizing whether
	a cell would be able to get all the oxygen and food it needs?
3.	Which model cell is likely to be most efficient at getting nutrients to all of the cell parts? Explain your answer in terms of surface area-to-volume ratios.
4.	What formula did you use to get the volume?
5.	What formula did you use for surface area?
6.	On back of this sheet, create the same table as in lab procedures above. Be
ur	e to use a ruler to make the table and include all the information that is in the
ab	le above. Highlight (in yellow) the model that would give you the MOST
T F	FICIENT ceil

Pre-Lab Questions

What are your predictions for this lab? Be specific.	
What is a ISOTONIC solution?	
What is a HYPERTONIC solution?	
What is a HYPOTONIC solution?	
What is distilled water? Is it different than tap water?	
What is the major component of grape flesh? What is grape j	
What is osmosis? If there is no water, can osmosis occur? Wheetween osmosis and diffusion.	
Why is it important to dry the grapes?	

Osmosis

You will observe the movement of water into or out of a grape under various conditions.

Procedure

- 1. Make a data table with four columns and three rows.
- 2. Fill one cup with a salt solution. Fill a second cup with grape juice. Fill a third jar with distilled water. Label each cup with the name of the solution that it contains.
- 3. Use a balance to find the mass of each of three grapes. Place one grape in each cup, and cover the cups with wrap. Place in your period's tray.
- 4. Predict whether the mass of each grape will increase or decrease over time. Explain your predictions on prelab sheet.
- 5. After 24 hours, remove each grape from its jar, and dry the grape gently with a paper towel. Using the balance, find each grape's mass again. Record your results.

	Identify the solutions in which osmosis occurred.
	How did you determine whether osmosis occurred in each of the three solutions?
•	Did the mass of each grape change as you had predicted? Why or why not?
ŀ.	Which solution was hypotonic? Hypertonic? Isotonic?
	157

	SALT WATER	GRAPE JUICE	DISTILLED WATER
DAY 1			
DAY 2			
			:
CHANGE IN MASS (+ IF IT	·		
INCREASED, -,IF DECREASED NO			
CHANGE)			

Why do cells divide?

When cells grow to a certain size, their rate of growth slows until they stop growing. At this point, they have reached their size limit. A cell that has reached its size limit divides into two smaller cells. In this lab, you will explore one of the factors that limit cell size: the relationship between the size of the cell—specifically, its surface area and volume—and how efficiently substances diffuse across its cell membrane.

Objectives

- · Model cells of different sizes with agar cubes.
- Model the diffusion of materials across a cell membrane.
- Calculate the surface area-to-volume ratio for model cells.
- Form a hypothesis about how cell division affects a cell's ability to absorb materials.

Materials

agar
beaker
timer
calculator
plastic ruler
100 mL 0.1M solution of hydrochloric acid
kitchen knife
plastic spoons
paper towels

Safety Precautions

WARNING: Use caution when handling hydrochloric acid.

Procedure

Part A. Setting Up the Experiment

- 1. Read and complete the lab safety form.
- 2. Obtain a block of agar containing phenolphthalein from your teacher. Recall that phenolphthalein turns pink in the presence of a base. It will become colorless in an acid.
- 3. Use a ruler to measure and a kitchen knife to cut three blocks out of the agar. One should be 3 cm on each side, one should be 2 cm on each side, and one should be 1 cm on each side.
- 4. Figure 1 Place the three agar cubes inside the beaker. Cover with 100 mL dilute hydrochloric acid solution.

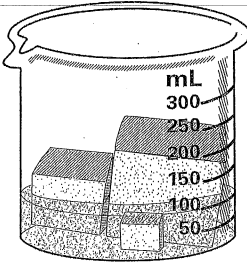


Figure 1

- 5. Leave the agar blocks in the dilute hydrochloric acid for a total of 10 min. Use a spoon to turn them every few minutes to ensure that they are soaking evenly.
- 6. Complete the data table on the next page.

Why do cells divide?

Part B. Measuring Diffusion

- 1. After 10 min, carefully use the plastic spoons to remove the agar blocks. Blot them dry with paper towels. Use care not to splash HCl on skin; it will cause burns.
- 2. Use the edge of the plastic ruler to cut each block in half. Measure the depth of the uncolored area in centimeters, recording the measurement to the nearest millimeter. This shows the depth of diffusion. Record these values in Table 1.
- 3. Complete Table 1, and answer the questions that follow.
- **4.** You might need the following formulas: surface area = length × width × number of surfaces
 - volume of a cube = $length \times width \times height$ Use a calculator for your calculations if necessary.
- 5. Wash your hands with soap and water, and dispose of the materials as instructed by your teacher.

Data and Observations

Table 1

Table 1 Agar Data				
Cube Size	Surface Area	Volume	Ratio	Depth of Diffusion
3 cm/side				
2 cm/side				
1 cm/side			,	

Analyze and Conclude

1.	Is the distance of diffusion the same for all of the blocks? Explain.
2.	Based on your answer to the question above, do you think that the depth of diffusion is the same in all cells? Explain.
	List the agar cubes in order of size, from largest to smallest. Then list them in order of surface area-to-volume ratio (from largest ratio to smallest ratio). How do these lists compare?

Why do cells divide?

5.	Which block has the greatest surface area-to-volume ratio—the onion cube or the 3 cm/side cube you used in this lab?
6,	What is the relationship between surface area-to-volume ratio and diffusion across a cell?
7.	What happens to diffusion as a cell grows?
8.	Error Analysis What are some possible sources of error in your experiment?
	Form a hypothesis to explain how cell division affects a cell's ability to absorb the material necessary

Inquiry Extensions

- 1. Which cells in the human body divide most frequently? Why is this? What activities or conditions spur cell division? What slows it down?
- 2. During adolescence the human body grows at a rate faster than at any other time after infancy. Explain how what you learned in this lab plays out in the human body during adolescence.

Osmosis and Diffusion

Student Study and Analysis Sheets

Introduction

Every plant and animal cell has a membrane which acts as a barrier between the "outside" environment and the cell's cytoplasm. Membranes are selectively permeable, allowing only certain molecules to enter and exit the cytoplasm freely.

In 1827, Scottish scientist Robert Brown found that tiny particles suspended in water moved in small, quick movements. This phenomenon, known as Brownian movement or random motion, illustrates that molecules are in a state of constant, random motion in all liquids and gases; they move in an undirected fashion, bouncing off other molecules.

Because molecules are in constant motion, they bounce off each other and move toward an area of fewer molecules. This action, known as diffusion, is the movement of molecules from an area with a high concentration of molecules to an area of low concentration of molecules.

When a concentration gradient (a high concentration of molecules in one area and a low concentration in another) exists, diffusion will take place, and molecules will move until an equilibrium is reached. For example, when a bottle of hydrogen sulfide, which smells like rotten eggs, is opened on one side of a room, the smell can quickly be detected on the other side. The bottle has a high concentration of hydrogen sulfide; the room has a low concentration. The hydrogen sulfide diffuses to the less-concentrated area until an equilibit room.

Like all molecules, water molecules are in constant motion, moving from areas of high concentration to areas of low concentration. Water moves through a selectively permeable membrane whenever there is an unequal concentration of water on either side of the membrane, until an equilibrium is reached. This process is called cosmosis. The osmotic process is a special case of diffusion involving the movement of a solvent, such as water, rather than substances dissolved in the solvent (solutes).

Sometimes the water molecules carry other molecules along with them. The action of the cell transporting substances in and out of its cell membrane is called active transport. The cell uses energy derived from ATP or a protein to move the solutes into or out of the cell.

Objective

To create a model of a cell membrane to observe osmosis and diffusion.

Materials Needed per Lab Group

Cup, 9 oz.
 Glucose Testing Strips
Dialysis Tubing, 1 ft.
Graduated Cylinder
Goggles
Gloves
Aprons

Shared Materials

Glucose Solution Starch Solution Iodine Potassium Iodide

Procedure

Safety: Wear goggles, gloves, and apron when conducting this investigation.

- 1. Fill the plastic cup three-quarters full with water.
- 2. Test the water for glucose by dipping a glucose test strip in the water. Record the data in the table below.
- 3. Add 20 drops of IKI solution to the cup of water. Note the color of the water and record it in the table. Note: lodine potassium lodide is a corrosive/irritant. Store away from other chemicals.

Read MSDS before use.

- 4. Hold the section of dialysis tubing under running water until it is pliable.
- 5. Once the tubing is pliable, tie a knot in one end.

Note: Handle the tubing carefully; make sure that you do not rip the tube.

- 6. Open the tubing by rubbing the untied end between your fingers.
- 7. Pour 15ml soluble starch solution into the tubing.
- 8. Pour 15ml glucose solution into the tubing.
- 9. Carefully tie a knot in the open end to form a bag. Note the color of the solution in the tubing and record the color in the table.
- 10. Rinse the tube thoroughly to wash off any glucose or starch that may have spilled onto the outside of the tube
- 11. Place the dialysis tube in the cup of water-IKI solution.
- 12. Observe the tube for 15 minutes. Record the final color of the solutions in the tubing and in the cup.
- 13. Test the water-IKI solution once again for glucose with the second glucose test strip. Record the results in the table.

Analysis

•	1	Color Glucose			se	Starch	
C -trinor	Contents	Initial	Final	initial	Final	Initial	Final
Container	COntents				,		
Cup							
Dialysis Tubing							. !
		<u>, , , , , , , , , , , , , , , , , , , </u>		•			

Note: Both glucose and starch are present, and will remain, in the tubing; only enough will move until an equilibrium is reached between the outside and the inside of the membrane, leaving a certain amount inside the tubing.

Questions

1. Which substance(s) migrated into or out of the dialysis tubing? How do you know?

2. Which, if any, substance(s) did not diffuse through the membrane? How do you know?

3. What is osmosis? How can you tell if osmosis occurred in the dialysis tube?

4. What is selective permeability?

5. Molecules of similar substances are about the same size, whereas molecules of different substances are different sizes. From the results of the experiment, is it possible to determine the relative sizes of molecules that did or did not diffuse across the dialysis membrane?

6. Can it be said that the dialysis membrane is similar to a plasma membrane?

Diffusion and Cell Membranes

Procedure

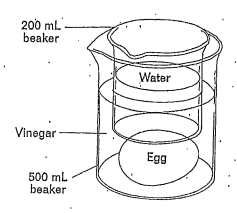
- 1. Label one 600 mL beaker "Egg 1: water" and the other 600 mL beaker "Egg 2: syrup." Also label the beakers with the initials of each member of your group. Measure the mass of each of two eggs to the nearest 0.1 g, and record your measurements in Table 1 below. CAUTION: When handling raw eggs, clean up any material from broken eggs immediately. Wash your hands with soap and water after handling the eggs.
- 2. Put on safety goggles and a lab apron. Pour 200 mL of vinegar into each labeled beaker. Using a tablespoon or tongs, place an egg into each beaker. Note: Always return each egg to the same beaker.

TABLE 1: EGGS IN VINEGAR

Egg	Mass of fresh egg with shell	Observations after 24 h	Mass after 24 h in vinegar
1			,
2			

- 3. Place a 250 mL beaker containing 100 mL of water on each egg to keep it submerged as shown in Figure 1 below. Add more vinegar if the egg is not covered by the vinegar already in the beaker. If some vinegar spills over when the 250 mL beaker is placed on the egg, carry the 600 mL beaker carefully to a sink and pour vinegar some out. Store your beakers for 24 hours in the area specified by your teacher.
- 4. Clean up your work area and wash your hands before leaving the lab.

FIGURE 1



Observing Plasmolysis in Onion Skin Cells

Materials

onion section

scalpel

forceps

15% NaCl solution (dissolve 15 g NaCl in 85 mL of water)

microscope slides

coverslips

distilled water

dropping pipets

absorbent paper

microscopes

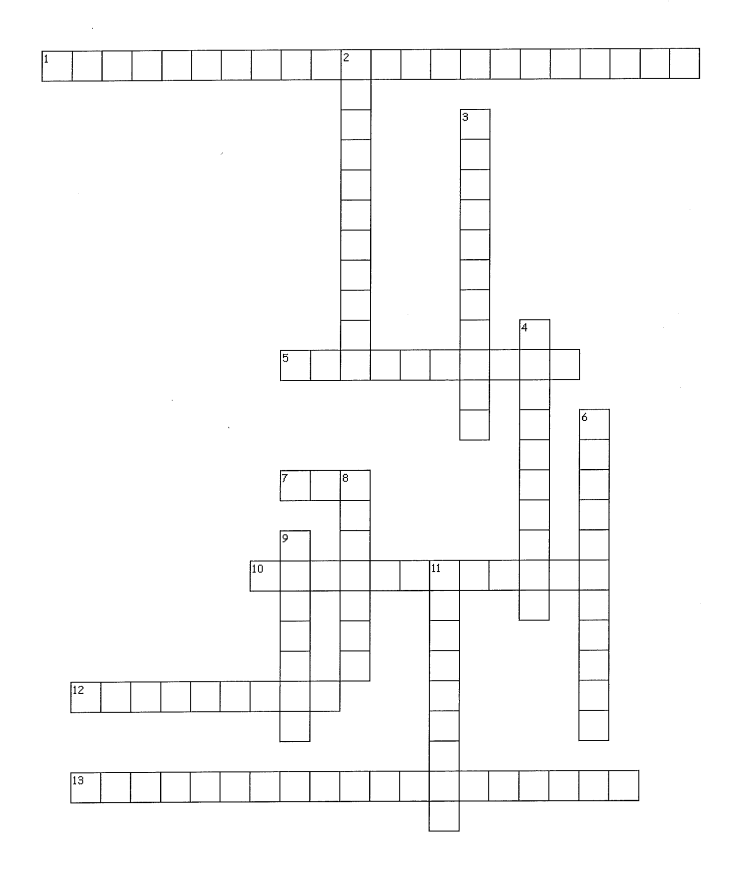
Procedure

- 1. An onion is made up of several layers of thick scale leaves. You will need one portion of scale from a cut section of an onion. Working on the concave surface of the scale, cut out a section about 1 cm². Use forceps to remove the epidermal layer (onion skin) from the concave surface of the section you have cut. Place the onion epidermis on a microscope slide and smooth it to remove as many wrinkles as possible. Add one or two drops of distilled water and a coverslip, and observe under a microscope.
- 2. You have probably observed onion cells before, but reacquaint yourself with their structure and appearance. Note especially the cell walls and the location of the nuclei.
- 3. Remove the slide from the microscope. Add a drop or two of 15% NaCl solution to one edge of the coverslip. Use a piece of absorbent paper to absorb water from the opposite edge of the coverslip. This should "pull" the salt solution under the coverslip and bring it in contact with the onion cells.
- 4. Observe under the microscope for several minutes for signs of change in the onion cells. If after several minutes you have seen no change, remove the coverslip, blot away excess water, and add one or two drops of 15% NaCl solution directly to the square of onion epidermis. Replace the coverslip and observe again.
- 5. Once you have observed plasmolysis, remove the coverslip and blot away the excess water. Flood the epidermis with distilled water and blot again. Add another drop of distilled water and replace the coverslip. Observe under the microscope. Do you see any change that would indicate that water is entering the cells?

photosynthesis		_		 TP.
cellular respiration				 N. Ave
ATP				
ATPsynthase	·			
electron transport chain				
thylakoid				
pigment				
chlorophyll				
Calvin cycle				
glycolysis				
anaerobic				
aerobic			=	
Krebs cycle				Alteria.
fermentation				

Ch 9 Crossword / Vocab Flash Cards- complete the crossword and make a flashcard for

each term with the word on one side and the definition on the back



Across

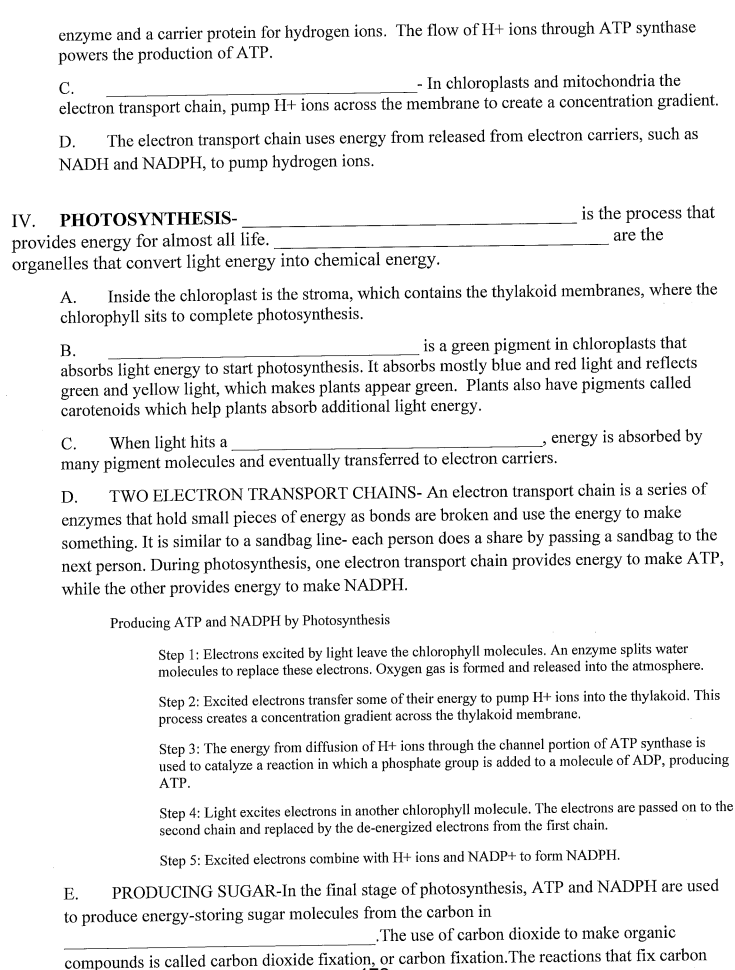
- 1. a series of molecules, found in the inner membranes of mitochondria and chloroplasts, through which electrons pass in a process that causes protons to build up on one side of the membrane
- 5. a series of biochemical reactions that convert pyruvate into carbon dioxide and water
- 7. adenosine triphosphate, an organic molecule that acts as the main energy source for cell processes; composed of a nitrogenous base, a sugar, and three phosphate groups
- 10. the breakdown of carbohydrates by enzymes, bacteria, yeasts, or mold in the absence of oxygen
- 12. describes a process that does not require oxygen
- 13. the process by which cells produce energy from carbohydrates

Down

- 2. an enzyme that catalyzes the synthesis of ATP
- 3. a green pigment that is present in most plant and algae cells and some bacteria, that gives plants their characteristic green color, and that absorbs light to provide energy for photosynthesis
- 4. the anaerobic breakdown of glucose to pyruvate, which makes a small amount of energy available to cells in the form of ATP
- 6. a biochemical pathway of photosynthesis in which carbon dioxide is converted into glucose using ATP and NADPH
- 8. a substance that gives another substance or a mixture its color
- 9. describes a process that requires oxygen, the most efficient respiration
- 11. a membrane system found within chloroplasts that contains the components for photosynthesis

Chapter 9 Photosynthesis and Cellular Respiration

		MICAL ENERGY- Organisms require a constant source of energy. Energy is
neede	d for o	organisms to maintain their homeostasis. is the process of maintaining internal order and
balanc	e ever	n when the environment changes.
	A. carbor	Photosynthesis is the process by which plants, algae, and some bacteria use sunlight, a dioxide, and water to produce and
		and oxide, and water to produce and Organisms that are able to perform
	photos on ear	synthesis, such as plants, are make all lood
	B. autotro	Organisms that cannot make their own food must absorb food molecules made by ophs, eat autotrophs, or eat organisms that consume autotrophs.
	C. bonds	Cells use molecules(from food ingested) to release the energy stored in the chemical of food.
Becau	organi ise org	ABOLISM AND THE CARBON CYCLE- Metabolism is either using energy to c molecules or breaking down organic molecules in which energy is stored. anic molecules contain (the definition an organism's metabolism is part of Earth's
	A. conve	Energy enters an ecosystem when organisms use sunlight during photosynthesis to rt molecules (gas) into
	В.	Through the process of cellular respiration, cells make the carbon in glucose into stable a dioxide molecules and produce
	C. molec	Energy is also released and used to make ATP (adenosine triphospate), an organic rule that is the main energy source for cell processes.
III. series		NSFERRING ENERGY- In cells, chemical energy is gradually released in a emical reactions that are assisted by enzymes.
	A.	ATP Energy is stored as glucose,
		stored as ATP. Energy is stored as glucose,
	starch	es and lipids. ATP is a nucleotide made up of a chain of
	broker	groups. When ATP is used, a phosphate group is is left.
		- In many cells, ATP synthase, an enzyme
	that m	volzog ATP and recycles ADP by adding a phosphate. ATP synthase acts as both an



dioxide are light-independent reactions, sometimes called dark reactions. The most common method of carbon fixation is the Calvin cycle.

- V. **FACTORS THAT AFFECT PHOTOSYNTHESIS-** Light intensity, carbon dioxide concentration, and temperature are three environmental factors that affect photosynthesis.
 - A. In general, the rate of photosynthesis increases as light intensity increases until all of the pigments in a chloroplast are being used.
 - B. The concentration of carbon dioxide affects the rate of photosynthesis in the same manner
 - C. Photosynthesis is most efficient in a certain range of temperatures.
- VI. **CELLULAR RESPIRATION** Respiration is not the cell breathing. Respiration is the process of breaking down food to release energy. There are 2 parts. Glycolysis is 1st. then if oxygen is present, aerobic respiration takes place (this includes the Krebs Citric Acid Cycle and electron transport chain. If oxygen is absent, anaerobic fermentation takes place.
 - A. **GLYCOLYSIS** In glycolysis, the first step, enzymes break down one six-carbon molecule of glucose into two three-carbon pyruvate molecules. The breaking of a sugar molecule by glycolysis results in a net gain of two ATP molecules.
 - 1. The primary fuel for cellular respiration is glucose. Fats can be broken down to make ATP. Proteins and nucleic acids can also be used to make ATP, but they are usually used for building important cell parts.
 - 2. In aerobic respiration, the pyruvate product of glycolysis undergoes another series of reactions to produce more ATP molecules. In anaerobic respiration, fermentation begins.
 - B. **AEROBIC RESPIRATION-** This is done with oxygen- the breakdown of stored sugars to release ATP for energy. Aerobic respiration is more efficient than anaerobic. The total yield of energy-storing products from one time through the Krebs cycle is one ATP, three NADH, and one FADH2.
 - 1. The first stage of aerobic respiration is the Krebs cycle, a series of reactions that produce electron carriers. The total yield of energy-storing products from one time through the Krebs cycle is one ATP, three NADH, and one FADH2.
 - 2. The electron carriers then enter an electron transport chain
 - 3. Up to 34 ATP molecules can be produced from one glucose molecule in aerobic respiration.
 - 4. Krebs Cycle
 - a) Occurs in mitochondria
 - b) Starts with pyruvate

- c) Produces 1 ATP
- 5. Electron Transport Chain
 - a) Takes place in the inner membranes of mitochondria, where the enzyme ATP synthase is located.
 - b) Electron carriers, produced during the Krebs cycle, transfer energy through the electron transport chain.
 - c) Energy from the electrons is used to actively transport hydrogen ions out of the inner mitochondrial compartment.
 - d) Hydrogen ions diffuse through ATP synthase, providing energy to produce several ATP molecules from ADP.
- C. **FERMENTATION-** Only occurs under anaerobic conditions because it is not cost efficient for the cells. Fermentation enables glycolysis to continue supplying a cell with ATP in anaerobic conditions.
 - 1. To make ATP during glycolysis, NAD+ is converted to NADH. Organisms must recycle NAD+ to continue making ATP through glycolysis.
 - 2. In lactic acid fermentation, pyruvate is converted to lactic acid in a process. During vigorous exercise, lactic acid fermentation also occurs in the muscles of animals, including humans.
 - 3. During alcoholic fermentation, one enzyme removes carbon dioxide from pyruvate. A second enzyme converts the remaining compound to ethanol, recycling NAD+ in the process.

VII. Efficiency of Cellular Respiration

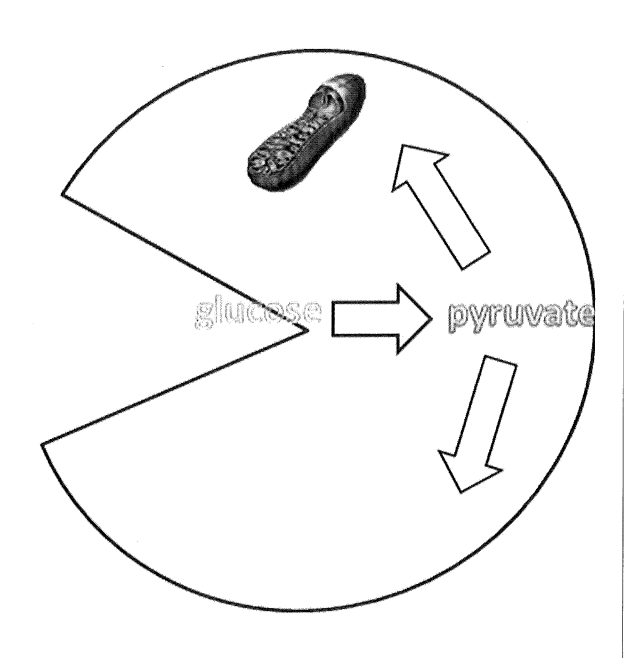
CELLULAR RESPIRATION		LOCATION	STARTING MATERIALS	PRODUCTS (NOT INCLUDING ATP)	NET ATP GAIN
GLYCOLYSIS (OCCURS UNDER ALL CONDITIONS)		CYTOPLASM	GLUCOSE	PYRUVATE	2
AEROBIC	KREBS CYCLE	MITOCHONDRIA	PYRUVATE	NADH	1
	ELECTRON TRANSPORT CHAIN	MITOCHONDRIAL MEMBRANE	HYDROGEN IONS, ATP SYNTHASE	2 WATER, NADH	32
ANAEROBIC	FERMENTATION	CYTOPLASM	PYRUVATE	ALCOHOL/LACTIC ACID, NADH	2

CHAPTER 9	SEC 1		DUE DA	TE		_•
1. How does ATP	synthase produc	e ATP?			: :	
2. How does the ca	arbon cycle deliv	er energy to org	ganisms?			
						.
3. How do organis	ms that are not a	autotrophs get e	nergy?			
4. In cells, glucose the reason for this		h oxygen in a s	eries of steps ir	nstead of all	at once. W	/hat is
The reason for this	!					
5. What happens i	n an electron tra					
			-11-2			
6. Name two mole	cules that can re	lease energy as	s part of the ele	ctron transp	ort chain.	
ellringer:Day M T W Th	F Date	Question				
nswer						
		175				

	SEC 3 DUI	E DATE .
Why is glycolysis consid	lered an anaerobic process, but the	electron transport chain is not
Fill in the blank spaces i	in the table below.	
Process	Description	Overall number of ATP molecules produced per molecule of glucose
	Glucose is broken down into two pyruvate molecules.	
	Pyruvate is used to produce NADH, ATP, and FADH2; carbon dioxide is produced as pyruvate breaks down.	
	Energy from electrons in NADH and FADH2 is used to produce ATP;	
	water is produced as hydrogen and oxygen accept electrons.	
s. Organism A can carry o	water is produced as hydrogen and	can carry out only glycolysis.
Which organism will be ab	water is produced as hydrogen and oxygen accept electrons. Out cellular respiration. Organism B on the energy in a management of the energy in a management.	olecule of glucose? Explain yo
Which organism will be ab	water is produced as hydrogen and oxygen accept electrons. out cellular respiration. Organism B	olecule of glucose? Explain yo
Which organism will be ab	water is produced as hydrogen and oxygen accept electrons. Out cellular respiration. Organism B on the energy in a management of the energy in a management.	olecule of glucose? Explain yo
Which organism will be ab answer. (Hint: Remember	water is produced as hydrogen and oxygen accept electrons. Out cellular respiration. Organism B on the energy in a management of the energy in a management of the energy in a management.	olecule of glucose? Explain yo
Which organism will be ab answer. (Hint: Remember	water is produced as hydrogen and oxygen accept electrons. Out cellular respiration. Organism B on the color of the energy in a manner that ATP is the main source of energy.	olecule of glucose? Explain yo
Which organism will be ab answer. (Hint: Remember	water is produced as hydrogen and oxygen accept electrons. Out cellular respiration. Organism B of the to use more of the energy in a material that ATP is the main source of energy portant? Does your body ever unde	rgy for cellular processes.) rgo fermentation? When?
Which organism will be abanswer. (Hint: Remember 4. Why is fermentation imp	water is produced as hydrogen and oxygen accept electrons. Out cellular respiration. Organism B on the color of the energy in a manner that ATP is the main source of energy.	rgy for cellular processes.) rgo fermentation? When?
Which organism will be abanswer. (Hint: Remember 4. Why is fermentation imp	water is produced as hydrogen and oxygen accept electrons. Out cellular respiration. Organism B of the to use more of the energy in a manner that ATP is the main source of energy in a manner that Poes your body ever unde	rgy for cellular processes.) rgo fermentation? When?
Which organism will be abanswer. (Hint: Remember 4. Why is fermentation imp	water is produced as hydrogen and oxygen accept electrons. Out cellular respiration. Organism B of the to use more of the energy in a more that ATP is the main source of energy in a more that ATP is the main source of energy in a more than the portant? Does your body ever unde	rgo fermentation? When?

Ch 9 Review

1.	Most of the energy used by life on Earth comes from the
2,	Light energy is converted to chemical energy through the process of
3.	During the final stage of photosynthesis, sugars are produced from
4.	As light intensity increases, the rate of photosynthesis does what?
5.	Low temperatures may cause photosynthesis to what?
6.	What environtal factors does not affect the rate of photosynthesis?
7.	The name of the process that takes place when organic compounds are broken down in the absence of
	oxygen is
8.	Fermentation enables glycolysis to continue under conditions
9.	If oxygen is absent during the second stage of cellular respiration, what happens?
10.	Cells produce ATP most efficiently in the presence of what?
11.	what is an autotroph?
12.	What is a heterotroph?
13.	Are the following auto or hetero- trophs: plants, algae, prokaryotes, animals
14.	ATP is what biomolecule? What is it composed of?
15.	When cells break down food molecules, what happens?
16.	The space inside the inner membrane of a chloroplast is called the
17.	The major atmospheric by-product of photosynthesis is
	The source of oxygen produced during photosynthesis is
19.	Carbon dioxide is converted into organic compounds in what part of the cell?
20.	what is glysolysis?
21.	. What is respiration?
	. What is photosynthesis?
23.	. The total amount of ATP that a cell gains for each glucose molecule depends on the presence of



Ch 9 PreLab

1. What are stomata? What is their function?
2. Write the formula for photosynthesis
3. Write the formula for respiration
4. Where does photosynthesis occur?
5. Where does respiration occur?
6. What are you using to make the slide? Explain how you will visualize the stomata
7. What are you painting on the leaf and what do you do when that is dry?

Ch 9 lab Stomata and Photosynthesis

Introduction

Plants and animals both have a layer of tissue called the epidermal layer. Plants have special pores called stomata to allow passage of material. The stomata pores are surrounded on both sides by jellybean shaped cells called guard cells. Unlike other plant epidermal cells, the guard cells contain chlorophyll to do photosynthesis. This allows the cells to expand/ contract to open or close the stomata. Guard cells also close when dehydrated. This keeps water in the plant from escaping. The opening or closing of guard cells can be viewed in a microscope by adding different water concentration to the leaf tissue.

Most stomata are on the lower epidermis of the leaves on plants (bottom of the leaf). The number of stomata on the epidermal surface can tell you a lot about a plant. Usually, a high concentration of stomata indicates fast growth and wet climate. Lower concentrations of stomata indicate lower rates of photosynthesis and growth or adaptations for dry weather.

Purpose:

To view and compare the stomata from the leaves of several species of plant

Materials:

3 leaves (1 from 3 different species), compound light microscope, 3 microscope slides, clear nail polish, transparent tape

Procedure:

- 1. Obtain three leaves from different types of plants.
- 2. Paint a thick patch (at least one square centimeter) of clear nail polish on the underside of the leaf surface being studied.
- 3. Allow the nail polish to dry completely.
- 4. Tape a piece of clear cellophane tape to the dried nail polish patch.
- 5. Gently peel the nail polish patch from the leaf by pulling on a corner of the tape and "peeling" the fingernail polish off the leaf. This is the leaf impression you will examine.
- 6. Tape your peeled impression to a very clean microscope slide. Use scissors to trim away any excess tape. Label the slide with plant name.
- 7. Examine the leaf impression under a light microscope at 400X.
- 8. Search for areas where there are numerous stomata, and where there are no dirt, thumb prints, damaged areas, or large leaf veins. Draw the leaf surface with stomata.
- 9. Count all the stomata in one microscopic field. Record the number on your data table.
- 10.Repeat counts for at least three other distinct microscopic fields. Record all the counts. Determine an average number per microscopic field.
- 11. From the average number/400X microscopic field, calculate the stomata per mm² by multiplying by 8.
- 12. Follow procedures 2 11 with the other leaves.

Data:

	Leaf 1	Leaf 2	Leaf 3
Name of Leaf			
Drawing in 400x (with several stomata)			
Stomata in field 1			
Stomata in field 2			
Stomata in field 3			
Average Stomata in field			
Stomata/ mm2			

Conclusion:

- 1. Which leaf had the most stomata? Why do you think this was so?
- 2. Explain, in detail, how guard cells open and close stomata?
- 3. At what time of day would stomata be closed and why?
- 4. Why does the lower epidermis have more stomata than the upper epidermis of a leaf?
- 5. Define transpiration.
- 6. What two gases move in and out of the leaf stomata?
- 7. What does a larger number of leaf stomata indicate about the growing climate of that plant?

Ch 9 Lab report help

Purpose- what are stomata and how are they involved in photosynethesis

Background: what is photosynthesis? What is the formula for photosynthesis? What is respiration? What is the formula for respiration? Where does photosynthesis and respiration take place in each cell? What are guard cells? What is transpiration? What is their role in both photosynthesis and respiration? What two gases move in and out of the leaf stomata?

Data-Include the chart above and three microscope drawings under high power

Conclusion- were the number of stomata per unit area different in each type of plant? Were the size of the stomata the same for each plant? What does a larger number of leaf stomata indicate about the growing climate of that plant? How would a plant be affected if the stomata became permanently closed?