AL-Mustansiriyah University
College of Science
Chemistry Department

## Practical Physical Chemistry

Second stage
Second course

## Subject teacher

Assistant Professor/Athraa Salman Ahmed

## Second Course Experiments

1-Finding the molecular weight of an unknown solid by the method of decreasing the degree of freezing.

2- Findingthe molecular weight of an unknown solid by boiling point method.

3- Phase diagram of a binary group consisting of (solid solid).

4- Mutual solubility between (phenol- water).
5- Determine the solubility of sodium sulfate and determine the transition point.

6-Solubility as a function of temperature.


## Introduction

## The effect of the solute on some properties of the solvent

## The solvent has the most important properties :

1-Vapor pressure: It is a measure of a liquid's tendency to vaporize and it changes directly with temperature.

2-Boiling point: It is the temperature at which the vapor pressure of a liquid is equal to the pressure on it.

3- Freezing point: It is the temperature at which a liquid turns into a solid under atmospheric pressure.
-*It has been observed that any solution that contains a (non volatile solute) will exhibit the following bonding properties:

1- Decreased vapor pressure of the solvent.
2- High boiling point.
3- A decrease in the degree of freezing.
4- Osmosis and osmotic pressure of the solution.
Bonding properties: It is that property that depends only on the number of solute particles present in the solution and does not depend in any way on the chemical nature of these particles. * Note: The present solute must not occur in any union or dissociation. If there is a union of solute particles, the actual number of particles added will become small and as a result, these properties will decrease. Similarly, if the solute particles dissociated, as in the case of the electrolytes, the number of particles becomes greater than the original number and the value of the bonding properties becomes higher. In these solutions in which the particles of the solute occur to unite or dissociate, there must be some basic modifications to the laws producing solutions in which the solute particles do not unite or dissociate While discussing these properties it should be assumed that solutions They are very dilute and ideal, that is, the unions between solute particles are small To the point of neglect. The
study of these properties helped a lot in calculating the molecular weight of non-volatile solutes.

We will take the first associative property which is (no experiment only explanation)

## 1-The decrease in the vapor pressure of the solvent

When a non-volatile solute is added to the (liquid) solvent, the vapor pressure of the solvent decreases because the solute molecules impede the escape of the solvent molecules from the surface of the liquid. In the vapor pressure of the solvent, from this we conclude that the decrease in the vapor pressure of the solvent depends on the number of dissolved particles and not on the chemical nature of the solute, and this can be explain according to Raoult's law.

$$
\mathrm{P}=\mathrm{p}^{\mathrm{o}} \cdot \mathrm{x}_{1} \longrightarrow(1)
$$

X $1=$ moL fraction of the solvent
$\mathrm{P}^{\mathrm{o}}=$ vapor pressure of the pure solvent
$\mathrm{P}=$ vapor pressure of a solvent when a substance is dissolved in it (above the solution)

Since $\mathrm{X}_{1}$ Positive quantity less than one unit, then the vapor pressure of the solvent ( $P$ ) is less than ( $p o$ )

The decrease in vapor pressure $(\Delta \mathrm{P})$ is given by the following equation :-

$$
\Delta \mathrm{P}=\mathrm{P}^{\mathrm{o}}-\mathrm{P} \longrightarrow \text { (2) }
$$

Substituting (1) into (2), we get

$$
\begin{equation*}
\Delta \mathrm{P}=\mathrm{p}^{\mathrm{o}}-\left(\mathrm{p}^{\mathrm{o}} \cdot \mathrm{x}_{1}\right) \tag{3}
\end{equation*}
$$

$\Delta \mathrm{P}=\mathrm{p}^{\circ}\left(1-\mathrm{x}_{1}\right) \longrightarrow$
$\mathrm{X}_{1}+\mathrm{X}_{2}=1$
$\mathrm{X}_{2}=1-\mathrm{x}_{1}$
$\mathrm{X} 2=\mathrm{mol}$ fraction of the solute
Substituting (4) into (3), we get:

$$
\Delta \mathrm{P}=\mathrm{p}^{\circ} \mathrm{x}_{2}
$$

Equation (2) can be written as follows:

$$
\frac{\Delta \mathrm{P}}{\mathrm{p}^{\circ}}=\frac{\mathrm{p}^{\circ}-\mathrm{p}}{\mathrm{p}^{\circ}}=\mathrm{X}_{2}
$$

## That :

$p^{0}-p$
$P^{0} \quad$-:It is the amount of decrease in vapor pressure and this indicates that the decrease the relative vapor pressure is a correlative property of being dependent on $x_{2}$ For the solute and nothing else.

## The first experiment

## Measurement of the molecular weight of a solid by the decrease in freezing point

## 2-Decrease in freezing point

The reason is that the presence of solute particles between the solvent molecules impedes the convergence of the solvent molecules until it freezes, for example, at the normal freezing point of water. The solute particles work to keep the water molecules apart, so the water does not freeze, and to bring the water molecules closer together, the temperature must be lowered to less than the normal freezing point.

Raoult found that the freezing point varies directly with the molality of the solution

Amount of decrease in freezing point $\alpha$ molality of the solution
$\Delta t=K f \times$ The molality of the solution
$\Delta t=\operatorname{Kf} \times$ number of moles of solute $/$ weight of solvent $(\mathrm{kg})$

$$
\frac{\text { Solute (M.wt } / \mathrm{Wt})}{\text { Weight of solvent (kg) }} \quad x K_{f}=\Delta t
$$

Solute W1


Solvent $W_{2} \quad$ Solute M.Wt

Note: You must pay attention to the weight of the solvent converted by law directly to Kg

$$
\mathbf{M . W t}=\frac{K_{f} X \text { solute } W_{1} \times 1000}{\text { Solvent } W_{2} \mathbf{X} \Delta t}
$$

Where $\mathrm{K}_{\mathrm{f}}$ It is the constant of decrease in the degree of freezing (molality) and it is a constant specific to each solvent.

Kf : It is a constant specific to each solvent and is the amount of decrease that causes one mole of solute to dissolve in 1000 grams of the solvent.

The following table shows the values of (k) for some known solvents

| solvent | K |
| :---: | :---: |
| water | 1.86 |
| Acetic acid | 3.90 |
| Benzen | $4.9-5.10$ |
| Formic acid | 2.8 |
| Phenol | 7.3 |
| Camphor | 40.0 |

## Working method:

The simplified device consisting of two glass tubes, one inside the other is taken and placed inside a glass container containing ice and salt (ice bath). The solid substance and solvent are placed in the inner tube with a thermometer to measure the temperature. This device is called (Beckman device) and this method is called (Beckman method).

1- $(10 \mathrm{ml})$ Benzene is weighed and the weight is recorded. Then it is placed in the inner tube with the thermometer while stirring, and the temperature is measured every ( 30 seconds) while recording the time until the solidification of the Benzene (solvent).
(Super Freezing phenomenon must be exceeded) In the event that it occurs, the tube is removed from the ice bath immediately.

2- The inner tube containing the pure solvent is taken out of the ice bath and held by hand to dissolve the Benzene
$3-(0.5 \mathrm{~g})$ of the unknown solid is weighed and placed in the inner tube with benzene, and the inner tube is placed inside the
second tube in the ice bath with stirring, and the temperature is measured every ( 30 seconds) with time recording until the mixture freezes and the freezing point recorded.


## Beckman device

## Calculations:

1 - Calculate the molecular weight from the law

$$
\mathrm{M} . \mathrm{Wt}=\frac{\mathrm{K}_{\mathrm{f}} \mathrm{X} \text { solute } \mathrm{W}_{1} \times 1000}{\text { Solvent } \mathrm{W}_{2} \times \Delta \mathrm{t}}
$$

## $\mathrm{M} . \mathrm{Wt}=$ Molecular weight of the solute $(\mathrm{g} / \mathrm{mol})$

$\mathrm{W}_{1}=$ Solute weight (gm)
$\mathrm{W}_{2}=$ Solvent Weight (Kg)
$\mathrm{K}_{\mathrm{f}}=$ Freezing point decrease constant( $5.19 \mathrm{~K} . \mathrm{Kg} / \mathrm{mol}$ )
$\Delta t=$ The difference in temperature between (freezing of a pure solvent and a solvent with an impurity)

2- We draw the graph between temperature and time


Time

## Second experience

## Measurement of the molecular weight of a solid by the height method boiling point

## 3 - High boiling point

As we studied previously, when adding a solute in a pure solvent will lead to a decrease in the freezing point of the solvent and at the same time lead to a rise in the boiling point and the reason is as we previously learned that the solute reduces the vapor pressure of the solvent and this needs heat more than the heat needed by the pure solvent until Their vapor pressures are equal, and thus the boiling point of the solvent rises in the presence of the solute when the pressure on it is constant. It was found that: The increase in the boiling point of the solvent varies directly with the molar concentration of the solution.

Amount of increase in boiling point $\alpha$ molality of the solution
$\Delta t \quad=K_{b}$ The molality of the solution
$\Delta t \quad=K_{b} \times$ number of moles of solute / weight of solvent $(\mathrm{kg})$

> Solute (M.wt /Wt)

Weight of solvent (kg)

$$
\frac{1000}{\text { Solvent } W_{2}} \times \frac{\text { Solute } W_{1}}{\text { Solute M.Wt }} \times K_{b} \quad=\Delta t
$$

Note: You must pay attention to the weight of the solvent converted by law directly toKg

$$
\mathrm{M} . \mathrm{Wt}=\frac{\mathrm{K}_{\mathrm{b}} \mathrm{X} \text { solute } \mathrm{W}_{1} \times 1000}{\text { Solvent } \mathrm{W}_{2} \mathrm{X} \Delta \mathrm{t}}
$$

Where kb :- is the boiling point constant, and it is a solvent-specific constant, and it varies with the change of the solvent.

## The method of work:

1- Install the device for this experiment.
2-Weight ( 10 ml ) of benzene and is placed in the beaker with two holes (one hole is placed in which a thermometer is placed and the other hole is connected to a condenser) and the boiling point of pure benzene is measured after we fix the temperature.

# 3 - weigh ( 0.5 gm ) of the unknown substance (solid) is 

 placed on benzene and we measure its boiling point after we fix the temperature.

## Calculations:

## 1-We calculate the molecular weight from the law:

$$
M . W t=\frac{K_{b} \times \text { solute } W_{1} \times 1000}{\text { solventW }_{2} \times \Delta t}
$$

M.Wt $=$ Molecular weight of the solute $(\mathrm{g} / \mathrm{mol})$
$\mathrm{W}_{1}=$ Solute weight( gm )
$\mathrm{W}_{2}=$ Solvent weight $(\mathrm{Kg})$
$\mathrm{K} \mathrm{b}=$ Boiling point increase constant ( $2.5 \mathrm{~K} . \mathrm{Kg} / \mathrm{mol}$ )
$\Delta t=$ The temperature difference between (pure solvent and solvent with impurity)

## The third experience

## Phase diagram of a binary group consisting of (solid (solid

## The theoretical part:

The phase rule was developed by CBS: It expresses the conditions for the occurrence of a state of equilibrium in a heterogeneous system, regardless of the number of components or phases in that system.
$\mathrm{F}+\mathrm{P}=\mathrm{C}+2$
$\mathrm{C}=$ Represent the number of components
$\mathrm{P}=$ represents the number of phases
$\mathrm{F}=$ Represents degrees of freedom

Phase: It is the homogeneous part that is physically distinguished and separated from the rest of the other parts of the system by separating boundaries.

Example (snow, liquid water and water vapor) Here each form is a separate phase

Solid system (homogeneous) where the phase is single, regardless of the number of components it contains

## Liquid

 system

That mean, the two liquids are not mixed, then there are two separate phase, such as (benzene and water)

The gaseous system (homogeneous) means the gases are always mixed and form a homogeneous mixture that is always a single phase, regardless of the number of components Number of components: It is the smallest number of variable components by which the composition of each phase can be expressed either directly or indirectly or in the form of the chemical equation.

An example of a water system that is (ice, water, and water vapor) in equilibrium is considered a system with one component, which is water.

Degrees of Freedom: The number of variables such as temperature, pressure, or concentration that must be specified to define the system completely.

The phase base is affected by its equilibrium only by temperature, pressure, and concentration, and not by any other forces such as gravity, electricity and magnetism.

Let's take some examples:
(One component system)
1- water $\left(\mathrm{H}_{2} \mathrm{O}\right)$ system (ice and water and water vapor)
$\mathrm{F}+\mathrm{P}=\mathrm{C}+2$
$\mathrm{F}+3=1+2$
F=3-3=0
2-Acetone (liquid and vapor acetone
$\mathrm{F}+\mathrm{P}=\mathrm{C}+2$
$\mathrm{F}+2=1+2$
$\mathrm{F}=3-2=1$
( G -S ) (G- L ) (LL)(SL)(SS) (Two-component system, twocomponent system)
(L- L) (SS) will be studied
In such systems experiments are conducted under atmospheric pressure, thus the pressure remains constant, and this will reduce the degree of freedom of the system by one degree, and the phase base becomes called (reduced phase rule).
$\mathrm{F}+\mathrm{P}=\mathrm{C}+1$
$\mathrm{F}=\mathrm{C}-\mathrm{P}+1$

In our experiment, the container is a two-component system $(\mathrm{P}=1)$ and single phase $(\mathrm{C}=2)$

The number of degrees of freedom according to the (reduced phase rule), then:
$\mathrm{F}+\mathrm{P}=\mathrm{C}+1$
$\mathrm{F}=2-1+1$
$\mathrm{F}=2$
That is, it needs two variables to define the system, which are temperature and concentration.

As for the (normal phase rule), the following:
$\mathrm{F}+\mathrm{P}=\mathrm{C}+2$
$\mathrm{F}=2-1+2$
$\mathrm{F}=3$
That is, it needs three variables to define the system: concentration, pressure and temperature. In such a case, it is difficult to draw in three dimensions on the graph, so one of the variables, which is pressure, is fixed, and the change in temperature and concentration is done for the purpose of defining the system.

Let's take a general case of concentrated systems, which are systems that do not have a two-component gas phase They
contain two components, B and A, which mix well in the liquid state, and their solutions produce only pure A or pure B as solid phases. As shown in the chart below


The figure shows the phase diagram

1- The components A and B mix completely in the liquid state and their solutions produce only pure $A$ and pure $B$ as solid phases.

2-(A , B ) The points represent the two melting points (degrees of freezing) for pure $A$ and pure $B$ respectively

3- When a quantity of $B$ is added to $A$, the degree of freezing of A decreases along the AC curve, and in the same way, when a quantity of A is added to B , the degree of freezing of B decreases along the curve BC .

4- The AC curve is a curve that represents the freezing point of component A and represents the composition of solutions saturated with solid A at temperatures between A and C , so we notice that the two phases are in equilibrium along AC (solid A and solution B are in A ).

If we apply the reduced phase rule, then:
$\mathrm{F}=\mathrm{C}-\mathrm{P}+1$
$\mathrm{F}=2-2+1$
$\mathrm{F}=1$
5- Similarly, the curve BC is the freezing point curve of component B and represents the composition of solutions saturated with solid $B$ at temperatures between $B$ and $C$, so we notice that the two phases are in equilibrium along BC (solid B and solution A are in B ).

If we apply the reduced phase rule, then:
$\mathrm{F}=\mathrm{C}-\mathrm{P}+1$
$\mathrm{F}=2-2+1$
$\mathrm{F}=1$
6-The two curves intersect at point $C$ where both solids A and B are in equilibrium with the liquid phase and since there are three phases in equilibrium at this point, the reduced phase rule is as follows at point c
$\mathrm{F}=\mathrm{C}-\mathrm{P}+1$
$\mathrm{F}=2-3+1$
$\mathrm{F}=0$
That is, at this point the temperature and composition of the solution are constant as long as the three phases are in equilibrium. If there is a change in one of these two variables, at least one of the phases will disappear

Point $C$ is known as the eutectic point. It is the temperature at which the solids are in equilibrium with the liquid phase.

7- The area above the lines AC and BC is related as the field for the presence of the unsaturated solution or the molten liquid (the two components exist as a homogeneous liquid solution) and thus there is only one phase and the system is bivariate and according to the reduced phase rule $\mathrm{F}=\mathrm{C}-\mathrm{P}+1$

$$
\mathrm{F}=2-1+1
$$

$\mathrm{F}=2$
That is, in order to define any point in this region, the temperature and composition must be specified.

## Working method :

1- (11) test tubes are taken and the percentage set in the table for each of two substances is placed in them
(B-Naphthol - Acetamide ). In our experiment (S-S ), we will take two solid materials, which are (Bethanphthol) B and (Astamide) A. The sum of the total weight of the two materials $=6 \mathrm{gm}$ in each tube

For example, if we take $\% \mathrm{~A}=90$ What is the percentage of B ? total sum of the two substances $=6 \mathrm{gm}$
(A)

$$
\frac{90}{100}=\frac{\text { Part }}{\text { Whole }} \times 100
$$



Weight (A) gram =5.4 Part A
$B=6-5.4=0.6 \operatorname{gram}(B)$ Weight

As for the percentage of substanceB

$$
\begin{gathered}
\%=\frac{0.6}{6} \times 100 \\
=10 \%
\end{gathered}
$$

Thus, the rest of the tubes are calculated in the same way

| A\% | weight ( A ) <br> acetamide | B weight <br> B-Naphthol | B \% | tube <br> number |
| :--- | :--- | :--- | :--- | :--- |
| $100 \%$ | 6 gm | 0 | $0 \%$ | 1 |
| $90 \%$ | 5.4 gm | 0.6 gm | $10 \%$ | 2 |
| $80 \%$ | 4.8 gm | 1.2 gm | $20 \%$ | 3 |
| $70 \%$ | 4.2 gm | 1.8 gm | $30 \%$ | 4 |
| $60 \%$ | 3.6 g | 2.4 gm | $40 \%$ | 5 |
| $50 \%$ | 3 gm | 3 gm | $50 \%$ | 6 |
| $40 \%$ | 2.4 gm | 3.6 g | $60 \%$ | 7 |
| $30 \%$ | 1.8 gm | 4.2 gm | $70 \%$ | 8 |
| $20 \%$ | 1.2 gm | 4.8 gm | $80 \%$ | 9 |
| $10 \%$ | 0.6 gm | 5.4 gm | $90 \%$ | 10 |
| $0 \%$ | 0 | 6 gm | $100 \%$ | 11 |

Each tube is placed in a hot water bath (except for tubes (1) and (11) and the beginning of melting and the end of melting are recorded and takes an average of two degrees. We take the average of the two readings.

As for the tube (1 and 11), it is placed on the (Hot Plate) directly because it contains the pure substance without adding the second substance to it, and the degrees are recorded in the same way mentioned above.

## Calculations:

We draw the graphical relationship between the average of the temperature (Tem) and the percentage of one of the two components and calculate the eutectic point from the graph.


The percentage of one of the two components

## Fourth experiment

## Finding the mutual solubility of phenol and water

## The theoretical part

## There are two types of liquids:

A-Fully mixed liquids: They are liquids that mix and dissolve with each other and in all proportions to form a homogeneous solution in one phase, such as (alcohol and water)

B-Partially mixed liquids: They are liquids that do not dissolve into each other and do not mix completely. That is they dissolve partially soluble forming a heterogeneous solution and two phases. When the mixture is shaken well and the bottle that contains it is left on the table. We find that after a period the two layers separate. An example of this (phenol and water) (ether and water). And these second types there are three types:

1- Liquid whose solubility in each other increase with increasing temperature (High Critical Temperature)


Such as (phenol and water). Which is the subject of our experiment, as the two liquids become completely miscible. That is, it will be one layer and one phase. Critical temperature:
Is the temperature at which if the mixture is exceeded each of the two solutions becomes dissolved in the other and in all proportions forming a homogeneous solution one phase.


The point (A) represents the composition of the (water-rich) layer.
The point (B) represents the composition of the (phenol-rich) layer.
All mixtures between (A) and (B) product two layers with the composition of (A) and (B).

When you add a quantity of phenol to a large amount of water, phenol is completely dissolved and be a solution (phenol in the water) then we have a fully mixed solution when using high percentage for one liquid.

$$
F=C-P+1=2, P=1
$$

When adding an increase of phenol that (exceeds the saturation limit) to the water, two layers will be produced, (layer of a saturated solution of phenol in water) and (a layer of a saturated solution of water in phenol). This case is inside the para bola.

$$
\mathrm{F}=\mathrm{C}-\mathrm{P}+1=1, \mathrm{P}=\mathbf{2}
$$

In a similar way, the water dissolved completely with a high percentage of phenol. So that the solution (water in phenol) then we have a fully mixed solution (when using a high percentage of one of the two liquids).
$\mathbf{F}=\mathbf{C}-\mathbf{P}+\mathbf{1}=\mathbf{2 , P}=\mathbf{1}$
When more water is added (exceeding the saturation limit) to phenol, two layers are formed.

## $\mathbf{F}=\mathbf{C}-\mathbf{P}+\mathbf{1 = 1 , P = 2}$

Upon reaching point (C) which is the critical temperature the two liquids become completely mixed, and one solution is formed in one phase.

$$
F=C-P+1=2, P=1
$$

The region above (C) will form a single homogeneous liquid phase.
There are many examples (water and aniline) (aniline and hexane) (benzene and sulfur) (methyl alcohol and cyclo hexane) (methyl alcohol and carbon disulfate).
2- Liquids whose solubility in each other increase with decreasing (Temperature)
(Lower Critical Temperature) such as (water and try ethyl amine).


3- Liquids that have (two critical temperatures) upper and lower such as (water and nicotine)


## The method of work:

1- They are placed in the tubes and as shown in the table below, we will take two substances, phenol and water, bearing in mind that the total volume of the two substances is 10 ml . If the percentage of phenol is $70 \%$, how do we calculate the percentage of water? And the volume of phenol and the volume of water?

$$
\begin{aligned}
& \frac{70}{100}=\frac{\text { Part }}{\text { Whol }} \times 100 \\
& \frac{70}{100}=\frac{\text { Part }}{10} \times 100
\end{aligned}
$$

$($ Volume of phenol $) 7 \mathrm{ml}=$ Part
$10-7=3 \mathrm{ml}$ ( volume of water)
3
\%Water $=$ X100 10
$\%$ water $=30 \%$

| water <br> volume | water \% | phenol <br> volume | phenol\% | tube <br> number |
| :--- | :--- | :--- | :--- | :--- |
| 1 | $\% 10$ | 9 | $\% 90$ | 1 |
| 2 | $\% 20$ | 8 | $\% 80$ | 2 |
| 3 | $\% 30$ | 7 | $\% 70$ | 3 |
| 4 | $\% 40$ | 6 | $\% 60$ | 4 |
| 5 | $\% 50$ | 5 | $\% 50$ | 5 |
| 6 | $\% 60$ | 4 | $\% 40$ | 6 |
| 7 | $\% 70$ | 3 | $\% 30$ | 7 |
| 8 | $\% 80$ | 2 | $\% 20$ | 8 |
| 9 | $\% 90$ | 1 | $\% 10$ | 9 |

2- Each tube is placed in a hot water bath, and the temperature is calculated when the two liquids become completely mixed, and then the tube is cooled and the temperature is recorded when the two liquids become incompletely mixed (i.e. the solution becomes turbid). Then we take the average of two degrees for each tube.


## Calculations:

We draw the graphic relationship between the average temperatures and the percentage of one of the two components for each tube and its color. The highest critical temperature for melting is determined from the graph


The percentage of one of the two components

## Fifth experience

# Determine the solubility of sodium sulfate in water and find the transition point 

## The theoretical part

Solubility : is the largest amount of a substance that can be dissolved in a fixed volume of a specific solvent to obtain a saturated solution at a known temperature.

## Typs of solution

1- Saturated solution : it is one in which equilibrium is established between dissolved and un-dissolved solute at a definite temperature.
2- Un saturated solution : it is a solution that contains the dissolved solute in a concentration below that necessary for complete saturation at definite temperature.
3- Super saturated solution : it is a solution contains more of the dissolved solute than it would normally contain in a saturated state at a definite temperature.

Some salts (e.g. Sodium thiosulfate) can be dissolved in a large amounts at an elevated temperature and upon cooling fail to crystallize from the solution. These supersaturated if disturbed in anyway (as cooling) the excess solute will precipitate out of solution and a saturated solution will result.

## Solubility expressions methods

1- Molarity
2- Normality
3- Molality
4- Mole fraction
5- W/W \%
6- V/V \%
7- W/V \%

## Factors effecting solubility of a solid in liquid

1- Nature of solute
2- Nature of solvent
3- Pressure
4- Temperature

## Temperature effect

Endothermic dissolution : Most solids dissolve with absorptions of heat ( $\Delta \mathrm{H}$ is positive) and the dissolution process is an endothermic process.

Exothermic dissolution : In case the less commonly accruing system that exhibit exothermic dissolution the solubility of exothermic dissolution is decrease by increasing the
temperature.


## Solubility curve

Plot of the solubility curves with temperature, which is referred to as solubility curves are often used describe the effect of temperature on a given system. Most of the curves are continuous. Abrupt changes in slop may be observed with some. Systems if a change in the nature of the dissolving solid accrues at a specific transition temperature. Example for abrupt change (Sodium sulfate) exists as the Decca hydrate ( $\mathrm{Na}_{2} \mathrm{SO}_{\mathbf{4}} \cdot \mathbf{1 0 \mathrm { H } _ { 2 } \mathrm { O } )}$ and its dissolution in water is an endothermic process. Its solubility therefore increases with rise in (temp.) above this (temp.), the solid is converted to anhydrous from ( $\mathrm{Na}_{2} \mathrm{SO}_{4}$ ) and the dissolution of this compound is an exothermic process.

The equation used to calculate solubility is the (Vant-Hoff) equation, which shows the change of solubility with (temp.) at constant pressure.

Ә LinS $/$ Ət $=\Delta H /$ RT $^{2}$
$\log S=-\Delta H / 2.303 R T+C$ (integral formation)
C= Proportionality constant
$\mathrm{R}=$ General constant for gases (J/mol.K)
$\Delta H=$ Heat of solubility ( $\mathbf{J} / \mathrm{mole}$ )
S= Solubility (W/W \%)

When the solid (the solute) is in contact with the liquid (the solvent) in which it is soluble, it diffuses in the solvent until the state of saturation occurs.

The temperature at which a substance changes from one crystalline form to another crystal form without any change in its composition is called a transition point.
$\mathrm{Na}_{2} \mathrm{SO}_{4} \cdot 10 \mathrm{H}_{2} \mathrm{O} \rightleftarrows \mathrm{Na}_{2} \mathrm{SO}_{4}+10 \mathrm{H}_{2} \mathrm{O}$

The solubility ( S ), it is calculated using the mathematical relationship:

$$
\text { Solubility }(S)=100 \times \frac{\text { Weight of solute }}{\text { Weight of solution }}
$$

## The method of work:

1- We take six clean, dry sacks and number them according to the temperatures $(55,50,45,40,35,30)$ and weigh them empty and record the weight (six weights).

1- We take ( 40 ml ) of distilled water and put it in a beaker and heat it to a temperature of $\left(60{ }^{\circ} \mathrm{C}\right)$, and then add sodium sulfate to the hot water until a saturated solution is prepared from it.

2- The solution is cooled to a temperature of ( $55^{\circ} \mathrm{C}$ ) and then withdrawn by a pipette ( 5 ml ) of the saturated solution and placed in the first beaker $\left(55^{\circ} \mathrm{C}\right)$ and the same step is repeated in the remaining degrees.
4- Weigh the six beakers with the solution to get (six weights), then leave them in the oven until the next day :

Weight of the beaker with the solution - the weight of the beaker when it is empty = weight of the solution

5 -On the second day, the beakers are weighed with the solute (the precipitate) and the weight (six weights) is recorded.

Weight of beaker with solute - weight of beaker when empty $=$ weight of solute

## Calculations:

A-We calculate the solubility ( S ) of the six beakers

$$
\text { Solubility }(S)=100 \times \frac{\text { Weight of solute }}{\text { Weight of solution }}
$$

| tc ${ }^{\circ}$ | T (K) | 1/T | weight of solution | solute weight | S | LogS |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| $\begin{aligned} & 55 \\ & 50 \\ & 45 \\ & 40 \\ & 35 \\ & 30 \end{aligned}$ | ( 55+273) |  |  |  |  |  |

B - We draw the graph between $\log S$ and $1 / T$ to calculate the heat of dissolution (solution):

It is the heat resulting from dissolving sodium sulfate in 40 ml of distilled water.

After that, we calculate the transition point, that is, the temperature at which a change in the crystalline form of sodium sulfate occurs.

$$
\begin{aligned}
& \log \mathrm{S} \\
& \Delta H=\text { Slop } \times 2.303 \times R^{\frac{1}{T}}
\end{aligned}
$$

The transition point will be $1 / \mathrm{T}$, we take its reciprocal, we get (K) T, subtract from it (-237), we get ( $\mathrm{t}^{\mathrm{o}} \mathrm{c}$ ), which is the transition point.

R : value can be used
$\mathrm{R}=0.082$ Lit. atm. $\mathrm{mol}^{-1} \cdot \mathrm{~K}^{-1}$
$\mathrm{R}=8.314 \mathrm{~J} . \mathrm{mol}^{-1} . \mathrm{K}^{-1}$
$\mathrm{R}=1.987 \mathrm{Cal} . \mathrm{atm} . \mathrm{mol}^{-1} \cdot \mathrm{~K}^{-1}$

## Sixth experiment

## Solubility as a function of temperature

## The theoretical part:

Similar to the theoretical part of the fifth experiment

## The method of work



1- Prepare three saturated solutions of oxalic acid at different temperatures (ice) (water with ice) (water)

2- Leave the three solutions for 5 minutes to take the temperature of the bath in which they were placed.

3- Take three clean, dry conical flasks and weigh them while they are empty

4- (10) of the clear solution is withdrawn from each solution by the pipette and placed in a clean, dry conical flask, weighed and recorded (the weight of the flask with the solution).

5- The solution is diluted with (10) of distilled water and two drops of phenophthalein are added to it and then it is titrated with $\mathrm{NaOH}(1 \mathrm{~N})$ and the volume of the three solutions is recorded.

## Calculations:

the method of work according to steps 3 and 2 From
The weight of the beaker when empty = known
The weight of the beaker with the solution ( 10 ml )= known

Three times for three solutions

1- Calculate the standard of oxalic acid according to step No. (5) for each solution according to the law


$$
\mathrm{V}=10 \mathrm{ml}=\mathrm{acid} \text { size }
$$

eq.wt $=\frac{90}{2}=\mathrm{H}_{2} \mathrm{C}_{2} \mathrm{O}_{4}$ of oxalic acid
3- We calculate the number of grams of solution for each solution (ie (the weight of the solution)
$\left[\begin{array}{l}\text { Weight of solution }=\text { Weight of beaker with solution }- \text { Weight of } \\ \text { beaker empty }\end{array}\right]$
Weight of solution $=?$

Three times for three solutions

3- Calculate the number of grams of solvent (water) for each solution (the weight of the solvent)
\(\left[\begin{array}{l}The number of grams of solvent=weight of the solution - weight of <br>
the acid <br>

Grams of solvent=?\end{array}\right]\)| Three |
| :--- |
| times |
| for |
| three |
| solution |

Grams of solvent $=$ ?

5-We calculate the number of grams of acid (weight) in (1000 gm ) of solvent for each solution (molality of acid).

Three times for
three solutions (calculate the molality)
$\left[\begin{array}{cc}\text { Weight of acid solute } & \text { Weight of solvent water } \\ W_{1} & W_{1} \\ m & 1000\end{array}\right]$

6 - work schedule

| m | Logm | T | $1 / \mathrm{T}$ |
| :--- | :--- | :--- | :--- |
|  |  |  |  |

7-We draw the graphic relationship between Logm and 1/T and then calculate the heat of the solution $\Delta \mathrm{H}$

$\mathrm{R}=8.314 \mathrm{~J}_{\mathrm{J}} \mathrm{mol}^{-1} \cdot \mathrm{~K}^{-1}$
$(\Delta \mathrm{H}=\mathrm{J} / \mathrm{mol}$

