# Identification of the common laboratory glassware, pipettes and Equipment 

## (1) Identification of the common laboratory glassware:

a. Conical flasks and beakers.
b. Graduated cylinders [measuring cylinder ].

Volumetric flasks.
Burettes.
Pipettes.


## (2) Identification of the common laboratory pipettes:

$\square$ Sometimes spelled pipet.
$\square$ Commonly used to transport a measured volume of liquid.
$\square$ Pipettes come in several designs for various purposes with differing levels of accuracy.
$\square$ There are three types of pipettes are used in biochemical laboratory:
(a) Volumetric or transfer pipettes.
(b) Graduated or measuring pipettes (Mohr and Serological Pipettes).
(c) Micropipettes.

Types of pipettes


## Comparison between types of pipettes

| Volumetric pipettes | Graduated pipettes |
| :---: | :---: |
| Transfer (designed to deliver accurately <br> fixed volume of liquid) | Measuring |
| Not graduated | Graduated |
| More accurate | Less accurate |
| Non-blown out | Some are blown out |
| Consists of a cylindrical bulb joined at <br> both ends to narrowed glass tubing. | Don’t contain a cylindrical bulb |

## Smallest division of graduated pipette



## Reading the meniscus:



## Steps of the Use of the pipettes:

1- Press the pipette into the pump with a slight twisting motion.
2- The pipette is first washed with water ,then rinsed several times with a little of the solution.
3- The pipette then filled to just above the mark , the liquid is allowed to fall to the mark .
4- The solution is allowed to drain into the appropriate vessel with the jet of the pipette touching the wall of the vessel.

5- After the flow of the liquid has stopped, the jet is held against the wall for some times and then removed.

## Note:

-For serological pipette, some are of the blown out type; the last drop being blown out against the vessel wall.
-For volumetric pipette a certain amount of liquid will remain at the tip and this must not be "blown out".

## Accuracy:

1- Volumetric flasks and volumetric pipettes $\rightarrow$ most accurate.
2- Burets and graduated pipets.
3- Graduated cylinders.
4- Beakers and conical flasks. $\rightarrow$ least accuracy - used only when a rough estimation of
volume is required-

## (3) Identification of the common laboratory Equipment:

A. pH meter.

B. Spectrophotometer.


Electronic Balance.


## pH and solution acidity:

- PH define as: the negative logarithm of the hydrogen ion concentration.

$$
\mathrm{pH}=-\log \left[\mathrm{H}^{+}\right]
$$

$\square$ So the term pH introduced as a way of expressing hydrogen ion concentration (acidity or alkalinity of a solution).
$\square \mathrm{PH}$ range value $(0-14) \rightarrow$ the higher PH number, the lower the hydrogen ion concentration and vice versa [ inverse relationship] .


## pH cont':

$\square$ PH determines many important aspects of the structure and activity of biological macromolecules and thus of the behaviour of the cell and organisms.
$\square$ There are many ways in biochemical laboratory to measure PH value such as :

1. litmus paper.

2. Test strips.

3. PH meter $\Rightarrow$ The most accurate and reliable method


## (1) pH meter:

- A pH meter is an electronic device used for measuring the pH (acidity or alkalinity) of a liquid.
- Before use it needs to be calibrated.

$\square \mathrm{pH}$ meter contain glass electrode which is very sensitive and readily responds to changes in hydrogen ion concentration .



The glass electrode

- A nice video show you how to use the pH meter: https://www.youtube.com/watch?v=vwY-xWMam7o


## （2）Spectrophotometer：

－Spectrophotometer is instrument used to measure the intensity of light that is transmitted or absorbed by a sample at a given wavelength．
－Wavelength in this instrument divided into：


I．Invisible range－ultraviolet－（from 100 to 360 nm ）$\rightarrow$ 【Qumirt cuvette ære used】


II．Visible range（above $360 \mathrm{~nm}-700 \mathrm{~nm}$ ）$\rightarrow$ 【Glass or plastic cuvette ære used］
－Blank ：contain everything except the compound to be measure．


## Spectrophotometer Principle



A nice video show you how dose spectrophotometer work:
http://www.youtube.com/watch?v=pxC6F7bK8CU

## (2) Spectrophotometer cont':

- By using the spectrophotometer, we can quantitatively measure absorbance, and this information can be used to determine the concentration of the absorbing molecule [concentration of unknown sample].

More concentrated solution will absorb more light and transmits less:
$\rightarrow$ So, the more concentrated solution $\longrightarrow$ high absorbance value.
$\rightarrow$ Less concentrated solution $\longrightarrow$ less absorbance value.


## (3) Electronic Balance:

- Electronic Balance is a device used to find accurate measurements of weight.
- It provide the results digitally, making them an easy tool for use.
- The weight can be displayed by different unites.
$\square$ Before waiting any substance, you should (Zero) the balance.

$\rightarrow$ What does mean zeroing of the electronic balance?
$($ mass of paper + substance $)-($ mass of paper $)=($ mass of substance $)$

$\square$ A nice video show you how to use the electronic balance: $h$ https://www.youtube.com/watch?v=0UymyTJATLc


## Practical Part

## Objective:

$\square$ To be familiar with most common biochemistry lab tools and equipment.

## Method and Results:

1. Identificetion of the common leboratory glessware:

| Glassware <br> number | Type of glassware | Final volume (capacity) |
| :---: | :---: | :---: |
| 1 |  |  |
| 2 |  |  |
| 3 |  |  |
| 4 |  |  |
| 5 |  |  |
|  |  |  |

## Method and Results:

## 2. Comparing between glessware accuracy:

1-Place a beaker in the electronic balance, and read the weight.
2-Remove the beaker from the balance, and add 5 ml of water using a graduated pipette (Mohr).
3-Record the weight.
4-Repeat the procedure again by using measuring cylinder this time.
5-Record the weight.

| Type of glassware | Weight of beaker <br> $(\mathrm{g})$ | Weight of beaker + water <br> $(\mathrm{g})$ | Weight of water <br> $(\mathrm{g})$ |
| :--- | :---: | :---: | :---: |
| Graduated pipette (Mohr) |  |  |  |
| Measuring cylinder |  |  |  |

Which one is more accurate?

## Method and Results:

## 3. \dentification of the common laboratory pipettes:

1-Examine the three pipettes placed on your laboratory bench. 2-Record their types and the volume of their smallest division.

|  | Type of pipette | Smallest division |
| :--- | :--- | :--- |
| A |  |  |
| B |  |  |
| C |  |  |

## Method and Results:

## Ao Identffication of the common leboratory equipmentr

## A. $\mathbf{p H}$ meter:

1-Standardize the PH meter by placing the electrode in a solution of known pH
(PH $4,7,9) \rightarrow$ Calibration.
2-Wash the electrode with distilled water and dry by tissue then put it into sample solution A then wash it again and place it in solution $\mathrm{B} \rightarrow$ Read pH .

Note: After use the electrode, you should storage it in distilled water and never be allowed to dry out. If the electrode get dry it will required reactivation.

| Solution | pH value | Neutral, acidic or basic |
| :--- | :--- | :--- |
| Standard 4 |  |  |
| Standard 7 |  |  |
| Standard 9 |  |  |
| Sample A |  |  |
| Sample B |  |  |

## Method and Results:

## 4o Identification of the common leboratory equipmento

## B. Spectrophotometer:

1- Adjust the spectrophotometer to zero using water as blank solution in the cuvette.
2-Read the absorbance of standard solution and the solution of unknown concentration at 280 nm . 3-Read your result.

| Solution | Absorbance at 280 mm |
| :--- | :--- |
| BSA standard solution $(0.5 \mathrm{~g} / 100 \mathrm{ml})$ |  |
| Solution of Unknown concentration |  |

## Calculation:

$\mathrm{C}_{\text {standard }} \rightarrow \mathrm{A}_{\text {standard }}$ $\mathrm{C}_{\text {unknown }} \rightarrow \mathrm{A}_{\text {unknown }}$

$$
\mathrm{C}_{\text {unknown }}=\mathrm{C}_{\text {standard }} \mathrm{XA} \mathrm{~A}_{\text {unknown }}
$$

$\mathrm{A}_{\text {standard }}$

## WVhere:

$\mathbf{C}_{\text {standard }}=$ concentration of standard solution, $\mathrm{C}_{\text {unknown }}=$ concentration of unknown solution, $\mathbf{A}_{\text {standard }}=$ Absorbance of standard solution, $\mathbf{A}_{\text {unknown }}=$ Absorbance of unknown solution.

## Example:

| Solution | Absorbance at 280 nm | $\mathrm{C}_{\text {unknown }}=\mathrm{C}$ | $\mathrm{C}_{\text {standard }} \mathrm{X} \mathrm{A}_{\text {unknown }}$ |  |
| :---: | :---: | :---: | :---: | :---: |
| BSA standard solution ( $0.5 \mathrm{~g} / 100 \mathrm{ml}$ ) | 0.675 |  | $\mathrm{A}_{\text {standard }}$ |  |
| Solution of Unknown concentration | 1.2 | So: $\quad \mathrm{C}_{\text {unknown }}=$ | $=\underline{0.5 \mathrm{~g} / 100 \mathrm{ml} \times 1.2}$ | $=0.889 \mathrm{~g} / 100 \mathrm{ml}$ |
|  |  |  | 0.675 |  |

## Homework:

1- What is the smallest division for the following:


2- Measuring cylinder cannot be a substitute for the pipette or a burette, why?
3- What is the meaning of Calibration?
4- There are three different solution have pH values 3,7 and 10 :

- solution 1 is basic ( T or F )
- solution 2 is neutral ( T or F )
- solution 3 is acidic (T or F)

5- Why in the invisible range wavelength quartz cuvette is used?

