My name is Subhashish Chattopadhyay. I have been teaching for IIT-JEE, Various International Exams (such as IMO [ International Mathematics Olympiad ], IPhO [ International Physics Olympiad ], IChO [ International Chemistry Olympiad ]), IGCSE ( IB ), CBSE, I.Sc, Indian State Board exams such as WB-Board, Karnataka PU-II etc since 1989. As I write this book in 2016, it is my 25 th year of teaching. I was a Visiting Professor to BARC Mankhurd, Chembur, Mumbai, Homi Bhabha Centre for Science Education (HBCSE) Physics Olympics camp BARC Campus.

I am Life Member of ...

- IAPT ( Indian Association of Physics Teachers )
The selection for National Camp (for Official Science Olympiads - Physics, Chemistry, Biology, Astronomy) happens in the following steps....

1) NSEP (National Standard Exam in Physics) and NSEC (National Standard Exam in Chemistry) held around 24th November. Approx 35,000 students appear for these exams every year. The exam fees is Rs 100 each. Since 1998 the IIT JEE toppers have been topping these exams and they get to know their rank/performance ahead of others.

2) INPhO (Indian National Physics Olympiad) and INChO (Indian National Chemistry Olympiad). Around 300 students in each subject are allowed to take these exams. Students coming from outside cities are paid fare from the Govt of India.

3) The Top 35 students of each subject are invited at HBCSE (Homi Bhabha Center for Science Education) Mankhurd, near Chembur, BARC, Mumbai. After a 2-3 weeks camp the top 5 are selected to represent India. The flight tickets and many other expenses are taken care by Govt of India.

Since last 50 years there has been no dearth of “Good Books”. Those who are interested in studies have been always doing well. This e-Book does not intend to replace any standard text book. These topics are very old and already standardized.
There are 3 kinds of Text Books

- The thin Books - Good students who want more details are not happy with these. Average students who need more examples are not happy with these. Most students who want to “Cram” quickly and pass somehow find the thin books “good” as they have to read less !!

- The Thick Books - Most students do not like these, as they want to read as less as possible. Average students are “busy” with many other things and have no time to read all these.

- The Average sized Books - Good students do not get all details in any one book. Most bad students do not want to read books of “this much thickness” also !!

We know there can be no shoe that’s fits in all.

Printed books are not e-Books! Can’t be downloaded and kept in hard-disc for reading “later” 

So if you read this book later, you will get all kinds of examples in a single place. This becomes a very good “Reference Material”. I sincerely wish that all find this “very useful”.

Students who do not practice lots of problems, do not do well. The rules of “doing well” had never changed …. Will never change !

In this e-Book I am trying to solve this problem. Those students who practice can learn.

No one can help those who are not studying, or practicing.

Learn more at http://skmclasses.weebly.com/iit-jee-home-tuitions-bangalore.html

Twitter - https://twitter.com/ZookeeperPhy

Facebook - https://www.facebook.com/IIT.JEE.by.Prof.Subhashish/

Blog - http://skmclasses.kinja.com
A very polite request:

I wish these e-Books are read only by Boys and Men. Girls and Women, better read something else; learn from somewhere else.
Preface

We all know that in the species “Homo Sapiens “, males are bigger than females. The reasons are explained in standard 10, or 11 ( high school ) Biology texts. *This shapes or size, influences all of our culture*. Before we recall / understand the reasons once again, let us see some random examples of the influence.

Random - 1

If there is a Road rage, then who all fight ? ( generally ? ). Imagine two cars driven by adult drivers. Each car has a woman of similar age as that of the Man. The cars “ touch “ or “ some issue happens”. Who all comes out and fights ? Who all are most probable to drive the cars?

( Men are eager to fight, eager to rule, eager for war. Men want to drive. Men want to win )

Random - 2

Heavy metal music artists are all Men. Metallica, Black Sabbath, Motley Crue, Megadeth, Motorhead, AC/DC, Deep Purple, Slayer, Guns & Roses, Led Zeppelin, Aerosmith ..... the list can be in thousands. All these are grown-up Boys, known as Men.

( Men strive for perfection. Men are eager to excel. Men work hard. Men want to win. )

Random - 3

Apart from Marie Curie, only one more woman got Nobel Prize in Physics. ( Maria Goeppert Mayer - 1963 ). So, ... almost all are men.

Random - 4

The best Tabla Players are all Men.


Random - 5

History is all about, which Kings ruled. Kings, their men, and Soldiers went for wars. History is all about wars, fights, and killings by men.
Boys start fighting from school days. Girls do not fight like this

( Men are eager to fight, eager to rule, eager for war. Men want to drive. Men want to win. )

Random - 6

The highest award in Mathematics, the “Fields Medal” is around since decades. Till date only one woman could get that. (Maryam Mirzakhani - 2014). So, ... almost all are men.

Actor is a gender neutral word. Could the movie like “Top Gun” be made with Female actors? The best pilots, astronauts, Fighters are all Men.

In my childhood had seen a movie named “The Tower in Inferno”. In the movie when the tall tower is in fire, women were being saved first, as only one lift was working....
Many decades later another movie is made. A box office hit. “ The Titanic “. In this also .... As the ship is sinking women are being saved. **Men are disposable.** Men may get their turn later...

Movies are not training programs. Movies do not teach people what to do, or not to do. Movies only reflect the prevalent culture. **Men are disposable, is the culture in the society.** Knowingly, unknowingly, the culture is depicted in Movies, Theaters, Stories, Poems, Rituals, etc. I or you can’t write a story, or make a movie in which after a minor car accident the Male passengers keep seating in the back seat, while the both the women drivers come out of the car and start fighting very bitterly on the road. There has been no story in this world, or no movie made, where after an accident or calamity, Men are being helped for safety first, and women are told to wait.

Random - 9

Artists generally follow the prevalent culture of the Society. In paintings, sculptures, stories, poems, movies, cartoon, Caricatures, knowingly / unknowingly, “ the prevalent Reality “ is depicted. The opposite will not go well with people. If deliberately “ the opposite “ is shown then it may only become a special art, considered as a special mockery.
Men go to “girl / woman’s house” to marry / win, and bring her to his home. That is a sort of winning her. When a boy gets a “Girl-Friend “, generally he and his friends consider that as an achievement. The boy who “ got / won “ a girl-friend feels proud. His male friends feel, jealous, competitive and envious. Millions of stories have been written on these themes. Lakhs of movies show this. Boys / Men go for “ bike race “, or say “ Car Race “, where the winner “ gets “ the most beautiful girl of the college.

( Men want to excel. Men are eager to fight, eager to rule, eager for war. Men want to drive. Men want to win. )

Prithviraj Chauhan ‘ went ` to “ pickup “ or “ abduct “ or “ win “ or “ bring “ his love. There was a Hindi movie ( hit ) song ... “ Pasand ho jaye, to ghar se utha laye “. It is not other way round. Girls do not go to Boy’s house or man’s house to marry. Nor the girls go in a gang to “ pick-up “ the boy / man and bring him to their home / place / den.

Random - 11

Rich people; often are very hard working. Successful business men, establish their business ( empire ), amass lot of wealth, with lot of difficulty. Lots of sacrifice, lots of hard work, gets into this. Rich people’s wives had no contribution in this wealth creation. Women are smart, and successful upto the extent to choose the right/rich man to marry. So generally what happens in case of Divorces ? Search the net on “ most costly divorces “ and you will know. The women; ( who had no contribution at all, in setting up the business / empire ), often gets in Billions, or several Millions in divorce settlements.


See http://skmclasses.kinja.com/save-the-male-1761788732
It was Boys and Men, who brought the girls / women home. The Laws are biased, completely favoring women. The men are paying for their own mistakes.

See https://zookeepersblog.wordpress.com/biased-laws/

( Man brings the Woman home. When she leaves, takes away her share of big fortune! )

Random - 12

A standardized test of Intelligence will never be possible. It never happened before, nor ever will happen in future; where the IQ test results will be acceptable by all. In the net there are thousands of charts which show that the intelligence scores of girls / women are lesser. Debates of Trillion words, does not improve performance of Girls.

I am not wasting a single second debating or discussing with anyone, on this. I am simply accepting ALL the results. IQ is only one of the variables which is required for success in life. Thousands of books have been written on “Networking Skills “, EQ ( Emotional Quotient ), Drive, Dedication, Focus, “ Tenacity towards the end goal “ ... etc. In each criteria, and in all together, women ( in general ) do far worse than men. Bangalore is known as “ ..... capital of India “. [ Fill in the blanks ]. The blanks are generally filled as “ Software Capital “, “ IT Capital “, “ Startup Capital “, etc. I am member in several startup eco-systems / groups. I have attended hundreds of meetings, regarding “ technology startups “, or “ idea startups “. These meetings have very few women. Starting up new companies are all “ Men’s Game “ / “ Men’s business “. Only in Divorce settlements women will take their goodies, due to Biased laws. There is no dedication, towards wealth creation, by women.
Many men, as fathers, very unfortunately treat their daughters as “Princess”. Every “non-performing” woman/wife was “princess daughter” of some loving father. Pampering the girls, in name of “equal opportunity”, or “women empowerment”, have led to nothing.

Let us recall the reasons, that we already learned in standard 10-11, Biology text books. In humans, women have a long gestation period, and also spends many years (almost a decade) to grow, nourish, and stabilize the child. (Million years of habit) Due to survival instinct Males want to inseminate. Boys and Men fight for the “facility (of womb + care)” the girl/woman may provide. Bigger size for males, has a winning advantage. Whoever wins, gets the “woman/facility”. The male who is of “Bigger Size”, has an advantage to win. Leading to Natural selection over millions of years. In general “Bigger Males”; the “fighting instinct” in men; have led to wars, and solving tough problems (Mathematics, Physics, Technology, startups of new businesses, Wealth creation, Unreasonable attempts to make things [such as planes], Hard work...)

So let us see the IIT-JEE results of girls. Statistics of several years show that there are around 17, (or less than 20) girls in top 1000 ranks, at all India level. Some people will yet not understand the performance, till it is said that...year after year we have around 980 boys in top 1000 ranks. Generally we see only 4 to 5 girls in top 500. In last 50 years not once any girl topped in IIT-JEE advanced. Forget about Single digit ranks, double digit ranks by girls have been extremely rare. It is all about “good boys “, “hard working “, “focused “, “Bel-esprit “ boys.

In 2015, Only 2.6% of total candidates who qualified are girls (upto around 12,000 rank). while 20% of the Boys, amongst all candidates qualified. The Total number of students who appeared for the exam were around 1.4 million for IIT-JEE main. Subsequently 1.2 lakh (around 120 thousands) appeared for IIT-JEE advanced.
IIT-JEE results and analysis, of many years is given at [https://zookeepersblog.wordpress.com/iit-jee-iseet-main-and-advanced-results/](https://zookeepersblog.wordpress.com/iit-jee-iseet-main-and-advanced-results/)

In Bangalore it is rare to see a girl with rank better than 1000 in IIT-JEE advanced. We hardly see 6-7 boys with rank better than 1000. Hardly 2-3 boys get a rank better than 500.

See [http://skmclasses.weebly.com/everybody-knows-so-you-should-also-know.html](http://skmclasses.weebly.com/everybody-knows-so-you-should-also-know.html)

Thousands of people are exposing the heinous crimes that Motherly Women are doing, or Female Teachers are committing. See [https://www.facebook.com/WomenCriminals/](https://www.facebook.com/WomenCriminals/)

Some Random Examples must be known by all
End violence against women...

North Carolina Grandma Eats Her Daughter’s New Born Baby After Smoking Bath Salts
Henderson, North Carolina—A North Carolina grandmother of 4 and recovering drug addict, is now in custody after she allegedly ate her daughter’s newborn baby...

Attractive Girl Gang Lured Men Into Alleyways Where Female Body Builder Would Attack Them
A Mexican street gang made up entirely of women has been accused of luring their feminine wiles to lure men into alleyways and then beating them up and...

Women are raping boys and young men
Rape advocacy has been reclaimed and twisted into a political agenda controlled by radicalized activists. Tim Patten takes a macro lens and well-informed look into the manufactured rape culture and...

End violence against women...

28-Year-Old Texas Teacher Accused of Sending Nude Picture to 14-Year-Old Former Student
http://www.wtmy.com/.../youngstown-woman-convicted-of-rape...-

Youngstown woman convicted of raping a 13-year-old is back in jail
A Youngstown woman who went to prison for raping a 13-year-old fifteen years ago is in trouble with the law again.

Bronx Woman Convicted of Poisoning and Drowning Her Children
Lisette Barrengao researched methods on the Internet before she killed her son and daughter in 2012.

NYTIMES.COM | BY MARC SANTORA.
In several countries or rather in several regions of the world, family system has collapsed, due to bad nature and naughty acts of women. Particularly in Britain, and America, almost 50% people are alone, lonely, separated, divorced or failed marriages. In 2013, 48% children were born out of wedlock. It was projected that by 2016, more than 51% children will be born, to unmarried mothers. In these developed countries “paternity fraud” by women, are close to 20%. You can see several articles in the net, and in wikipedia etc. This means 1 out of 5 children are calling a wrong man as dad. The lonely, alone “mothers” are frustrated. They see the children as burden. Love in the Society in general is lost, long time ago. The types of “Mothers” and “Women” we have now

............
This is the type of women we have in this world. These kind of women were also someone's daughter.

Mother Stabs Her Baby 90 Times With Scissors After He Eats Her While Breastfeeding Him!

Eight-month-old Raeb Ritu was discovered by his uncle in a pool of blood needed 100 stitches after the incident; he is now recovering in hospital. Reports say hit...
HURT FEMINISM BY DOING NOTHING

Don’t help women
Don’t fix things for women
Don’t support women’s issues
Don’t come to women’s defense
Don’t speak for women
Don’t value women’s feelings
Don’t portray women as victims
Don’t protect women

Without white knights feminism would end today

*Don’t even navalt (“Not All Women Are Like That”) 1 For example from criticism or insults

How Society prioritize Men

High Priority
Rich women
Women
Rich Men
Girls
Boys
Animals
Prisoners
Men
Poor Men

Low Priority

They can get away with murder.
They get all the rights with no responsibility and shelter for homeless women.
They get bail outs and short prison sentence.
They get educational benefits but no violence against kids Act.
They have some support but don’t have any education that fits boys.
They have animal rights and PETA.
They get conjugal visits and 3 squares and a roof.
Paid slaves.
Nothing.

Who pays the most Taxes?
This is why MGTOW exist.

Professor Subhashish Chattopadhyay
We must first understand the most important graph or image of Chemistry

It is very strange, that so many Chemistry books exist in this world, which do not even have a single Periodic table in them. Almost all books do not have this graph either. This is the graph of electrode potential of various elements with respect to (standard) Hydrogen electrode.

All chemical reactions happen because of this electrode potential difference. When a Solid is added to a solution of another compound such as a Gold Spoon in CuSO₄ Solution, of a Copper container having AgNO₃ Solution, then the displacement reaction happens or not is decided by these potential values. Battery and Electrochemistry chapter uses this knowledge. Oxidation-Reduction reactions, titrations etc all need this understanding.

**ELECTROCHEMICAL CELLS**

An electrochemical cell consists of two electrodes (metallic conductors) in contact with an electrolyte (an ionic conductor). An electrode and its electrolyte comprise an Electrode Compartment.

Electrochemical Cells can be classified as:

(i) Electrolytic Cells - in which a non-spontaneous reaction is driven by an external source of current.

(ii) Galvanic Cells - which produce electricity as a result of a spontaneous cell reaction a galvanic cell, cathode is positive with respect to anode.
In an electrolytic cell, anode is made positive with respect to cathode.

**Galvanic Cell**

Spontaneous rxn. drawn into cell from cathode where reduction occurs and release them at anode where oxidation occurs.

Example:

\[ \text{Cu}^{2+} + \text{Zn} \rightarrow \text{Cu} + \text{Zn}^{2+} \]

**Electrolytic Cell**

Current supplied by external source drive nonspontaneous oxidation/reduction reaction.

Anode + and cathode −, opposite of galvanic cell.

Example:

\[ 2\text{Cl}^- + \text{Mg}^{2+} \rightarrow \text{Cl}_2 + \text{Mg}(\text{I}) \]
ELECTROLYSIS (Where a Positive current, of holes, is forced into the Anode. So electrons will come out of the Anode.)

The decomposition of electrolyte solution by passage of electric current, resulting into deposition of metals or liberation of gases at electrodes is known as electrolysis. Or in another words, Electrolysis is the chemical deposition of a compound into its constituent elements brought about by a flow of electric current. In the Electrolyte Cations (positively charged) move to Cathode as it is with excess electrons.

- Direct current is passed through the compound (the compound can be in molten or aqueous state).
- Electrical energy (From the direct current) is changed into chemical energy (the decomposition of the compound).
- One common example is the electrolysis of water, where water decomposes to hydrogen and oxygen.
ELECTROLYTIC CELL

This cell converts electrical energy into chemical energy. The entire assembly except that of the external battery is known as the electrolytic cell. So **Main components** of an electrolytic cell: Electrolytes, Electrodes and External Battery (or Source of DC Voltage).

ELECTRODES

The metal strip at which positive current enters is called anode; anode is positively charged in electrolytic cell. On the other hand, the electrode at which current leaves is called cathode. Cathodes are negatively charged. So The positive terminal of the battery is connected to the positive electrode (anode) while the negative terminal of the battery is connected to the negative electrode (cathode).

Spoon Feeding

Anode is positive. Here electrons are lost. So it is getting oxidized. Positive current enters.

Cathode is negative. Here electrons are gained. Positive Current leaves the cathode.

Electrolytes are usually made up of an ionic compound in solution or molten state OR aqueous solutions of acids or alkalis.

- Electrolytes conduct electricity with decomposition at the electrodes as it does so.
- Electrolytes can be classified into three categories: Non-electrolytes, weak electrolytes and strong electrolytes.
- Strong electrolytes = lots of ions to carry the charges from one electrode to the other. Examples of strong electrolytes: Strong acids or alkalis (sulphuric acid, aqueous sodium hydroxide), salt solutions (aqueous sodium chloride)
- Weak electrolytes = few ions to carry the charges from one electrode to the other. Examples of weak electrolytes: Weak acids or bases (Ethanoic acid, aqueous ammonia)
- Non-electrolytes = no ions available to carry the charges from one electrode to the other. Examples of non-electrolytes: Pure water, organic liquids or solvents

Electrodes are conductors through which electrons enter and leave the electrolyte.

- Anode is the POSITIVE (+) electrode from which electrons leave the electrolyte (or the electric current enters the electrolyte)
- Cathode is the NEGATIVE (-) electrode from which electrons enter the electrolyte (or the electric current leaves the electrolyte)
- Electrodes are typically made up of inert materials (which do not participate in any redox reactions), such as carbon, platinum, titanium, or stainless steel. (NOTE: There are some cases where the electrodes undergoes redox reactions during electrolysis.)

What happens during electrolysis?

- Negative ions (Anions) are attracted to the anode while the positive ions (cations) are attracted to the cathode.
- When the ions reaches their respective electrodes, they will be discharged. (They lose or gain electrons and form neutral atoms)
- Hence, at anode, anions are discharged through loss of electrons; while at the cathode, cations are discharged through gain of electrons.
ELECTROLYSIS OF MOLTEN SODIUM CHLORIDE

NaCl(molten) $\rightarrow$ Na$^+$ + Cl$^-$

Reactions at anode (oxidation) 2 Cl$^-$ $\rightarrow$ Cl$_2$ (gas) + 2e$^-$ Recall Loosing electron is being oxidized

Reactions at cathode (reduction) Na$^+$ + e$^-$ = Na

There are two types of electrodes used in the electrolytic cell, namely attackable and non-attackable.

The attackable electrodes participate in the electrode reaction. They are made up of reactive metals like Zn, Cu, Ag etc. In such electrodes, atom of the metal gets oxidised into the corresponding cation, which is passed into the solution. Thus, such anodes get dissolved and their mass decreases. On the other hand, non-attackable electrodes do not participate in the electrode reaction as they made up of unreactive elements like Pt, graphite etc. Such electrodes do not dissolve and their mass remain same.

FARADAY'S LAWS OF ELECTROLYSIS:

(i) First law of electrolysis:

Amount of substance deposited or liberated at an electrode is directly proportional to amount of charge passed (utilized) through the solution.

$W \propto Q$

$W = \text{weight liberated, } Q = \text{charge in coulomb}$

$W = ZQ$
Thus, weight deposited by 1 coulomb charge is called electrochemical equivalent.

Let 1 ampere current is passed till ‘t’ seconds.

Then, $Q = I \cdot t$

∴ $w = Z \cdot I \cdot t$  

Some teachers prefer to write current in small case “i”

1 Faraday = 96500 coulomb = Charge of one mole electrons

One faraday is the charge required to liberate or deposit one gm equivalent of a substance at corresponding electrode.

Let ‘E’ is equivalent weight then ‘E’ gm will be liberated by 96500 coulomb.

∴ 1 Coulomb will liberate $E / 96500$ gm

So by definition $Z = E / 96500$

$W = I \cdot t \cdot E / 96500$

When a gas is evolved at an electrode, then above formula changes as

$$V = \frac{I \cdot t \cdot V_e}{96500}$$

where $V = \text{volume of liberated gas}$, $V_e = \text{equivalent volume of gas}$.

Equivalent volume may be defined as: The volume of gas liberated by 96500 coulomb at STP.

(ii) Second law of electrolysis:
When same amount of charge is passed through different electrolyte solutions connected in series then weight of substances deposited or dissolved at anode or cathode are in ratio of their equivalent weights. i.e. $W_1 / W_2 = E_1 / E_2$

QUALITATIVE ASPECTS OF ELECTROLYSIS

In the electrolysis process we have discussed above, we have taken molten salt as electrolyte, which contains only one cation and anion. Now, if the electrolyte taken contains more than one cation and anion (for example, aqueous solution of the ionic electrolyte), then the cation and anion that will get discharged depends on the ability of cation to get reduced and the ability of anion to get oxidised.

The ability of an ion to get oxidised or reduced depends upon the size, mass, positive charge, negative charge etc. Thus, it is not possible to predict qualitatively that which ion would be discharged first, as one factor might enhance the ability to discharge while the other factor may hamper it. This can only be predicted on the basis of quantitative value assigned based on the cumulative effect of all the factors responsible for an ion’s ability to discharge. The value is referred as standard potential, which is determined by keeping the concentration of ion as 1 M, pressure of gas at 1 atm, and the measurement done at 25°C. For a cation, the standard reduction potential (SRP) values are compared. The cation having higher standard reduction potential value is discharged in preference to cation with lower SRP value provided the ions are at 1 M concentration. For an anion, the standard oxidation potential (SOP) values are compared and anion having higher SOP is preferentially discharged, if the concentration is 1 M for each of the ion. The SRP values at 25°C for some of the reduction half reactions are given in the tables.
When solution of an electrolyte contains more than one type of cations and anions at concentrations different than 1 M, the discharge of an ion does not depend solely on standard potentials but also depends on the concentration of ion in the solution. This value is referred as potential, called as reduction potential for cation and oxidation potential for anion. The relation between reduction potential and standard reduction potential is given by Nernst equation, as

$$E_{RP} = E^\circ_{RP} - \frac{RT}{nF} \ln \left( \frac{\text{concentration of product}}{\text{concentration of reactant}} \right)$$

Where $E_{RP}$ = Reduction potential of cation

$E^\circ_{RP}$ = Standard Reduction potential of cation

Thus, it is possible that a cation ($A^+$) with lower standard reduction potential getting discharged in preference to cation ($B^+$) having higher standard reduction potential because their concentration might be such that the reduction potential of $A^+$ is higher than that of $B^+$. When two metal ions in the solution have identical values of their reduction potentials, the simultaneous deposition of both the metals will occur in the form of an alloy.
Walther Hermann Nernst (1864 - 1941) German Physicist & Chemist received the Nobel prize in 1920 Chemistry.

Note on Nernst Equation also uses Electrode potential

\[
E_{\text{cell}} = E^0_{\text{cell}} - \frac{0.059}{n} \log \left( \frac{[M^{n+}]}{[M^n]} \right)
\]

At 25°C:

\[
E^0_{\text{cell}} = \frac{0.059}{n} \log \left( \frac{\text{Products}}{\text{Reactants}} \right)
\]

\[
E^0_{\text{cell}} = E^0_{\text{anode}} - E^0_{\text{cathode}} = \frac{0.059}{n} \log \left( \frac{[P]}{[R]} \right)
\]

GALVANIC CELL

This cell converts chemical energy into electrical energy.

---

\[
\text{Cu}^{2+} + 2 \text{e}^- \rightarrow \text{Cu(s)} + 0.34 \text{ V}
\]

\[
\text{Zn(s)} \rightarrow \text{Zn}^{2+} + 2 \text{e}^- + 0.76 \text{ V}
\]

\[
\text{Cu}^{2+} + \text{Zn(s)} \rightarrow \text{Zn}^{2+} + \text{Cu(s)} + 0.76 \text{ V}
\]
Galvanic cell is made up of two half cells i.e., anodic and cathodic. The cell reaction is of redox kind. Oxidation takes place at anode and reduction at cathode. It is also known as voltaic cell. It may be represented as shown in Fig. Zinc rod immersed in ZnSO4 behaves as anode and copper rod immersed in CuSO4 behaves as cathode.

Remember:

In the electrode potential chart/list the elements higher up will act as Cathode compared to elements lower in the chart.

Observe that Cu has positive electrode potential, and is higher up compared to Hydrogen. So Cu rod acts as Cathode.

Zinc has negative electrode potential. Zinc is below SHE (Standard Hydrogen Electrode), so below Hydrogen, in the chart. So Zn rod acts as Anode.

- In galvanic cell like Daniell cell; electrons flow from anode (zinc rod) to the cathode (copper rod) through external circuit; zinc dissolves as Zn$^{2+}$; Cu$^{2+}$ ion in the cathode cell picks up two electron and become deposited at cathode.

- The e.m.f of the standard cell Zn | Zn$^{2+}$ || Ag$^+$ | Ag $E^{\circ}$ cell = 1.562 Volt

If the standard electrode potential of Zinc is -0.762 Volt, then that of Silver is?

Solution: $E^{\circ}$ cell = $E_{Right}$ - $E_{Left}$ = $E_{Ag^+ | Ag}$ - $E_{Zn^{2+} | Zn}$ = ($E_{Reduction (Right)}$ - $E_{Reduction (Left)}$)

So 1.562 = $E_{Ag^+ | Ag}$ - (-0.762) => $E_{Ag^+ | Ag}$ = 0.8 Volt

- What will happen if a solution of copper sulphate put into an iron pot?

Note: Electrode potential of Cu is more positive (so higher) than Fe (Iron).

A solution of copper sulphate put into an iron pot will damage. This is because the standard electrode potential for Fe$^{2+}$/Fe is more negative than that for Cu$^{2+}$/Cu. The iron oxidises first, while copper ions are reduced. Iron will corrode (rust). Or in more simple words it is easier to replace Cu ions by Fe ions from the pot. Any element lower in the series tends to replace the element which is higher up the series.

This is called a redox reaction, since the iron is being oxidised, and copper ions reduced.

Note: A solution of Iron Salt in a Copper Vessel would have just done fine. In this case copper is in solid form, as vessel. So will not replace the Iron ions.
<table>
<thead>
<tr>
<th>Half-Reaction</th>
<th>E (V)</th>
</tr>
</thead>
<tbody>
<tr>
<td>$\text{F}_2(g) + 2\text{e}^- \rightarrow 2\text{F}^-(aq)$</td>
<td>$+2.87$</td>
</tr>
<tr>
<td>$\text{Cl}_2(g) + 2\text{e}^- \rightarrow 2\text{Cl}^-(aq)$</td>
<td>$+1.36$</td>
</tr>
<tr>
<td>$\text{MnO}_2(g) + 4\text{H}^+(aq) + 2\text{e}^- \rightarrow \text{Mn}^{2+}(aq) + 2\text{H}_2\text{O}(l)$</td>
<td>$+1.23$</td>
</tr>
<tr>
<td>$\text{NO}_3^-(aq) + 4\text{H}^+(aq) + 3\text{e}^- \rightarrow \text{NO}(g) + 2\text{H}_2\text{O}(l)$</td>
<td>$+0.96$</td>
</tr>
<tr>
<td>$\text{Ag}^+(aq) + \text{e}^- \rightarrow \text{Ag}(s)$</td>
<td>$+0.80$</td>
</tr>
<tr>
<td>$\text{Fe}^{3+}(aq) + \text{e}^- \rightarrow \text{Fe}^{2+}(aq)$</td>
<td>$+0.77$</td>
</tr>
<tr>
<td>$\text{O}_2(g) + 2\text{H}_2\text{O}(l) + 4\text{e}^- \rightarrow 4\text{OH}^-(aq)$</td>
<td>$+0.40$</td>
</tr>
<tr>
<td>$\text{Cu}^{2+}(aq) + 2\text{e}^- \rightarrow \text{Cu}(s)$</td>
<td>$+0.34$</td>
</tr>
<tr>
<td>$2\text{H}^+(aq) + 2\text{e}^- \rightarrow \text{H}_2(g)$</td>
<td>$0.00$</td>
</tr>
<tr>
<td>$\text{N}_2(g) + 5\text{H}^+(aq) + 4\text{e}^- \rightarrow 2\text{H}_5\text{N}^+(aq)$</td>
<td>$-0.23$</td>
</tr>
<tr>
<td>$\text{Fe}^{2+}(aq) + 2\text{e}^- \rightarrow \text{Fe}(s)$</td>
<td>$-0.44$</td>
</tr>
<tr>
<td>$2\text{H}_2\text{O}(l) + 2\text{e}^- \rightarrow \text{H}_2(g) + 2\text{OH}^-(aq)$</td>
<td>$-0.83$</td>
</tr>
<tr>
<td>$\text{Na}^+(aq) + \text{e}^- \rightarrow \text{Na}(s)$</td>
<td>$-2.71$</td>
</tr>
<tr>
<td>$\text{Li}^+(aq) + \text{e}^- \rightarrow \text{Li}(s)$</td>
<td>$-3.05$</td>
</tr>
</tbody>
</table>
A more detailed table

<table>
<thead>
<tr>
<th>Half-Reaction</th>
<th>$E^\circ$ (V)</th>
<th>Half-Reaction</th>
<th>$E^\circ$ (V)</th>
</tr>
</thead>
<tbody>
<tr>
<td>$F_2 + 2e^- \rightarrow 2F^-$</td>
<td>2.87</td>
<td>$O_3 + 2H_2O + 4e^- \rightarrow 4OH^-$</td>
<td>0.40</td>
</tr>
<tr>
<td>$Ag_2O + e^- \rightarrow Ag^+$</td>
<td>1.99</td>
<td>$Cu^{2+} + 2e^- \rightarrow Cu$</td>
<td>0.34</td>
</tr>
<tr>
<td>$Co^{4+} + e^- \rightarrow Co^{3+}$</td>
<td>1.82</td>
<td>$HgCl_2 + 2e^- \rightarrow Hg + 2Cl^-$</td>
<td>0.27</td>
</tr>
<tr>
<td>$H_2O_2 + 2H^+ + 2e^- \rightarrow 2H_2O$</td>
<td>1.78</td>
<td>$AgCl + e^- \rightarrow Ag + Cl^-$</td>
<td>0.22</td>
</tr>
<tr>
<td>$Cu^{2+} + e^- \rightarrow Cu^{+}$</td>
<td>1.70</td>
<td>$SO_3^{2-} + 4H^+ + 2e^- \rightarrow H_2SO_3 + H_2O$</td>
<td>0.20</td>
</tr>
<tr>
<td>$Ph_2O_4 + 4H^+ + SO_2^{3-} + 2e^- \rightarrow PbSO_4 + 2H_2O$</td>
<td>1.69</td>
<td>$Ca^{2+} + e^- \rightarrow Ca^{+}$</td>
<td>0.16</td>
</tr>
<tr>
<td>$MnO_4^- + 4H^+ + 3e^- \rightarrow Mn^{2+} + 2H_2O$</td>
<td>1.68</td>
<td>$2H^+ + 2e^- \rightarrow H_2$</td>
<td>0.00</td>
</tr>
<tr>
<td>$2e^- + 3H^+ + IO_3^- \rightarrow IO_2^- + H_2O$</td>
<td>1.60</td>
<td>$Fe^{3+} + 3e^- \rightarrow Fe$</td>
<td>-0.036</td>
</tr>
<tr>
<td>$MnO_4^- + 8H^+ + 5e^- \rightarrow Mn^{2+} + 4H_2O$</td>
<td>1.51</td>
<td>$Fe^{3+} + 2e^- \rightarrow Fe^{2+}$</td>
<td>-0.13</td>
</tr>
<tr>
<td>$Ar^{3+} + 3e^- \rightarrow Ar$</td>
<td>1.50</td>
<td>$Sn^{2+} + 2e^- \rightarrow Sn$</td>
<td>-0.14</td>
</tr>
<tr>
<td>$PSO_4 + 4H^+ + 2e^- \rightarrow Pb^{2+} + 2H_2O$</td>
<td>1.46</td>
<td>$Np^{2+} + 2e^- \rightarrow Np$</td>
<td>-0.23</td>
</tr>
<tr>
<td>$Cl_2 + 2e^- \rightarrow 2Cl^-$</td>
<td>1.36</td>
<td>$PbSO_4 + 2e^- \rightarrow Pb + SO_4^{2-}$</td>
<td>-0.35</td>
</tr>
<tr>
<td>$Co_2O_4^{2-} + 14H^+ + 6e^- \rightarrow 2Co^{3+} + 7H_2O$</td>
<td>1.33</td>
<td>$Cd^{2+} + 2e^- \rightarrow Cd$</td>
<td>-0.40</td>
</tr>
<tr>
<td>$O_2 + 4H^+ + 4e^- \rightarrow 2H_2O$</td>
<td>1.23</td>
<td>$Fe^{3+} + 2e^- \rightarrow Fe^{2+}$</td>
<td>-0.44</td>
</tr>
<tr>
<td>$MnO_4^- + 4H^+ + 2e^- \rightarrow Mn^{2+} + 2H_2O$</td>
<td>1.21</td>
<td>$Fe^{3+} + e^- \rightarrow Fe^{2+}$</td>
<td>-0.59</td>
</tr>
<tr>
<td>$IO_3^- + 6H^+ + 5e^- \rightarrow I^- + 3H_2O$</td>
<td>1.20</td>
<td>$C_2^{2+} + 3e^- \rightarrow Cr$</td>
<td>-0.73</td>
</tr>
<tr>
<td>$Br_2 + 2e^- \rightarrow 2Br^-$</td>
<td>1.09</td>
<td>$Zn^{2+} + 2e^- \rightarrow Zn$</td>
<td>-0.76</td>
</tr>
<tr>
<td>$VO_2^{+} + 2H^+ + e^- \rightarrow VO^{3+} + H_2O$</td>
<td>1.00</td>
<td>$2H_2O + 2e^- \rightarrow H_2 + 2OH^-$</td>
<td>-0.83</td>
</tr>
<tr>
<td>$MnO_4^- + 3e^- \rightarrow Mn^{2+} + 4Cl^-$</td>
<td>0.99</td>
<td>$Mn^{2+} + 2e^- \rightarrow Mn$</td>
<td>-1.18</td>
</tr>
<tr>
<td>$NO_3^- + 4H^+ + 3e^- \rightarrow NO + 2H_2O$</td>
<td>0.96</td>
<td>$AP^{3+} + 3e^- \rightarrow Al$</td>
<td>-1.66</td>
</tr>
<tr>
<td>$ClO_3^- + e^- \rightarrow ClO_2^-$</td>
<td>0.94</td>
<td>$H_2 + 2e^- \rightarrow 2H$</td>
<td>-2.22</td>
</tr>
<tr>
<td>$2Hg^{+} + 2e^- \rightarrow Hg^2+$</td>
<td>0.91</td>
<td>$Mg^{2+} + 2e^- \rightarrow Mg$</td>
<td>-2.37</td>
</tr>
<tr>
<td>$Ag^+ + e^- \rightarrow Ag$</td>
<td>0.80</td>
<td>$La^{3+} + 3e^- \rightarrow La$</td>
<td>-2.37</td>
</tr>
<tr>
<td>$Hg^{2+} + 2e^- \rightarrow Hg$</td>
<td>0.80</td>
<td>$Na^+ + e^- \rightarrow Na$</td>
<td>-2.71</td>
</tr>
<tr>
<td>$Fe^{3+} + e^- \rightarrow Fe^{2+}$</td>
<td>0.77</td>
<td>$Ca^{2+} + 2e^- \rightarrow Ca$</td>
<td>-2.76</td>
</tr>
<tr>
<td>$O_2 + 2H^+ + 2e^- \rightarrow H_2O_2$</td>
<td>0.68</td>
<td>$Ba^{2+} + 2e^- \rightarrow Ba$</td>
<td>-2.90</td>
</tr>
<tr>
<td>$MnO_4^- + e^- \rightarrow MnO_4^{2-}$</td>
<td>0.56</td>
<td>$K^+ + e^- \rightarrow K^+$</td>
<td>-2.92</td>
</tr>
<tr>
<td>$I_2 + 2e^- \rightarrow 2I^-$</td>
<td>0.54</td>
<td>$Li^+ + e^- \rightarrow Li$</td>
<td>-3.05</td>
</tr>
<tr>
<td>$Cu^{+} + e^- \rightarrow Cu$</td>
<td>0.52</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Spoon Feeding

What will happen if a solution of Magnesium sulphate put into an Copper pot ?

Note : Magnesium looses electron very easily, metallic character, or basic character being high. Electrode potential is - 2.37 While we know Copper has +ve electrode potential. ( This is an important exception and the data is better remembered as 0.5 )

So Copper is solid in Copper pot ( or even if a copper spoon was inserted, the same logic ). Copper will not replace Magnesium ions. But if a Magnesium spoon was put into Copper-Sulphate solution then the Magnesium ions / atoms will start replacing the Cu ions and Copper will start getting precipitated. The Magnesium spoon, or container will start getting corroded or damaged.
SALT BRIDGE

Two electrolyte solutions in galvanic cells are separated using salt bridge joining the two sides. Saltbridge is a device to minimize or eliminate the liquid junction potential. Saturated solution of salts like KCl, KNO₃, NH₄Cl and NH₄NO₃ etc. in agar-agar gel is used in salt bridge. Salt bridge contains high concentration of ions viz. K⁺ and NO₃⁻ at the junction with electrolyte solution. Thus, salt bridge carries whole of the current across the boundary; more over the K⁺ and NO₃⁻ ions have same speed. Hence, salt bridge with uniform and same mobility of cations and anions minimize the liquid junction potential & completes the electrical circuit & permits the ions to migrate.
Representation of a cell (IUPAC conventions): Let us illustrate the convention taking the example of Daniel cell.

(i) Anodic half-cell is written on left and cathodic half-cell on right hand side.

\[ \text{Zn(s)} \mid \text{ZnSO}_4(\text{sol}) \mid \text{CuSO}_4(\text{sol}) \mid \text{Cu(s)} \]

(ii) Two half cells are separated by double vertical lines: Double vertical lines indicate salt bridge or any type of porous partition.

(iii) EMF (electromotive force) may be written on the right hand side of the cell.

(iv) Single vertical lines indicate the phase separation between electrode and electrolyte solution.

\[ \text{Zn} \mid \text{Zn}^{2+} \parallel \text{Cu}^{2+} \parallel \text{Cu} \]

(In Illustration of Phase boundary)

(v) Inert electrodes are represented in the bracket

\[ \text{Zn} \mid \text{ZnSO}_4 \parallel \text{H}^+ \parallel \text{H}_2, \text{Pt} \]

Consider the following set of half-reactions

\[
\begin{align*}
\text{Reduction} & : \quad \text{Fe}^{3+} + 2e^- \rightleftharpoons \text{Fe} \\
E_{\text{red}}^\circ & = -0.44 \text{ V} \\
\text{Oxidation} & : \quad \text{Zn} \rightleftharpoons \text{Zn}^{2+} + 2e^- \\
E_{\text{ox}}^\circ & = +0.76 \text{ V}
\end{align*}
\]
Le Châtelier’s principle gives us some ideas. Look at the overall reaction. Consider the reaction to be at equilibrium ($E_{\text{cell}} = 0 \text{ V}$). If we add more Fe$^{2+}$ to the reaction mixture, The equilibrium will shift to the right to use up the excess Fe$^{2+}$. This will, of course cause a positive cell potential to be measurable.

Conversely, if we removed Fe$^{2+}$ from the reaction mixture, the equilibrium would shift to the left to try to replace some of the missing Fe$^{2+}$. This would result in a measurable negative potential (reaction goes left).

If the reaction were not at equilibrium then these preceding two changes would me additive to the measured voltage (say standard voltage). So, if we add Fe$^{2+}$ to a mixture at Standard Conditions, then the voltage would be more positive and if we removed some Fe$^{2+}$ from a mixture at Standard Conditions the measured voltage would be lower (less positive).

The reverse results would be found for addition or removal of Zn$^{2+}$ (a product). More Zn$^{2+}$ would give a lower positive potential and vice versa.

Remember the relation: $\Delta G = \Delta G^\circ + RT \ln Q$ 

Nernst Equation for non-standard conditions

Substitute the new definition of $\Delta G$ and we get

$$-nFE = -nFE^\circ + RT \ln Q$$

$$E = E^\circ - \frac{RT}{nF} \ln Q$$

( This is again called the Nernst Equation of another form ).

Thus, we can calculate the cell potential of any cell, given only the initial conditions (to calculate Q) and the reduction potentials (to calculate $E^\circ$)

We can calculate the voltage for the situation where the concentrations are not standard. For example, consider $[\text{Fe}^{2+}] = 0.1 \text{ M}$ and $[\text{Zn}^{2+}] = 1.9 \text{ M}$

For this reaction, we also know that there are two electrons transferred for each equation ($n = 2$).

$$E = E^\circ - \frac{RT}{nF} \ln \left( \frac{[\text{Zn}^{2+}]}{[\text{Fe}^{2+}]} \right)$$

$$= 0.32V - \frac{8.3145 \text{ J K}^{-1} \text{ mol}^{-1} \times 298.15K}{2 \times 95485 \text{ C mol}^{-1}} \times \ln \frac{1.9}{0.10}$$

$$= 0.28V$$

Although the reaction is 90% complete (from Standard Conditions initial) the cell potential has only dropped by a small amount (0.04 V). This is normal. For example, the batteries in your flashlight will put out almost full voltage until the last traces of chemical are almost used up at which point the voltage will drop off rather sharply. This is an especially useful characteristic of cells powering electronic equipment.
In general, for a redox cell reaction involving the transference of \( n \) electrons \( aA + bB \rightarrow cC + dD \), the EMF can be calculated as:

\[
E_{\text{cell}} = E_0 - \frac{0.0591}{n} \log \frac{[C]^c[D]^d}{[A]^a[B]^b}
\]

Prediction and feasibility of spontaneity of a cell reaction.

Work done by the cell = \( nFE \);

It is equivalent to decrease in free energy \( \Delta G = -nFE \)

Under standard state \( \Delta G^0 = -nFE^0 \) \quad (i)

(i) From thermodynamics we know, \( \Delta G \) = negative for spontaneous process. Thus from eq. (i) it is clear that the EMF should be positive for a cell process to be feasible or spontaneous.

(ii) When \( \Delta G = \) positive, \( E = \) negative and the cell process will be non spontaneous.

(iii) When \( \Delta G = 0 \), \( E = 0 \) and the cell will attain the equilibrium.

<table>
<thead>
<tr>
<th>Reactions</th>
<th>( \Delta G )</th>
<th>( E )</th>
</tr>
</thead>
<tbody>
<tr>
<td>Spontaneous</td>
<td>(–)</td>
<td>(+)</td>
</tr>
<tr>
<td>Non- spontaneous</td>
<td>(+)</td>
<td>(–)</td>
</tr>
<tr>
<td>Equilibrium</td>
<td>0</td>
<td>0</td>
</tr>
</tbody>
</table>

Standard free energy change of a cell may be calculated by electrode potential data. Substituting the value of \( E_0 \) (i.e., standard reduction potential of cathode- standard reduction potential of anode) in eq. (i) we may get \( \Delta G^0 \).

THERMODYNAMIC TREATMENT OF NERNST EQUATION

Determination of equilibrium constant

\[
E^0 = \frac{0.0591}{n} \log K_{\text{eq}}
\]

\( : \) We know, that \( E = \) .... (1)

At equilibrium, the cell potential is zero because cell reactions are balanced, i.e. \( E = 0 \)

\( \therefore \) From Eq. (1), we have

\[
E^0 = \frac{0.0591}{n} \log K_{\text{eq}}
\]

Or

\[
K_{\text{eq}} = \text{antilog} \left( \frac{nE^0}{0.0591} \right)
\]

Recall antilog is 10 to the power.

Heat of Reaction inside the cell:

Let \( n \) Faraday charge flows out of a cell of e.m.f. \( E \), then
\[ -\Delta G = nFE \ldots (i) \]

Gibbs Helmholtz equation (from thermodynamics) may be given as,

\[ \Delta G = \Delta H + T \left[ \frac{\partial \Delta G}{\partial T} \right]_p \]  

(ii)

From Eqs. (i) and (ii), we have

\[ -nFE = \Delta H + T \left[ \frac{\partial (-nFE)}{\partial T} \right]_p = \Delta H - nFT \left[ \frac{\partial E}{\partial T} \right]_p \]

Thus

\[ \Delta H = nFE + nFT \left[ \frac{\partial E}{\partial T} \right]_p \]

Entropy change inside the cell:

We know that \( G = H - TS \) or
\[ \Delta G = \Delta H - T \Delta S \ldots (i) \]

where

\( \Delta G \) = Free energy change;
\( \Delta H \) = Enthalpy change and
\( \Delta S \) = entropy change.

According to Gibbs Helmholtz equation,

\[ \Delta G = \Delta H + T \left[ \frac{\partial \Delta G}{\partial T} \right]_p \]

\[ \Delta G = \Delta H = T \left[ \frac{\partial \Delta G}{\partial T} \right]_p \]

From Eqs. (i) and (ii), we have

\[ -T \Delta S = T \left[ \frac{\partial \Delta G}{\partial T} \right]_p \quad \text{or} \quad \Delta S = - \left[ \frac{\partial \Delta G}{\partial T} \right]_p \]

or

\[ \Delta S = nF \left[ \frac{\partial E}{\partial T} \right]_p \]

Where \( \left[ \frac{\partial E}{\partial T} \right]_p \) is called temperature coefficient of cell e.m.f.
Spoon Feeding Rule of Electrochemistry

\[
A \rightarrow A^{+2} + 2e^- \quad \Delta G_1 = -0.2 \text{kcal} \quad \cdots (i)
\]
\[
B \rightarrow B^{+2} + 2e^- \quad \Delta G_2 = +0.4 \text{kcal} \quad \cdots (ii)
\]

According to the rule of spontaneity, reaction (i) is spontaneous but reaction (ii) is non-spontaneous. To gain full cell reaction, we convert non-spontaneous reaction into spontaneous reaction.

\[
\begin{align*}
\text{Same:} & \quad A \rightarrow A^{+2} + 2e^- \quad \Delta G_1 = -0.2 \text{kcal} \\
\text{Change:} & \quad B^{+2} + 2e^- \rightarrow B \quad \Delta G_2 = +0.4 \text{kcal}
\end{align*}
\]

\[
\Delta G_{\text{rxn}} = \Delta G_1 + \Delta G_2 = (-0.2 \text{kcal}) + (+0.4 \text{kcal}) = +0.2 \text{kcal}
\]

Recall \(\Delta G = -nFE\) or \(E = -\frac{\Delta G}{nF}\)

Spoon feeding Rule of Electrochemistry

If both half cell reactions are non-spontaneous, then we convert more non-spontaneous reaction into spontaneous reaction to gain full cell reaction

\[
\begin{align*}
\text{A} & \rightarrow A^{+2} + 2e^- \quad \Delta G_1 = +0.2 \text{kcal} \quad \cdots (i) \\
\text{B} & \rightarrow B^{+2} + 2e^- \quad \Delta G_2 = +0.5 \text{kcal} \quad \cdots (ii)
\end{align*}
\]

Reaction (i) is less non-spontaneous and reaction (ii) is more non-spontaneous.

\[
\begin{align*}
\text{Change:} & \quad B^{+2} + 2e^- \rightarrow B \quad \Delta G_2 = +0.5 \text{kcal} \\
\text{Same:} & \quad A \rightarrow A^{+2} + 2e^- \quad \Delta G_1 = +0.2 \text{kcal}
\end{align*}
\]

\[
\Delta G_{\text{rxn}} = \Delta G_1 + \Delta G_2 = (+0.2 \text{kcal}) + (+0.5 \text{kcal}) = +0.7 \text{kcal}
\]

Recall Spontaneity of \(\Delta G\)

More Positive lesser spontaneous

More negative more spontaneous

So if both half cell reactions are spontaneous then we convert less spontaneous reaction into non-spontaneous reaction.
If $\Delta G^\circ$ given in the half cells then spontaneity of the reaction not determined by $\Delta G^\circ$. We convert $\Delta G^\circ$ into $\Delta G$ by following equation.

$$\Delta G = -nFE$$

Greater the reduction potential greater will be ease of reduction. Greater the oxidation potential greater will be ease of oxidation.
A voltaic cell is made of aqueous Bromine and Zinc. Given -

\[ \text{Br}_2(aq) + \text{Zn}(s) \rightarrow \text{Zn}^{2+}(aq) + 2\text{Br}^-(aq) \quad E_{\text{cell}} = 1.83 \ \text{V}. \]

Calculate \( E_{\text{bromine}} \) given that \( E_{\text{zinc}} = -0.76 \ \text{V} \)

**Solution:**

\[ \text{Br}_2(aq) + 2e^- \rightarrow 2\text{Br}^-(aq) \quad [\text{reduction; cathode}] \]
\[ \text{Zn}(s) \rightarrow \text{Zn}^{2+}(aq) + 2e^- \quad [\text{oxidation; anode}] \quad E_{\text{zinc}} = -0.76 \ \text{V} \]
What is the equilibrium constant for the reaction of copper metal with bromine to form copper(II) ions and bromide ions in aqueous solution at 25°C?

\[ \text{Cathode} \quad Br_2 + 2e^- \rightleftharpoons 2 Br^- \quad E^\circ_{\text{red}} = 1.09 \text{ V} \]

\[ \text{Anode} \quad Cu \rightleftharpoons Cu^{2+} + 2e^- \quad E^\circ_{\text{ox}} = -E^\circ_{\text{red}} = -0.34 \text{ V} \]

\[ \text{Cell} \quad Cu + Br_2 \rightleftharpoons Cu^{2+} + 2 Br^- \quad E^\circ_{\text{cell}} = 0.75 \]

\[ \ln K = \frac{nFE^\circ}{RT} = \frac{2 \times 96485 \text{ J mol}^{-1} \text{F} \times 0.75 \text{ V}}{8.3145 \text{ J K}^{-1} \text{mol}^{-1} \times 298.15 \text{ K}} = 58.38 \]

\[ K = e^{58.38} = 2.2 \times 10^{25} \]

Let us always remember the basics. If something, say M, looses electrons, it is getting oxidized. Metals are basic. Metals loose electrons and gets oxidized. The metals which loose electron very easily, get oxidized very easily. So these are better reducing agent.

Adding of Oxygen, or Halogens ( F, Cl, Br, I ) or other p-Block elements which accept electrons, is oxidation. So if Potassium ( K ) gets Oxygen ( attaches to O ) to form K₂O or KO₂ ( Superoxide ), K is getting oxidized. If Cs attaches with F then Cs is getting oxidized, while F is being reduced. Cs is a reducing agent as it is losing electrons. F is an oxidizing agent as it is gaining electrons.

Adding of Hydrogen is reduction. So if Ethene receives Hydrogen to become Ethane, then Ethene is being reduced. Raney Nickel is Nickel with molecular level pores. Raney Ni with HCl gives Hydrogen atoms, ( some teachers say this as Nascent Hydrogen ) locally, meaning near the molecule where the reaction is needed. So Raney Ni + HCl is a good reducing agent. Same for Zinc dust + HCl. But then what is Birch reduction ?

As my experience of teaching more than 25 years, almost every student understands these. Knowing this is essential but does not ensure success in chemistry. There are many kinds of details of oxidizing agent or Reducing agent that vary from reactions / situations.

Using half reactions write the spontaneous Redox reactions

\[ \text{Sn}^{2+}(aq) + 2e^- \rightarrow \text{Sn}(s) \quad E^\circ_{\text{Sn}} = -0.14 \text{ V} \]

\[ \text{Ag}^{+}(aq) + e^- \rightarrow \text{Ag}(s) \quad E^\circ_{\text{Ag}} = 0.80 \text{ V} \]

We have to multiply the Silver reaction with 2 to balance and match electrons

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Now we need to add the half reactions to eliminate the electrons, but subtract to get the Voltage correctly. Recall \( E^\circ_{\text{cell}} = E_{\text{Right}} - E_{\text{Left}} = E_{\text{Cathode}} - E_{\text{Anode}} = (E_{\text{Reduction (Right)}} - E_{\text{Reduction (Left)}}) \)

\[
\begin{align*}
\text{Cathode:} & \quad \text{Sn(s) → Sn}^{2+}(aq) + 2e^- \quad E^o_{\text{Sn}} = -0.14 \text{ V} \\
\text{Anode:} & \quad 2\text{Ag}^+(aq) + 2e^- → 2\text{Ag(s)} \quad E^o_{\text{Ag^+}} = 0.80 \text{ V} \\
\text{Overall:} & \quad \text{Sn(s) + 2Ag}^+(aq) → 2\text{Ag(s)} + \text{Sn}^{2+}(aq) \quad E^o_{\text{cell}} = 0.94 \text{ V}
\end{align*}
\]

\[E^o_{\text{cell}} = E^o_{\text{Ag^+}} - E^o_{\text{Sn}} = 0.80 - (-0.14) = 0.94 \text{ V}\]

Cathode provides electrons. Ag ions are receiving electrons at Cathode. Anode is positive where electrons are lost. See Tin is more metallic than Silver and is looses electrons in Anode.

It is easier to visualize the Voltage as gap in electrode potential. Silver is higher up than Hydrogen in the series. Thus has positive electrode potential. While Tin is more metallic, below Hydrogen in the series. The gap is of 0.94 Volts.

The concept of electron transfer can easily explain the redox reaction in case of ionic substances. However, for covalent compounds oxidation and reduction or redox reactions are explained using oxidation number.

The term oxidation was first used to mean the addition of oxygen to an element or compound, or the removal of hydrogen from a compound. Reduction meant the addition of hydrogen to an element or compound, or the removal of oxygen from a compound. Such definitions have been extended and now many oxidation-reduction, or redox, reactions are best interpreted in terms of transfer of electrons.

- Oxidation is defined as the loss of electrons by a chemical species (atom, ion or molecule).
- Reduction is the gain of electrons by a chemical species (atom, ion or molecule).
- An oxidising agent that chemical species which takes electrons thus it is an electron acceptor.
- A reducing agent is the chemical species that gives electrons and thus acts as an electron donor.
- Redox reactions are those chemical reactions in which both oxidation as well as reduction occur simultaneously.
- The substance which undergo reduction is called oxidising agent while the substance which undergo oxidation is called reducing agent. One can say that the substance that causes the oxidation of any substance in reaction is called the oxidizing agent while the substance that causes the reduction is called the reducing agent.
- Neither reduction nor oxidation occurs alone. Both of them occur simultaneously. Since both these reactions must occur at the same time they are often termed as 'redox reactions'. The oxidation or reduction portion of a redox reaction, including the electrons gained or lost can be determined by means of a Half-Reaction.
Two Types of Redox Reactions

(i) Inter molecular Redox Reactions:

In these kinds of redox reactions, one molecule of reactant is oxidized whereas molecule of other reactant is reduced.

\[ \text{NO}_3^- + 3H_2S + H_2O + H^+ \rightarrow NH_4^+ + HSO_4^- \]

Oxidised Reduced

(ii) Intra molecular Redox Reactions:

One atom of a molecule is oxidized and other atom of same molecule is reduced then it is intra molecular redox reaction.

\[ 2\text{KClO}_3 \rightarrow 2\text{KCl}^- + 3\text{O}_2 \]

(1) Molecular Equations:

When the reactant and products involved in a chemical change are written in molecular form in a chemical equation, it is termed as molecular equation.

Example: \( \text{MnO}_2 + 4\text{HCl} \rightarrow \text{MnCl}_2 + 2\text{H}_2\text{O} + \text{Cl}_2 \)

In above example the reactant and products have been written in molecular forms, thus it is a molecular equation.

(2) Ionic Equation:

When the reactant and products involved in a chemical change are ionic compounds, these will be present in the form of ions in the solution. The chemical change is written in ionic forms in the chemical equation, it is termed as ionic equation.

Example: \( \text{MnO}_2 + 4\text{H}^+ + 4\text{Cl}^- \rightarrow \text{Mn}^{2+} + 2\text{Cl}_2^- + 2\text{H}_2\text{O} + \text{Cl}_2 \)

In the above example, the reactant and products have been written in ionic forms, thus the equation is termed as ionic equation.

The rules to be followed for writing ionic equations are:
All soluble ionic compounds involved in a chemical change are expressed in ionic symbols and covalent substances are written in molecular form. \( \text{H}_2\text{O}, \text{NH}_3, \text{NO}_2, \text{NO}, \text{SO}_2, \text{CO}, \text{CO}_2 \), etc., are expressed in molecular form.

The ionic compound which is highly insoluble is expressed in molecular form.

The ions which are common and equal in number on both sides, i.e., spectator ions, are cancelled.

Besides the atoms, the ionic charges must also be balanced on both the sides.

**3) Spectator Ions:**

Species that are present in the solution but not take part in the reaction and are also omitted while writing the net ionic equation are called spectator ions or bystander ions.

\[
\text{Zn} + 2\text{H}^+ + 2\text{Cl}^- \rightarrow \text{Zn}^{2+} + 2\text{Cl}^- + \text{H}_2
\]

In this reaction, ions are omitted and are called as spectator ions and appear on the reactant as well as product side.

**4) Oxidising agent:**

The substance (atom, ions or molecules) that gain electrons and is thereby reduced to a low valency state is called an oxidising agent.

**5) Reducing agent:**

The substance that loses electrons and its valency thereby oxidised to a higher valency state is called a reducing agent.

It is not always possible to balance redox equations using the simple “inspection” technique. The following unbalanced net ionic equation provides an example.

\[
\text{Au}^{3+}(\text{aq}) + \text{I}^- (\text{aq}) \rightarrow \text{Au}(\text{s}) + \text{I}_2(\text{s})
\]

At first glance, it seems that this equation can be balanced by placing a 2 in front of the \( \text{I}^- \).

\[
\text{Au}^{3+}(\text{aq}) + 2\text{I}^- (\text{aq}) \rightarrow \text{Au}(\text{s}) + \text{I}_2(\text{s})
\]

Note, however, that although the atoms are now balanced, the charge is not. The sum of the charges on the left is +1, and the sum of the charges on the right is zero, as if the products could somehow have one more electron than the reactants. To correctly balance this equation, it helps to look more closely at the oxidation and reduction that occur in the reaction. The iodine atoms are changing their oxidation number from −1 to 0, so each iodide ion must be losing one electron. The \( \text{Au}^{3+} \) is changing to \( \text{Au} \), so each gold(III) cation must be gaining three electrons.

The half-reactions are:

\[
\text{I}^- (\text{aq}) \rightarrow \frac{1}{2}\text{I}_2(\text{s}) + \text{e}^-
\]

\[
\text{Au}^{3+}(\text{aq}) + 3\text{e}^- \rightarrow \text{Au}(\text{s})
\]
We know that in redox reactions, the number of electrons lost by the reducing agent must be equal to the number of electrons gained by the oxidizing agent; thus, for each $\text{Au}^{3+}$ that gains three electrons, there must be three $\text{I}^-$ ions that each lose one electron. If we place a 3 in front of the $\text{I}^-$ and balance the iodine atoms with a $3/2$ in front of the $\text{I}_2$, both the atoms and the charge will be balanced.

$$\text{Au}^{3+} (\text{aq}) + 3\text{I}^- (\text{aq}) \rightarrow \text{Au}(s) + 3/2\text{I}_2(s)$$

or

$$2\text{Au}^{3+} (\text{aq}) + 6\text{I}^- (\text{aq}) \rightarrow 2\text{Au}(s) + 3\text{I}_2(s)$$

**Rules For Writing Ionic Equations:**

(i) All soluble electrolytes involved in a chemical change are expresses in ionic symbols and covalent substances are written in molecular form.

(ii) The electrolyte which is highly insoluble, is expressed in molecular form.

(iii) The ions which are common and equal in number on both sides (spectator ions) are cancelled.

(iv) Besides the atoms, the ionic charges must also balance on both the sides.

**Stock's Notation:**

Generally used for naming compounds of metals, some non-metals also. eg. Cr$_2$O$_3$ Chromium (III) oxide and P$_2$O$_5$ Phosphorous (V) oxide.

**Oxidising Agents (oxidants / oxidizers):**

They oxidise others, themselves are reduced & gain electrons. eg. O$_2$, O$_3$, HNO$_3$, MnO$_2$, H$_2$O$_2$, halogens, K$\text{MnO}_4$, K$_2$Cr$_2$O$_7$, KIO$_3$, Cl(SO$_4$)$_3$, FeCl$_3$, NaOCl, hydrogen ions.

[Atoms present in their higher oxidation state.]

**Reducing Agents ( reductants / reducers):**

They reduce others, themselves get oxidised & lose electrons. H$_2$, molecule is weak but Nascent hydrogen is powerful. C, CO, H$_2$S, SO$_2$, SnCl$_2$, Sodium thio Sulphate, Al, Na, CaH$_2$, NaBH$_4$, LiAlH$_4$

[Atoms present in their lower oxidation state].

**Both Oxidising & Reducing Agents:** SO$_2$, H$_2$O$_2$, O$_3$, NO$_2$, etc.
Types Of Redox Reduction:
Intermolecular redox, disproportion, Intra molecular redox.

Disproportion:
In such reactions the oxidising and reducing agents (atom) are the same.

\[
\text{H}_2\text{O}_2 + \text{H}_2\text{O}_2 \rightarrow 2\text{H}_2\text{O} + \text{O}_2
\]

To identify whether a reaction is redox or not, find change in oxidation number or loss and gain of electrons. If there is no change in oxidation number, the reaction is not a redox reaction.

**NOTE:** To predict the product of reaction remember:
(a) Free halogen on reduction gives halide ion (F₂ → F⁻)
(b) Alkali metals on oxidation give metallic ion with +1 oxidation state.
(c) Conc. HNO₃ on reduction gives NO₂, while dilute HNO₃ can give NO, N₂, NH₄⁺ or other products depending on the nature of reducing agent and on dilution.
(d) In acid solution KMnO₄ is reduced to Mn²⁺ while in neutral or alkaline, it gives MnO₂ or K₂MnO₄.
(e) H₂O₂ on reduction gives water and on oxidation gives oxygen.
(f) Dichromate ion in acid solution is reduced to Cr³⁺.

**Nature of oxides based on oxidation number:**

- Lowest oxidation state Basic -> (MnO)
- Intermediate oxidation state Amphoteric -> (Mn₃O₄, MnO₂)
- Highest oxidation state Acidic -> (Mn₂O₇)

**Metathesis Reactions:**
Never redox reactions. In these two compounds react to form two new compounds and no change in oxidation number occurs. eg.
(i) Pb (NO₃)₂ + K₂CrO₄ -> Pb CrO₄ + 2 KNO₃ (ii) HCl + NaOH -> NaCl + H₂O
Rules For Assigning Oxidation Number:
(i) Oxidation number of free elements or atoms is zero.
(ii) Oxidation number of allotropes is zero.
(iii) Oxidation number of atoms in homo-nuclear molecules is zero.
(iv) Oxidation number of mono-atomic ions is equal to the algebraic charge on them.
(v) Oxidation number of F in compounds is -1.
(vi) Oxidation number of H in its compounds is +1, except in metalhydrides where it is -1.
(vii) Oxidation number of O is -2 in its compounds, but in F2O it is +2 and in peroxides it is -1 and -0.5 in KO2.
(viii) Oxidation number of alkali metals in their compounds +1.
(ix) Oxidation number of alkaline earth metals in their compounds is +2.
(x) Oxidation number of an ion is equal to its charge.
(xi) Oxidation number of a molecule as a whole is zero.
(xii) The sum of oxidation number of all the atoms in a molecule should be zero and in an ion equal to its charge.

Average Oxidation Number: Find Oxidation Number of Fe in Fe3O4 Fe3O4 is FeO. Fe2O3.
O. N. of Fe in FeO is +2; O. N. of Fe in Fe2O3 is +3.
Therefore average O. N. of three Fe atoms = (2 + 2 X 3)/3 = 8/3

Concept of Equivalent Weight

(A) Volumetric analysis: This mainly involve titrations based chemistry. It can be divided into two major category. (I) Non-redox system (II) Redox system

(I) Non-redox system: This involve following kind of titrations:
1. Acid-Base titrations
2. Back titration
3. Precipitation titration
4. Double indicator acid base titration

Titrimetric Method of Analysis: A titrimetric method of analysis is based on chemical reaction such as. aA + tT -> Product
Where ‘a’ molecules of “analysis”, A, reacts with t molecules of reagent T.
T is called Titrant normally taken in buret in form of solution of known concentration. The solution of titrant is called “standard solution”.

The addition of titrant is added till the amount of T, chemically equivalent to that of ‘A’ has been added.
It is said equivalent point of titration has been reached. In order to know when to stop addition of titrant, a chemical substance is used called indicator, which respond to appearance of excess of titrant by changing colour precisely at the equivalence point. The point in the titration where the indicator changes colour is termed the ‘end point’. It is possible that end point be as close as possible to the equivalence point.

The term titration refer’s to process of measuring the volume of titrant required to reach the end point.

For many years the term volumetric analysis was used rather than titrimetric analysis. However from a rigorous stand point the term titrimetric is preferable because volume measurement may not be confirmed to titration. In certain analysis, for example one might measure the volume of a gas.

We can adopt mole method in balanced chemical reactions to relate reactant and products but it is more easier to apply law of equivalents in volumetric calculations because it does not require knowledge of balanced chemical reactions involved in sequence.
Law of equivalents refers to that, equivalents of a limiting reactant is equal to equivalent of other reactant reacting in a chemical reaction or equal to equivalents of products formed. \( n \) factor here we mean a conversion factor by which we divide molar mass of substance to get equivalent mass and it depends on nature of substance which vary from one condition to another condition.

We can divide \( n \)-factor calculations in two category.
(A) when compound is not reacting.
(B) when compound is reacting.

**Acid-Base titration**

To find out strength or concentration of unknown acid or base it is titrated against base or acid of known strength. At the equivalence point we can know amount of acid or base used and then with the help of law of equivalents we can find strength of unknown.

\[
\text{Meq of acid at equivalence point} = \text{Meq of base at equivalence point}
\]

**Back titration**

Back titration is used in volumetric analysis to find out excess of reagent added by titrating it with suitable reagent. It is also used to find out percentage purity of sample. For example in acid-base titration suppose we have added excess base in acid mixture. To find excess base we can titrate the solution with another acid of known strength.

**Precipitation titration**

In ionic reaction we can know strength of unknown solution of salt by titrating it against a reagent with which it can form precipitate. For example NaCl strength can be known by titrating it against AgNO3 solution with which it form white ppt. of AgCl.

\[
\text{Meq of NaCl at equivalence point} = \text{meq of AgNO3 used} = \text{meq of AgCl formed}
\]

**Double indicator acid-base titration:**

In the acid-base titration the equivalence point is known with the help of indicator which changes its colour at the end point. In the titration of polyacidic base or polybasic acid there are more than one end point for each step neutralization. Sometimes one indicator is not able to give colour change at every end point. So to find out end point we have to use more than one indicator. For example in the titration of Na2CO3 against HCl there are two end points.

\[
\text{Na2CO3} + \text{HCl} \rightarrow \text{NaHCO3} + \text{NaCl}
\]

\[
\text{NaHCO3} + \text{HCl} \rightarrow \text{H2CO3} + \text{NaCl}
\]

When we use phenolphthalein in the above titration it changes its colour at first end point when NaHCO3 is formed and with it we cannot know second end point. Similarly with methyl orange it changes its colour at second end point only and we cannot know first end point. It is because all indicator changes colour on the basis of pH of medium. So in titration of NaHCO3, KHCO3 against acid phenolphthalein cannot be used.

So we can write with phenolphthalein, if total meq of Na2CO3 = 1 then ½ meq of Na2CO3 = meq of HCl with methyl orange, meq of Na2CO3 = meq of HCl
**n-factor in non-redox system**

<table>
<thead>
<tr>
<th>Titration</th>
<th>Indicator</th>
<th>pH Range</th>
<th>n factor</th>
</tr>
</thead>
<tbody>
<tr>
<td>Na₂CO₃</td>
<td>Phenolphthalein</td>
<td>8.3 – 10</td>
<td>1</td>
</tr>
<tr>
<td>K₂CO₃</td>
<td>Methyl orange</td>
<td>3.1 – 4.4</td>
<td>2</td>
</tr>
</tbody>
</table>

*a product formed in reaction.*

**Note:** When we carry out dilution of solution, meq eq, milli mole or mole of substance does not change because they represent amount of substance, however molar concentration may change.

**Solubilities of some important salt’s:**

1. **Chloride:**
   - AgCl - White ppt.
   - Hg₂Cl₂ - White ppt. All other chlorides are soluble in water.
   - PbCl₂ - White ppt.
   - CuCl - Insoluble ppt.
   - BiOCl - White ppt.
   - SbOCl - White ppt.
   - Hg₂OCl₂ - White ppt.

2. **Bromide:** AgBr - Pale yellow ppt.
   - PbBr₂ - White ppt. All other bromides are soluble in water.
   - Hg₂Br₂ - White ppt.
   - CuBr - White ppt.

3. **Iodide:** AgI - Yellow ppt.
   - Pbi₂ - Yellow ppt.
   - Hg₂I₂ - Green ppt.
   - Hgl₂ - Red ppt.
   - CuI - White ppt.
   - BI₃ - Black ppt.

4. **Some important oxides and hydroxides:**
   - Ag₂O - Brown ppt.
   - Pb(OH)₂ - White ppt.
   - HgO - Yellow ppt.
   - CuO - Red ppt.
   - Cu(OH)₂ - Blue ppt.
   - Cd(OH)₂ - White ppt.
   - Fe(OH)₂ - White ppt.
   - Fe(OH)₃ - Red ppt.
   - Sn(OH)₂ - White ppt.
   - Sn(OH)₄ - White ppt.
   - Al(OH)₃ - White gelatinous
   - Cr(OH)₃ - Grey-Green
   - Co(OH)₂ - Pink
   - Co(OH)₃ - Brownish black
   - Ni(OH)₂ - Green
Ni(OH)₃ - Black  
Mn(OH)₂ - White  
MnO(OH)₂ - Brown  
Zn(OH)₂ - White  
Mg(OH)₂ - White

**Carbonates**: Except Alkali metals and NH₄⁺, all other carbonates are insoluble.

- Ag₂CO₃ -> White ppt. -> Ag₂O + CO₂
- 3HgO.HgCO₃ -> basic murcuric carbonate White ppt.
- CuCO₃ -> Green ppt.
- CaCO₃ -> White ppt.

**Sulphites** (SO₃⁻²): Except Alkali metal and Ammonium, all other sulphite are generally insoluble.

- Examples: Ag₂SO₃, PbSO₃
- BaSO₃ -> White ppt.
- CaSO₃

**Thiosulphates**: Mostly soluble except

- Ag₂S₂O₃ -> White ppt. [Ag(S₂O₃)₂]⁻ - soluble
- PbS₂O₃ -> White ppt.
- BaS₂O₃ -> White ppt.

**Thiocyanate (SCN⁻)**: Hg(SCN)₂ - White ppt. (Pharaoh’s serpent)

- Ag(SCN) - White ppt.
- Cu(SCN)₂ - Black ppt.
- Cu(SCN) - White ppt.
- Fe(SCN)₃ - Red complex.
- [Co(SCN)₄]²⁻ - Blue complex
- Co[Hg(SCN)₄] - Blue ppt.

**Cynoïdes (CN⁻)**: Except Alkali metal Alkaline earth metal cyanides are soluble in water.

- Hg(CN)₂ - soluble in water in undissociated form
- Ag(CN) - White ppt. [Ag(CN)₂]⁻ - soluble
- Pb(CN)₂ - White ppt.
- Fe(CN)₃ - Brown ppt. [Fe(CN)₆]³⁻ - soluble
- Co(CN)₂ - Brown ppt. [Co(CN)₆]⁴⁻ - soluble
- Ni(CN)₂ - Green [Ni(CN)₄]²⁻ - soluble

**Sulphides**: Except Alkali metals and ammonium salt’s all other sulphides are insoluble. Some insoluble sulphides with unusual colour are

- CdS -> Yellow
- MnS -> Pink
- ZnS -> White
- SnS -> Brown
- SnS₂ -> Yellow
- As₂S₃ -> Yellow
- Sb₂S₃ -> Orange

**Chromates**: Ag₂CrO₄ -> Red ppt.  
PbCrO₄ -> Yellow ppt.  
BaCrO₄ -> Yellow ppt.  
FeCrO₄ -> Green ppt.

Dichromates are generally soluble.
Phosphates: Are generally insoluble:
- $\text{Ag}_3\text{PO}_4$ -> Yellow ppt.
- $\text{AlPO}_4$ -> Yellow ppt.
- $\text{ZrO(HPO}_4\text{)}$ -> White ppt.
- $\text{Mg(NH}_4\text{)PO}_4$ -> White ppt.
- (NH4)3[P Mo12O40] -> Canary yellow ppt.

Phosphite ($\text{HPO}_4^{2-}$): Except Alkali metals all other phosphites are insoluble

Hypo phosphite: All hypophosphites are soluble in water.

- All Acetate are soluble except $\text{Ag(CH}_3\text{COO)}$.
- All formates are soluble except $\text{Ag(HCOO)}$.
- Tatarate, Citrate, Salicylate, Succinate of Silver-are all insoluble white ppt.

Some Important ppt.: KH (Tartarate) -> White ppt.
- $\text{NH}_4\text{H(Tartarate)}$ -> White ppt.
- $\text{K}_2[\text{PtCl}_6]$ -> White ppt.
- $\text{K}_3[\text{Co(NO}_2\text{)_6}]$ -> Yellow ppt.
- (NH4)3[Co(NO2)6] -> Yellow ppt.
- (NH4)2[PtCl6] -> Yellow ppt.

Heating Effect

Heating effect of carbonate & bicarbonate salts
- All carbonates except (Na, K, Rb, Cs) decompose on heating giving CO2
  - $\text{Li}_2\text{CO}_3$ -> $\text{Li}_2\text{O} + \text{CO}_2$
  - $\text{MCO}_3$ -> MO + CO2 $[\text{M} = \text{Be, Mg, Ca, Sr, Ba}]$
  - $\text{Cu(OH}_2\text{.CuCO}_3$ -> 2CuO + CO2 + H2O

Basic Cu(II) carbonate

- White $\text{ZnCO}_3$ -> ZnO + CO2;  
  - ZnO is white(cold) Yellow(hot)
- $\text{PbCO}_3$ -> Yellow PbO + CO2
- $\text{Ag}_2\text{CO}_3$ -> 2Ag + CO2 + $\frac{1}{2}$ O2  
  - Ag is Black
- $\text{HgCO}_3$ -> Hg + $\frac{1}{2}$ O2 + CO2
- (NH4)2CO3 -> 2NH3 + H2O + CO2
- All bicarbonates decompose to give carbonates and CO2. eg.
  - 2NaHCO3 -> Na2CO3 + CO2 + H2O
General reaction:

\[ 2\text{HCO}_3^- \rightarrow \text{CO}_3^{2-} + \text{H}_2\text{O} + \text{CO}_2 \]

**Heating effect of ammonium salts**

\[
\begin{align*}
\text{NH}_4\text{NO}_2 & \rightarrow \text{N}_2 + 2\text{H}_2\text{O}; \quad \text{NH}_4\text{NO}_3 \rightarrow \text{N}_2\text{O} + 2\text{H}_2\text{O} \\
(\text{NH}_4)\text{Cr}_2\text{O}_7 & \rightarrow \text{N}_2 + \text{Cr}_2\text{O}_3 + 4\text{H}_2\text{O} \\
2\text{NH}_4\text{ClO}_4 & \rightarrow \text{N}_2 + \text{Cl}_2 + 2\text{O}_2 + 4\text{H}_2\text{O} \\
2\text{NH}_4\text{IO}_3 & \rightarrow \text{N}_2 + \text{I}_2 + \text{O}_2 + 4\text{H}_2\text{O} \\
\text{[If anionic part is oxidising in nature, then N}_2\text{ will be the product (sometimes N}_2\text{O).]} \]
\[
\begin{align*}
(\text{NH}_4)_2\text{HPO}_4 & \rightarrow \text{HPO}_3 + \text{H}_2\text{O} + 2\text{NH}_3 \\
(\text{NH}_4)_2\text{SO}_4 & \rightarrow \text{NH}_3 + \text{H}_2\text{SO}_4 \\
2(\text{NH}_3)_3\text{PO}_4 & \rightarrow 2\text{NH}_3 + \text{P}_2\text{O}_5 + 3\text{H}_2\text{O} \\
(\text{NH}_4)_2\text{CO}_3 & \rightarrow 2\text{NH}_3 + \text{H}_2\text{O} + \text{CO}_2 \\
\text{[If anionic part weakly oxidising or non-oxidising in nature then NH}_3\text{ will be the product.]} \\
\end{align*}
\]

**Heating effect of nitrate salts**

\[
\begin{align*}
\text{MNO}_3 & \rightarrow \text{KNO}_2 + \frac{1}{2} \text{O}_2 \quad [\text{M} = \text{Na, K, Rb, Cs}] \\
2\text{LiNO}_3 & \rightarrow \text{Li}_2\text{O} + 2\text{NO}_2 + \frac{1}{2} \text{O}_2 \\
2\text{M(NO}_3)_2 & \rightarrow 2\text{MO} + 4\text{NO}_2 + \text{O}_2 \quad [\text{M} = \text{all bivalent metal's ions eg. Zn}^{+2}, \text{Mg}^{+2}, \text{Sr}^{+2}, \text{Ca}^{+2}, \text{Ba}^{+2}, \text{Cu}^{+2}, \text{Pb}^{+2}] \\
\text{Hg(NO}_3)_2 & \rightarrow \text{Hg} + 2\text{NO}_2 + \text{O}_2 \\
2\text{AgNO}_3 & \rightarrow 2\text{Ag} + 2\text{NO}_2 + \text{O}_2
\end{align*}
\]

**Heating effect of Halides salts**

\[
\begin{align*}
2\text{FeCl}_3 & \rightarrow 2\text{FeCl}_2 + \text{Cl}_2 \\
\text{AuCl}_3 & \rightarrow \text{AuCl} + \text{Cl}_2 \\
\text{HgCl}_2 & \rightarrow \text{HgCl}_2 + \text{Hg} \\
\text{NH}_4\text{Cl} & \rightarrow \text{NH}_3 + \text{HCl} \\
\text{PbX}_4 & \rightarrow \text{PbX}_2 + \text{X}_2 \quad [\text{X} = \text{Cl, Br, SCN}^-]
\end{align*}
\]

**Heating effect of hydrated chloride salts**

\[
\begin{align*}
\text{MgCl}_2 \cdot 6\text{H}_2\text{O} & \rightarrow \text{MgO} + 2\text{HCl} + 5\text{H}_2\text{O} \\
2\text{FeCl}_3 \cdot 6\text{H}_2\text{O} & \rightarrow 2\text{FeCl}_2 + 6\text{HCl} + 9\text{H}_2\text{O} \\
2\text{AlCl}_3 \cdot 6\text{H}_2\text{O} & \rightarrow 2\text{Al}_2\text{O}_3 + 6\text{HCl} + 9\text{H}_2\text{O}
\end{align*}
\]

\[
\begin{align*}
\text{Pink CoCl}_2\cdot 6\text{H}_2\text{O} \quad (50^\circ \text{C}, \text{loss of 2H}_2\text{O}) & \rightarrow \text{Pink CoCl}_2\cdot 4\text{H}_2\text{O} \quad (58^\circ \text{C}, \text{loss of 2H}_2\text{O}) \rightarrow \text{Red Violet CoCl}_2\cdot 2\text{H}_2\text{O} \quad (140^\circ \text{C}, \text{loss of 2H}_2\text{O}) \rightarrow \text{CoCl}_2 \quad (\text{Blue})
\end{align*}
\]
Heating effect of hydrated Sulphate salts

\[
\begin{align*}
\text{CuSO}_4 \cdot 5\text{H}_2\text{O} \xrightarrow{100^\circ\text{C}} & \text{CuSO}_4 \cdot \text{H}_2\text{O} \xrightarrow{220^\circ\text{C}} \text{CuSO}_4 \xrightarrow{>800^\circ\text{C}} \text{CuO} + \text{SO}_2 + \frac{1}{2}\text{O}_2 \\
\text{FeSO}_4 \cdot 7\text{H}_2\text{O} \xrightarrow{300^\circ\text{C}} & \text{FeSO}_4 \xrightarrow{\Delta} \text{Fe}_2\text{SO}_3 + \text{SO}_2 + \text{SO}_3 \quad \text{Very Important} \\
\text{Fe}_2(\text{SO}_4)_3 \xrightarrow{\Delta} & \text{Fe}_2\text{O}_3 + 3\text{SO}_3 \\
\text{MgSO}_4 \cdot 7\text{H}_2\text{O} \xrightarrow{\Delta} & \text{MgSO}_4 \downarrow \quad \text{[Same as ZnSO}_4\text{]} \\
\text{CaSO}_4 \cdot 2\text{H}_2\text{O} \xrightarrow{120^\circ\text{C}} & (\text{CaSO}_4 \cdot \frac{1}{2}\text{H}_2\text{O}) + 1\frac{1}{2}\text{H}_2\text{O} \\
\text{CaSO}_4 \cdot 2\text{H}_2\text{O} \xrightarrow{\Delta} & \text{CaSO}_4 + \frac{1}{2}\text{H}_2\text{O} \\
\text{Na}_2\text{S}_2\text{O}_3 \cdot 5\text{H}_2\text{O} \xrightarrow{220^\circ\text{C}} & \text{Na}_2\text{S}_2\text{O}_3 + 5\text{H}_2\text{O} \\
\text{ZnSO}_4 \cdot 7\text{H}_2\text{O} \xrightarrow{70^\circ\text{C}} & \text{ZnSO}_4 \cdot 6\text{H}_2\text{O} \xrightarrow{70^\circ-200^\circ\text{C}} \text{ZnSO}_4 \cdot \text{H}_2\text{O} \xrightarrow{>200^\circ\text{C}} \text{ZnSO}_4 \xrightarrow{\Delta >800^\circ\text{C}} \text{ZnO} + \text{SO}_2 + \frac{1}{2}\text{O}_2 \\
2\text{NaHSO}_3 & \xrightarrow{\Delta} \text{Na}_2\text{S}_2\text{O}_5 + \text{H}_2\text{O} \\
2\text{NaHSO}_4 & \xrightarrow{\Delta} \text{Na}_2\text{SO}_4 + \text{H}_2\text{O} + \text{SO}_3
\end{align*}
\]
Heating effect of Oxide salts

\[ 2\text{Ag}_2\text{O} \xrightarrow{\Delta \text{300}^\circ\text{C}} 4\text{Ag} + \text{O}_2 \]

\[ \text{PbO}_2 \xrightarrow{\Delta} \text{PbO} + \frac{1}{2} \text{O}_2 \]

\[ 3\text{MnO}_2 \xrightarrow{900^\circ\text{C}} \text{Mn}_3\text{O}_4 + \text{O}_2 \]

\[ 2\text{CrO}_5 \rightarrow \text{Cr}_2\text{O}_3 + \frac{5}{2} \text{O}_2 \]

I$_2$O$_5$ $\rightarrow$ I$_2$ + $\frac{5}{2}$ O$_2$

Heating effect of dichromate & chromate salts

\[ (\text{NH}_4)_2\text{Cr}_2\text{O}_7 \xrightarrow{\Delta} \text{N}_2 + \text{Cr}_2\text{O}_3 + 4\text{H}_2\text{O} \]

\[ \text{K}_2\text{Cr}_2\text{O}_7 \xrightarrow{\Delta} 2\text{K}_2\text{CrO}_4 + \text{Cr}_2\text{O}_3 + \frac{7}{2} \text{O}_2 \]

Heating effect of phosphate salts

\[ \text{Na}_2\text{HPO}_4 \xrightarrow{\Delta} \text{H}_2\text{O} + \text{NaPO}_3 \]

\[ \text{Na}_3\text{PO}_4 \xrightarrow{\Delta} \text{No effect} \]

\[ \text{Na(NH}_4\text{)HPO}_4 \cdot 4\text{H}_2\text{O} \rightarrow \text{NaNH}_4\text{HPO}_4 \rightarrow (\text{High Temp}) \text{NaPO}_3 + \text{NH}_3 + \text{H}_2\text{O} \]

Heating effects of acetate, formate, oxalate salts

\[ \text{CH}_3\text{CO}_2\text{K} \xrightarrow{\Delta} \text{K}_2\text{CO}_3 + \text{CH}_3\text{COCH}_3 \]

\[ \text{Na}_2\text{C}_2\text{O}_4 \xrightarrow{\Delta} \text{Na}_2\text{CO}_3 + \text{CO} \]

\[ \text{Pb(OAc)}_2 \xrightarrow{\Delta} \text{PbO} + \text{CO}_2 + \text{CH}_3\text{COCH}_3 \]
SnC2O4 $\xrightarrow{\Delta} \text{SnO + CO2 + CO}$

Mg(OAc)2 $\xrightarrow{\Delta} \text{MgO + CO2 + CH3COCH3}$

FeC2O4 $\rightarrow \text{FeO + CO + CO2}$

Be(OAc)2 $\rightarrow \text{BeO + CO2 + CH3COCH3}$

Ag2C2O4 $\rightarrow 2\text{Ag + 2CO2}$

Ca(OAc)2 $\rightarrow \text{CaCO3 + CH3COCH3}$

HgC2O4 $\rightarrow \text{Hg + 2CO2}$

Ba(OAc)2 $\rightarrow \text{BaCO3 + CH3COCH3}$

(HCOO)2Hg $\rightarrow \text{HCOOH + Hg + CO2}$

2HCOOAg $\rightarrow \text{HCOOH + 2Ag + CO2}$

$\text{HCO}_2\text{Na} \xrightarrow{350^\circ C} \text{Na}_2\text{C}_2\text{O}_4 + \text{H}_2 \uparrow$

**Heating effect of Acids**

\[2\text{HNO}_3 \xrightarrow{\Delta} \text{H}_2\text{O} + 2\text{NO}_2 + \text{S}_2\text{O}_2\]

\[\text{H}_2\text{SO}_4 \xrightarrow{444^\circ C} \text{H}_2\text{O} + \text{SO}_3:\]

\[3\text{H}_2\text{SO}_3 \rightarrow 2\text{H}_2\text{SO}_4 + \text{S} \downarrow + \text{H}_2\text{O}\]

\[3\text{HNO}_2 \rightarrow \text{HNO}_3 + 2\text{NO} + \text{H}_2\text{O}\]

\[\text{HClO}_3 \rightarrow \text{HClO}_4 + \text{ClO}_2 + \text{H}_2\text{O}\]

\[3\text{HClO} \rightarrow 2\text{HCl} + \text{HClO}_3\]

\[4\text{H}_3\text{PO}_3 \xrightarrow{200^\circ C} 3\text{H}_3\text{PO}_4 + \text{PH}_3\]

\[2\text{H}_3\text{PO}_2 \rightarrow \text{H}_3\text{PO}_4 + \text{PH}_3\]

\[2\text{NaH}_2\text{PO}_2 \rightarrow \text{Na}_2\text{HPO}_4 + \text{PH}_3\]

\[\text{H}_3\text{PO}_4 \xrightarrow{220^\circ C} \text{H}_4\text{P}_2\text{O}_7 \xrightarrow{320^\circ C} 4\text{HPO}_3 \xrightarrow{320^\circ C} 2\text{P}_2\text{O}_5 + 2\text{H}_2\text{O}\]

\[\text{H}_2\text{SO}_4 \xrightarrow{>800^\circ C} \text{H}_2\text{O} + \text{SO}_2 + \frac{1}{2}\text{O}_2\]

Undergoes disproportionation reaction
Some reactions of important oxidising agents

(I) Potassium dichromate (K₂Cr₂O₇):
Cr₂O₇²⁻ ion takes electrons in the acidic medium and is reduced to Cr³⁺ ion. Thus Cr₂O₇⁻² acts as an oxidising agent in acidic medium.

\[
\text{K}_2\text{Cr}_2\text{O}_7 + 4\text{H}_2\text{SO}_4 (\text{dil.}) \rightarrow \text{K}_2\text{SO}_4 + \text{Cr}_2(\text{SO}_4)_3 + 4\text{H}_2\text{O} + 3\text{O}
\]

or

\[
\text{Cr}_2\text{O}_7^{2-} + 14\text{H}^+ + 6\text{e}^- \rightarrow 2\text{Cr}^{3+} + 7\text{H}_2\text{O}
\]

(1) \(\text{K}_2\text{Cr}_2\text{O}_7 + 7\text{H}_2\text{SO}_4 + 6\text{KI} \rightarrow 4\text{K}_2\text{SO}_4 + \text{Cr}_2(\text{SO}_4)_3 + 7\text{H}_2\text{O} + 3\text{I}_2\)

or

\(\text{Cr}_2\text{O}_7^{2-} + 14\text{H}^+ + 6\text{I}^- \rightarrow 2\text{Cr}^{3+} + 7\text{H}_2\text{O} + 3\text{I}_2\)

(2) \(\text{K}_2\text{Cr}_2\text{O}_7 + 7\text{H}_2\text{SO}_4 + 6\text{FeSO}_4 \rightarrow \text{K}_2\text{SO}_4 + \text{Cr}_2(\text{SO}_4)_3 + 7\text{H}_2\text{O} + 3\text{Fe}_2(\text{SO}_4)_3\)

or

\(\text{Cr}_2\text{O}_7^{2-} + 14\text{H}^+ + 6\text{Fe}^{2+} \rightarrow 2\text{Cr}^{3+} + 7\text{H}_2\text{O} + 6\text{Fe}^{3+}\)

(3) \(\text{K}_2\text{Cr}_2\text{O}_7 + 4\text{H}_2\text{SO}_4 + 3\text{H}_2\text{S} \rightarrow \text{K}_2\text{SO}_4 + \text{Cr}_2(\text{SO}_4)_3 + 7\text{H}_2\text{O} + 3\text{S}\)

or

\(\text{Cr}_2\text{O}_7^{2-} + 8\text{H}^+ + 3\text{H}_2\text{S} \rightarrow 2\text{Cr}^{3+} + 7\text{H}_2\text{O} + 3\text{S}\)

(4) \(\text{K}_2\text{Cr}_2\text{O}_7 + \text{H}_2\text{SO}_4 + 3\text{SO}_2 \rightarrow \text{K}_2\text{SO}_4 + \text{Cr}_2(\text{SO}_4)_3 + \text{H}_2\text{O}\)

or

\(\text{Cr}_2\text{O}_7^{2-} + 2\text{H}^+ + 3\text{SO}_2 \rightarrow 2\text{Cr}^{3+} + 3\text{ MnO}_4^- + \text{H}_2\text{O}\)

(5) \(\text{K}_2\text{Cr}_2\text{O}_7 + 4\text{H}_2\text{SO}_4 + 3\text{Na}_2\text{SO}_3 \rightarrow \text{K}_2\text{SO}_4 + \text{Cr}_2(\text{SO}_4)_3 + 3\text{Na}_2\text{SO}_3 + 4\text{H}_2\text{O}\)

or

\(\text{Cr}_2\text{O}_7^{2-} + 8\text{H}^+ + 3\text{SO}_3^{2-} \rightarrow 2\text{Cr}^{3+} + 3\text{SO}_4^{2-} + 4\text{H}_2\text{O}\)

(6) \(\text{K}_2\text{Cr}_2\text{O}_7 + 14\text{HCl} \rightarrow 2\text{KCl} + 2\text{CrCl}_3 + 7\text{H}_2\text{O} + 3\text{ Cl}_2\)

or

\(\text{Cr}_2\text{O}_7^{2-} + 14\text{H}^+ + 6\text{Cl}^- \rightarrow 2\text{Cr}^{3+} + 7\text{H}_2\text{O} + 3\text{ Cl}_2\)

(7) \(\text{K}_2\text{Cr}_2\text{O}_7 + \text{H}_2\text{SO}_4 + 4\text{H}_2\text{O}_2 \rightarrow \text{ether} \rightarrow \text{K}_2\text{SO}_4 + 2\text{CrO}_5 + 5\text{H}_2\text{O}\)

or

\(\text{Cr}_2\text{O}_7^{2-} + 2\text{H}^+ + 4\text{H}_2\text{O}_2 \rightarrow \text{ether} \rightarrow 2\text{CrO}_5 + 5\text{H}_2\text{O}\)

(II) Manganese dioxide (MnO₂):

In presence of excess of H⁺ ions, MnO₂ acts as a strong oxidising agent. In showing this behaviour Mn⁴⁺ changes to Mn²⁺ ion.

\[
\text{MnO}_2 + 4\text{H}^+ + 4\text{e}^- \rightarrow \text{Mn}^{2+} + 2\text{H}_2\text{O}
\]

(1) \(\text{MnO}_2 + 4\text{H}^+ + \text{C}_2\text{O}_4^{2-} \rightarrow \text{Mn}^{2+} + 2\text{H}_2\text{O} + 2\text{CO}_2\)

(2) \(\text{MnO}_2 + 4\text{H}^+ + 2\text{Fe}^{2+} \rightarrow \text{Mn}^{2+} + 2\text{H}_2\text{O} + 2\text{Fe}^{3+}\)

(3) \(\text{MnO}_2 + 4\text{H}^+ + 2\text{Cl}^- \rightarrow \text{Mn}^{2+} + 2\text{H}_2\text{O} + \text{Cl}_2\)
(III) Potassium permanganate (KMnO₄):

(A) In acidic medium: The reduction of MnO₄⁻ ion into Mn²⁺ ion can be represented by the following ionic equation:
(8) **Halogen acid, HX (X = -1)** to the corresponding halogen, X₂(X = 0) (X⁻ → X₂)  
2KMnO₄ + 3H₂SO₄ + 10HCl → K₂SO₄ + 2MnSO₄ + 8H₂O + 5Cl₂  
or  
2 MnO₄⁻ + 16H⁺ + 10Cl⁻ → 2 Mn²⁺ + 3H₂O + 5Cl₂

(9) **H₂O₂ (O = -1) to O₂ (O = 0)**  
or  
2KMnO₄ + 3H₂SO₄ + 5H₂O₂ → K₂SO₄ + 2MnSO₄ + 8H₂O + 5 O₂  
2 MnO₄⁻ + 6H⁺ + 5H₂O₂ → 2 Mn²⁺ + 8H₂O + 5H₂O

(10) **SO₂ (S = +4) to H₂SO₄ (S = +6)** (SO₂ → SO₄²⁻)  
or  
2KMnO₄ + 2 H₂O + 5SO₂ → K₂SO₄ + 2 MnSO₄ + 2 H₂SO₄  
2 MnO₄⁻ + 2 H₂O + 5SO₂ → 2 Mn²⁺ + 5 SO₄²⁻

(11) **Hydrazine, N₂H₄ (N = -2)** to N₂ (N = 0)  
or  
4KMnO₄ + 6 H₂SO₄ + 5 N₂H₄ → 2 K₂SO₄ + 4MnSO₄ + 16H₂O + 5N₂  
or  
4 MnO₄⁻+12 H⁺ + 5N₂H₄ → 2Mn²⁺ + 16H₂O + 5N₂

(12) **Hydrazoic acid, HN₃ (N = -1/3)** to N₂ (N = 0)  
or  
2 KMnO₄ + 3 H₂SO₄ + 10HN₃ → K₂SO₄ + 2MnSO₄ + 8H₂O + 15N₂  
or  
2 MnO₄⁻ + 6H⁺ + 5HN₃ → 2Mn²⁺ + 8H₂O + 15N₂

(13) **Nitric oxide, NO (N = +2)** to HNO₃ (N = +5)  
or  
6KMnO₄ + 9H₂SO₄ + 10NO → 3K₂SO₄ + 6MnSO₄ + 4H₂O + 10HNO₃  
or  
3 MnO₄⁻ + 9H⁺ + 5NO → 3Mn²⁺ + 2H₂O + 5HNO₃

(14) **Potassium ferrocyanide, K₃[Fe(CN)₆]** to potassium ferricyanide, K₃[Fe(CN)₆]⁻  
or  
2KMnO₄ + 3H₂SO₄ + 10K₃[Fe(CN)₆] → K₂SO₄ + 2MnSO₄ + 2H₂O  
+ 10K₃[Fe(CN)₆]⁻ + 10KOH  
or  
MnO₄⁻ + 8H⁺ + 5[Fe(CN)₆]⁴⁻ → Mn²⁺ + 4H₂O + 5[Fe(CN)₆]⁻

(15) **Sodium thiosulphate, Na₂S₂O₃ (S = +3)** to sodium dithionate, Na₂S₂O₆ (S = +5)  
or  
6KMnO₄ + 9H₂SO₄ + 5Na₂S₂O₃ → 3K₂SO₄ + 6MnSO₄ + 9H₂O + 5Na₂S₂O₆  
or  
6 MnO₄⁻ + 18H⁺ + 5S₂O₃²⁻ → 6Mn²⁺ + 9H₂O + 5S₂O₆²⁻
(B) In alkaline medium: In alkaline solution MnO₄⁻ ion is reduced to colourless & insoluble MnO₂ according to the following equations:

\[
2 \text{MnO}_4^- + \text{H}_2\text{O} \overset{\text{alkali}}{\longrightarrow} 2\text{MnO}_2 + 2\text{OH}^- + 3\text{O}_2
\]

or

\[
\text{MnO}_4^- + 2\text{H}_2\text{O} + 3\text{e}^- \longrightarrow \text{MnO}_2 + 4\text{OH}^-;
\]

(I) Iodides (I⁻) to iodates (IO₃⁻)

\[
2\text{KMnO}_4 + \text{H}_2\text{O} + \text{KI} \longrightarrow 2\text{MnO}_2 + 2\text{KOH} + \text{KIO}_3
\]

or

\[
2\text{MnO}_4^- + \text{H}_2\text{O} + \text{I}^- \longrightarrow 2\text{MnO}_2 + 2\text{OH}^- + \text{IO}_3^-;
\]

(2) NH₃ (N = -3) to N₂ (N = 0)

\[
2\text{KMnO}_4 + 2\text{NH}_3 \longrightarrow 2\text{MnO}_2 + 2\text{KOH} + \text{N}_2 + 2\text{H}_2\text{O}
\]

or

\[
2\text{MnO}_4^- + 2\text{NH}_3 \longrightarrow 2\text{H}_2\text{O} + 2\text{OH}^- + \text{N}_2 + 2\text{MnO}_2;
\]

(3) Nitrotoluene to nitrobenzoic acid

\[
2\text{KMnO}_4 + \text{C}_6\text{H}_4\text{NO}_2 \longrightarrow 2\text{MnO}_2 + 2\text{KOH} + \text{C}_6\text{H}_4\text{COOH}
\]

Nitrotoluene

\[
\text{Nitrobenzoic acid}
\]

or

\[
2\text{MnO}_4^- + \text{C}_6\text{H}_4\text{NO}_2 \longrightarrow \text{MnO}_2 + \text{C}_6\text{H}_4\text{COOH} + \text{OH}^-;
\]

(4) Ethylene (H₂C = CH₂) to ethylene glycol (HO-CH₂-CH₂-OH)

\[
2\text{KMnO}_4 + 4\text{H}_2\text{O} + 3\text{H}_2\text{C} = \text{CH}_2 \longrightarrow 2\text{MnO}_2 + 2\text{KOH} + 3\text{HO-CH}_2\text{CH}_2\text{-OH}
\]

or

\[
2\text{MnO}_4^- + 4\text{H}_2\text{O} + 3\text{H}_2\text{C} = \text{CH}_2 \longrightarrow 2\text{MnO}_2 + 2\text{OH}^- + 3\text{HO-CH}_2\text{CH}_2\text{-OH};
\]

(C) In neutral medium: In neutral solution, KMnO₄ is directly reduced to MnO₂

\[
2\text{KMnO}_4 + \text{H}_2\text{O} \overset{\text{¼O}}{\longrightarrow} 2\text{KOH} + 2\text{MnO}_2 + 3\text{O}_2
\]

\[
\text{MnO}_4^- + \text{H}_2\text{O} \overset{\text{¼O}}{\longrightarrow} 2\text{OH}^- + 2\text{MnO}_2 + 3\text{O}_2
\]

\[
\text{MnO}_4^- + 2\text{H}_2\text{O} + 3\text{e}^- \overset{\text{¼O}}{\longrightarrow} 2\text{MnO}_2 + 4\text{OH}^-;
\]
Stoichiometry

Many teachers approach Stoichiometry in 2 parts
1 ) Theoretical Stoichiometry
2 ) Experimental Stoichiometric analysis

In Theoretical Stoichiometry we have
1.1 ) Stoichiometric proportions
1.2 ) non-Stoichiometric proportions
1.3 ) Concept of Limiting Reagent

In Experimental Stoichiometric analysis we have
2.1 ) Gravimetric Analysis
2.2 ) Titrimetric Methods
2.3 ) Gas Analysis ( Reactions where Reactants and Products are gaseous )

Titrimetric methods are of ( Reaction in aqueous solution )
2.2.1 ) Acid-Base Reactions
2.2.2 ) Redox Reactions
2.2.3 ) Precipitation Reactions
2.2.4 ) Complex Titrimetric Reactions

Aliquot. A portion of the whole, usually a simple fraction. A portion of a sample withdraw from a volumetric flask with a pipet is called an aliquot.

Analytical concentration. The total number of moles per litre of a solute regardless of any reactions that might occur when the solute dissolves. Used synonymously with formality.

Equivalent. The amount of a substance which furnishes or reacts with 1 mol of H+ (acid-base), 1 mol of electrons (redox), or 1 mol of a univalent cation (precipitation and complex formation).

Equivalent weight. The weight in grams of one equivalent of a substance.
Equivalence point. The point in a titration where the number of equivalents of titrant is the same as the number of equivalents of analyte.
End point. The point in a titration where an indicator changes color.

Formula weight. The number of formula weights of all the atoms in the chemical formula of a substance.

Formality. The number of formula weights of solute per litre of solution; synonymous with analytical concentration.

Indicator. A chemical substance which exhibits different colors in the presence of excess analyte or titrant.

Normality. The number of equivalents of solute per litre of solution.

Primary standard. A substance available in a pure form or state of known purity which is used in standardizing a solution.

Standardization. The process by which the concentration of a solution is accurately ascertained.

Standard solution. A solution whose concentration has been accurately determined.

Titrant. The reagent (a standard solution) which is added from a buret to react with the analyte.

**2008 EAMCET**

Match the following

<table>
<thead>
<tr>
<th>List – I</th>
<th>List – II</th>
</tr>
</thead>
<tbody>
<tr>
<td>A) 10g. CaCO₃ ( \xrightarrow{\text{D \ decomposition}} )</td>
<td>i) 0.224 lit. CO₂</td>
</tr>
<tr>
<td>B) 1.06 g Na₂CO₃ ( \xrightarrow{\text{excess HCl}} )</td>
<td>ii) 4.48 lit. CO₂</td>
</tr>
<tr>
<td>C) 10g. CaCO₃ ( \xrightarrow{\text{excess O₂ \ combustion}} )</td>
<td>iii) 0.448 lit. CO₂</td>
</tr>
<tr>
<td>D) 0.056g. CO ( \xrightarrow{\text{excess O₂ \ combustion}} )</td>
<td>iv) 2.24 lit. CO₂</td>
</tr>
</tbody>
</table>

A B C D | A B C D
| iv i ii iii | 2) v i ii iii |
| iv i iii ii | 4) i iv ii iii |

Ans : ( 1 )

Solution : ( A ) CaCO \( \rightarrow \) CaO + CO
100 gm of CaCO₃ gives 22.4 lit CO₂ at STP \( \rightarrow \) 10 gm of CaCO₃ gives 2.24 lit of CO₂

(B ) Na CO + 2HCl \( \rightarrow \) 2NaCl + H O + CO
106 gm of Na₂CO₃ gives 22.4 lit of CO₂ at STP
1.06 gm of Na₂CO₃ gives 0.224 lit of CO₂

(C ) C+O \( \rightarrow \) CO
12 gm of ‘C’ on combustion gives 22.4 lit of CO₂ at STP 2.4 gm of ‘C’ give 4.4 lit of CO₂

(D ) 2C+O \( \rightarrow \) 2CO
2x 28 gm of CO gives 2 x 22.4 lit of CO₂ at STP \( \rightarrow \) 0.56 gm of CO gives 0.448 lit CO₂
2008 EAMCET

40 grams of a sample of carbon on combustion left 10% of it unreacted. The volume of oxygen required at STP for this combustion reaction is  
1) 22.4 / (2) 67.2 / (3) 11.2 / (4) 44.8 l  
Ans: 2  
Solution: C+O → CO  
Amount of 'C' reacts = 40 - 4 = 36 gm  
According to equation 12 gm of C requires 22.4 lit of O2 -> 36 gm of 'C' requires 3× 22.4 = 67.2 lit of O2

Question :  
In an oxidation reduction reaction, dichromate \( \text{Cr}_2\text{O}_7^{2-} \) ion is reduced to \( \text{Cr}^{3+} \) ion. The equivalent weight of \( \text{K}_2\text{Cr}_2\text{O}_7 \) in this reaction is ?  
1) \( \frac{3}{\text{Molecular weight}} \)  
2) \( \frac{6}{\text{Molecular weight}} \)  
3) \( \frac{1}{\text{Molecular weight}} \)  
4) \( \frac{2}{\text{Molecular weight}} \)  
Answer : (2)

Solution: change in oxidation number of Cr = 6 units  
So \[ \text{Equivalent weight} = \frac{6}{\text{Molecular weight}} \] 

Question : In the redox reaction,  
\[ 2\text{KMnO}_4 + 3\text{H}_2\text{SO}_4 + 5\text{H}_2\text{C}_2\text{O}_4 \rightarrow \text{K}_2\text{SO}_4 + 2\text{MnSO}_4 + 8\text{H}_2\text{O} + 10\text{CO}_2 \]  
the volume of 0.1 M. \text{KMnO}_4 required to oxidises 25mL of 0.25 M \text{H}_2\text{C}_2\text{O}_4 solution  
1) 25 L  
2) 125 mL  
3) 25 mL  
4) 1.25 L  
Ans : 3  
Solution :  
\[ \frac{M_1v_1}{n_1} (\text{KMnO}_4) = \frac{M_2v_2}{n_2} (\text{H}_2\text{C}_2\text{O}_4) \]  
\[ 0.1 \times v_1 = 0.25 \times 25 \]  
\[ \frac{2}{5} \]  
\[ 0.5 \times 25 \]  
\[ 0.5 \]  
\[ 25 \text{ml} \]
2005 EAMCET
'X' grams of calcium carbonate was completely burnt in air. The weight of the solid residue formed is 28 g. What is the value of 'X' (in grams)
1) 44  2) 200  3) 150  4) 50
Ans: 4
Solution:
\[
\text{CaCO}_3 \rightarrow \text{CaO} + \text{CO}_2
\]
56 gm of CaO is formed from 100 gm of CaCO₃
28 gm of CaO is formed from \( \frac{28 \times 100}{56} = 50 \text{gm} \)

2004 EAMCET
An organic compound containing C and H has 92.3% of carbon. Its empirical formula is?
1) CH  2) CH₃  3) CH₂  4) CH₄
Ans: 1
Solution:
Element Atomic Ratio = \( \frac{\%}{\text{Atomic Weight}} \)
Carbon Ratio = \( \frac{92.3}{12} = 7.7 \)
Hydrogen Ratio = \( \frac{7.7}{1} = 7.7 \)
The Simplest Ratio of these is 1 : 1
So the empirical Formula is CH

2003 EAMCET
Question: What is the volume (in litres) of oxygen required at S.T.P. to completely convert 1.5 moles of sulphur into sulphur dioxide?
1) 11.2  2) 22.4  3) 33.6  4) 44.8
Ans: 3
Solution:
\[
\text{S} + \frac{1}{2}\text{O}_2 \rightarrow \text{SO}_2
\]
1 mole of S required \( \rightarrow \) 22.4 lit of \( \text{O}_2 \) at STP
1.5 mole of S is required \( \rightarrow \) ?
\[
\frac{1.5 \times 22.4}{1} = 33.6 \text{lit}
\]

2002 EAMCET
'X' litres of carbon monoxide is present at STP. It is completely oxidized to CO₂. The volume of CO₂ formed is 11.207 litres at STP. What is the value of 'X' in litres?
1) 22.414  2) 11.207  3) 5.6035  4) 44.828
Ans: 2
Solutions:
\[
\text{CO} + \frac{1}{2}\text{O}_2 \rightarrow \text{CO}_2
\]
According to equation 22.4 lit of CO₂ is formed from \( \rightarrow \) 22.4 lit of CO
11.207 lit of CO₂ is formed from \[ \text{How much?} \]
\[
= \frac{11.207 \times 22.4}{22.4} = 11.207 \text{ lit of CO} = X
\]

### 2002 EAMCET

**Question**: 4gms of a hydrocarbon on complete combustion give 12.571 gms of CO₂ and 5.143 gms of water. What is the empirical formula of the Hydrocarbon?

1) CH  
2) CH₂  
3) CH₃  
4) C₂H₃

**Ans**: 2

**Solution**:

Carbon Atomic Ratio = \( \frac{\% \text{C}}{\text{Atomic Weight}} = \frac{85.71}{12} = 7.14 \)

Hydrogen Atomic Ratio = \( \frac{\% \text{H}}{\text{Atomic Weight}} = \frac{14.28}{1} = 14.28 \)

So Empirical Formula is (1 : 2) CH₂

### 2002 EAMCET

**Question**: What is the minimum quantity (in grams) of Methyl iodide required for preparing 1 mole of ethane by Wurtz reaction? (Atomic weight of iodine = 127)

1) 142  
2) 568  
3) 326  
4) 284

**Ans**: 4

**Solution**:

\[ 2\text{CH}_3\text{I} + 2\text{Na} \xrightarrow{\text{dry ether}} \text{C}_2\text{H}_6 + 2\text{NaI} \]

According to equation 1 mole of \( \text{C}_2\text{H}_6 \) is formed from 2 mole of \( \text{CH}_3\text{I} \)

\[ \therefore \text{Wt of 2 moles of CH}_3\text{I} = 2 \times 142 = 284\text{g} \]

### 2002 EAMCET

**Question**: What is the volume (in litres) of oxygen at STP required for complete combustion of 32 gms of CH₄?

1) 44.8  
2) 89.6  
3) 22.4  
4) 179.2

**Ans**: 2

\[ \text{CH}_4 + 2\text{O}_2 \longrightarrow \text{CO}_2 + 2\text{H}_2\text{O} \]

16 gm of CH₄ is required \( \longrightarrow \) 2 \times 22.4 lit of O₂ at STP

32 gm of CH₄ is required \( \longrightarrow \) ?

\[ \text{lit of O}_2 = \frac{32 \times 2}{16} \times 22.4 = 44.8 \]
2002 EAMCET

What are the oxidation nos. of nitrogen in NH4NO3?

1) +3, -5  2) -3, +5  3) +3, -6  4) +2, +2

Ans: 2

\[ \text{NH}_4\text{NO}_3 \rightarrow \text{NH}_4^+ + \text{NO}_3^- \]

In \( \text{NH}_4^+ \) oxidation state of N is -3

\[ x + 4 = +1 \quad x = -3 \]

In \( \text{NO}_3^- \) oxidation state of N is +5

\[ x - 6 = -1 \quad x = +5 \]

---

2002 EAMCET

How many litres of oxygen (at STP) are required for complete combustion of 39 gms of liquid Benzene?

1) 84  2) 22.4  3) 42  4) 11.2

Ans: 1

\[ \text{C}_6\text{H}_6 + \frac{15}{2}\text{O}_2 \rightarrow 6\text{CO}_2 + 3\text{H}_2\text{O} \]

78 gm of \( \text{C}_6\text{H}_6 \) requires \( \frac{15}{2} \times 22.4 \text{ lit of } \text{O}_2 \) at STP

39 gm of \( \text{C}_6\text{H}_6 \) requires \( ? \)

\[ 39 \times \frac{15}{2} \times \frac{1 \times 22.4}{78} = 84 \]

---

2005 EAMCET

Four grams of hydro carbon (C\( \text{xH}_y \)) on complete combustion gave 12 grams of CO\(_2\). What is the empirical formula of the hydro carbon?

1) CH\(_3\)  2) C\(_4\)H\(_9\)  3) CH  4) C\(_3\)H\(_8\)

Ans: 4

\[ \text{C} + 5\text{O} \rightarrow 3\text{CO} + 4\text{H}_2\text{O} \]

44gm of C\(_3\)H\(_8\) gives 3\times44 gm of CO\(_2\)

So 4gm of C\(_3\)H\(_8\) gives 12 gm of CO\(_2\)

---

2004 EAMCET

10 grams of CaCO\(_3\) is completely decomposed to \( x \) and CaO. \( x \) is passed into an aqueous solution containing 0.1 mole of sodium carbonate. What is the number of moles of sodium bicarbonate formed?

(mol. wts : CaCO\(_3\) = 100; NaCO\(_3\) = 106; NaHCO\(_3\) = 84)

1) 0.2  2) 0.1  3) 0.01  4) 10

Ans: 1

Solution:

\[ \text{CaCO}_3 \rightarrow \text{CaO} + \text{CO}_2 \]

100 gm \( \text{CaCO}_3 \) gives 1 mole of CO\(_2\) -> 10 gm \( \text{CaCO}_3 \) gives 0.1 mole of CO\(_2\)

\[ \text{Na}_2\text{CO}_3 + \text{H}_2\text{O} + \text{CO}_2 \rightarrow 2 \text{NaHCO}_3 \]

1 mole of CO\(_2\) gives 2 moles of NaHCO\(_3\) -> 0.1 mole of CO\(_2\) gives 0.2 mole of NaHCO\(_3\)

---
2004 EAMCET

Study the following table: Which two compounds have least weight of oxygen (molecular weights of compounds are given in brackets)

<table>
<thead>
<tr>
<th>Compound (mol. wt.)</th>
<th>Weight of compound (in g) taken</th>
</tr>
</thead>
<tbody>
<tr>
<td>I. CO2 (44)</td>
<td>4.4</td>
</tr>
<tr>
<td>II. NO2 (46)</td>
<td>2.3</td>
</tr>
<tr>
<td>III. H2O2 (34)</td>
<td>6.8</td>
</tr>
<tr>
<td>IV. SO2 (64)</td>
<td>1.6</td>
</tr>
</tbody>
</table>

1) II & IV  
2) I & III  
3) I & II  
4) III & IV

Ans: 1

\[ \%O = \frac{\text{wt of organic compound}}{\text{gram molecular wt of compound}} \times 100 \]

hence, (1) is correct.

2004 EAMCET

Assertion (A) : 8 grams of methane occupies 11.207 litres of volume at 273 K and 1 atm. pressure.

Reason (R) : One mole of any gas at S.T.P. occupies 22.414 litres of volume.

The correct answer is
1) Both (A) and (R) are true and (R) is the correct explanation of (A)
2) Both (A) and (R) are true, but (R) is not the correct explanation of (A)
3) (A) is true and (R) is not true
4) (A) is not true but (R) is true

Ans: 1

Solution :

8 gms of methane = ½ mole of methane
1 mole occupies 22.4 lit at STP
hence half mole occupies 11.2 lit
hence (1) is correct

2003 EAMCET

In a balanced equation H2SO4 + xHI = H2S + yI2 + zH2O the values of x, y and z are

1) x = 3, y = 5, z = 2  
2) x = 4, y = 8, z = 5  
3) x = 8, y = 4, z = 4  
4) x = 5, y = 3, z = 1

Ans: 3

Solution :

\[ \text{H}_2\text{SO}_4 + 8\text{HI} \rightarrow \text{H}_2\text{S} + 4\text{I}_2 + 4\text{H}_2\text{O} \]

2002 EAMCET

0.078 grams of a hydrocarbon occupy 22.414 ml volume at STP. The empirical formula of the hydrocarbon is CH. The molecular formula of the hydrocarbon is ............

1) C2H2  
2) C6H6  
3) C8H8  
4) C4H4

Ans: 2

Solution :

22.4 ml volume of a hydrocarbon weight is 0.078 gm
22.400 ml volume of a hydrocarbon weight ——>

\[
\frac{22.400 \times 0.078}{22.4} = 78 \text{ gm (molecular weight)}
\]

Molecular weight of C6H6 = 78 g
2002 EAMCET
When ammonium nitrate is gently heated, an oxide of nitrogen is formed. What is the oxidation state of nitrogen in this oxide?
1) +4                         2) +2                         3) +3                                 4) +1
Ans: 4
Solution :
NH\textsubscript{4}NO\textsubscript{3} \rightarrow \text{N}_2O + 2\text{H}_2\text{O}
oxidation state of nitrogen in N\textsubscript{2}O is + 1

---

2002 EAMCET
Two grams of sulphur is completely burnt is oxygen to form SO\textsubscript{2}. In this reaction, what is the volume (in litres) of oxygen consumed at STP?
(At. Wts. of sulphur and oxygen are 32. and 16 respectively)
1) \frac{16}{22.414}              2) \frac{22.414}{16}               3) \frac{22.414}{30}                              4) \frac{32}{22.414}
Ans: 2
Solution :
S + O\textsubscript{2} \rightarrow \text{SO}_2
32 gm of S consumes 22.4 lit of O\textsubscript{2} at STP
2 gm of S consumes = \frac{22.4 \times 2}{32} lit of O\textsubscript{2} at STP
= \frac{22.4}{16}

---

2002 EAMCET
Which one of the following is an example for exothermic reaction?
1) H\textsubscript{2}(g) + Cl(g) \rightarrow 2\text{HCl}(g) \Delta H = -184.6 KJ
2) N\textsubscript{2}(g) + O\textsubscript{2} (g) \rightarrow 2\text{NO} (g) \Delta H = +180.80
3) C(graphite) + H\textsubscript{2}O(g) \rightarrow \text{CO}(g) + \text{H}_2(g) \ - 181.4 KJ
4) C(graphite) + 2S (s) + 91.9 KJ \rightarrow \text{CS}_2 (l)
Ans: 1
Solution :
(1) choice is a correct answer, Because exothermic reaction have \Delta H value negative

---

2002 EAMCET
What is the oxidation state of Fe in the product formed when acidified potassium ferrocyanide is treated with H\textsubscript{2}O\textsubscript{2}?
1) +2                    2) +6                 3) +1                     4) +3
Ans: 4
\[ 2\text{K}_4\left[\text{Fe(CN)}_6\right] + \text{H}_2\text{O}_3 + \text{H}_2\text{SO}_4 \rightarrow 2\text{K}_3\left[\text{Fe(CN)}_6\right] + \text{K}_2\text{SO}_4 + 2\text{H}_2\text{O} \]
Oxidation state of Fe in \[ \text{K}_3\left[\text{Fe(CN)}_6\right] \] is +3
\[ x + 3 - 6 = 0 \Rightarrow x = +3 \]
2001 EAMCET
What the oxidation state of chlorine in hypochlorous acid?
1) +7  2) +5  3) +3  4) +1
Ans: 4
Solution:
Hypochlorous acid = HOCl
\[ x +1- 2 = 0 \]
x = +1

2001 EAMCET
50 grams of calcium carbonate was completely burnt in air. What is the weight (in grams) of the residue? Atomic weights of Ca, C and O are 40, 12 and 16 respectively)
1) 2.8  2) 28  3) 4.4  4) 44
Ans: 2
Solution:
CaCO\(_3\) → CaO + CO
100 gm of CaCO\(_3\) gives 56 gm of CaO
\[
\frac{50 \times 56}{100} = 28
\]

Find the equivalent weights in each case
(a) \(\text{As}_2\text{O}_3 + 5\text{H}_2\text{O} \rightarrow 2\text{AsO}_4^{3-} + 10\text{H}^+ + 4e\)
(b) \(\text{MnO}_4^- + 8\text{H}^+ + 5e \rightarrow \text{Mn}^{2+} + 4\text{H}_2\text{O}\)
(c) \(\text{Cr}_2\text{O}_7^{2-} + 14\text{H}^+ + 6e \rightarrow 2\text{Cr}^{3+} + 7\text{H}_2\text{O}\)

\[ E_{\text{red/oxi}} = \frac{\text{Mol. weight of reductant or oxidant}}{\text{Number of electrons gained or lost by one molecule of reductant or oxidant or valence factor}} \]

(a) \(E_{\text{As}_2\text{O}_3} = \frac{M_{\text{As}_2\text{O}_3}}{4} \quad (\text{As}_2^{3+} \rightarrow 2\text{As}^{5+} + 4e)\)
(b) \(E_{\text{MnO}_4^-} = \frac{M_{\text{MnO}_4}}{5} \quad (\text{Mn}^{7+} + 5e \rightarrow \text{Mn}^{2+})\)
(c) \(E_{\text{Cr}_2\text{O}_7^{2-}} = \frac{M_{\text{Cr}_2\text{O}_7}}{6} \quad (\text{Cr}_2^{6+} \rightarrow 2\text{Cr}^{3+} + 6e)\)

Question: Find the equivalent weights in each case
(d) \(\text{C}_2\text{O}_4^{2-} \rightarrow 2\text{CO}_2 + 2e\)
(e) \(\text{FeC}_2\text{O}_4 \rightarrow \text{Fe}^{3+} + 2\text{CO}_2 + 3e\)
(f) \(2\text{CuSO}_4 + 2e \rightarrow \text{Cu}_2^{1+} + \text{SO}_4^{2-}\)
Solution

\[
E_{C_2O_4^{2-}} = \frac{M_{\text{CuS}}}{2} \quad (C_2^{3+} \rightarrow 2C^{4+} + 2e) \\
E_{FeC_2O_4} = \frac{M_{\text{Fe}} + C_{\text{O}} + C_{\text{H}}} {3} \quad (Fe^{2+} + C_2^{3+} \rightarrow Fe^{3+} + 2C^{4+} + 3e) \\
E_{CuSO_4} = \frac{M_{\text{CuSO_4}}}{1} \quad (2Cu^{2+} + 2e \rightarrow Cu_2^+) 
\]

Question

Reaction, \(2Br^-_{(aq)} + Cl_2_{(aq)} \rightarrow 2Cl^-_{(aq)} + Br_2_{(aq)}\), is used for commercial preparation of bromine from its salts. Suppose we have 50 mL of a 0.060 \(M\) solution of NaBr. What volume of a 0.050 \(M\) solution of Cl_2 is needed to react completely with the Br^-?

Solution:

\[
2Br^- + Cl_2 \rightarrow 2Cl^- + Br_2 \\
\text{or} \quad 2Br^- \rightarrow Br_2^0 + 2e \quad \text{(Valence factor for Br}^{-1} = 1) \\
\text{Cl}_2^0 + 2e \rightarrow 2Cl^- \quad \text{(Valence factor for Cl}_2 = 2) \\
\text{Meq of Cl}_2 = \text{Meq of Br}^- \\
0.05 \times 2 \times V = 50 \times 0.60 \times 1 \\
V = 300 \text{ mL}
\]
Question:

What mass of Na₂S₂O₃.5H₂O is needed to make 500 cm³ of 0.200 N solution for the reaction:

\[ 2S_2O_3^{2-} + I_2 \rightarrow S_4O_6^{2-} + 2I^- \]

Solution:

\[ \text{Meq. of } Na_2S_2O_3 = 0.2 \times 500 = 100 \quad (\because \text{Meq.} = N \times V_{(\text{in mL})}) \]

\[ \frac{w}{M/1} \times 1000 = 100 \quad (\because \text{Meq.} = \frac{wt}{E.wt} \times 1000) \]

or \[ \frac{w}{248} \times 1000 = 100 \]

(Mol. wt. Na₂S₂O₃.5H₂O = 248)

\[ w = 24.8 \text{ g} \]

Question:

How many equivalents are there per mole of H₂S in its oxidation to SO₂?

Solution:

\[ \text{H}_2\text{S} \rightarrow \text{SO}_2 \]

\[ \text{S}^{2-} \rightarrow \text{S}^{4+} + 6e^- \]

\[ N_{\text{H}_2\text{S}} = M_{\text{H}_2\text{S}} \times 6 \]

\[ \therefore \quad \frac{\text{Moles}}{\text{Equivalent}} = \frac{N}{M} = 6 \]

Thus one mole of H₂S has 6 equivalent in it.

Question:

12.53 mL of 0.0509 M SeO₂ reacted with 25.52 mL 0.1 M CrSO₄ solution. In the reaction Cr²⁺ was oxidized to Cr³⁺. To what oxidation state selenium was converted in the reaction? Write the redox change for SeO₂.
Solution:

\[ \text{Se}^{4+} + (a - 4)e \rightarrow \text{Se}^2+ \]
\[ n = a - 4 \]
\[ \text{or} \quad \text{Cr}^{2+} \rightarrow \text{Cr}^{3+} + e \]

\[ \text{Meq. of SeO}_2 = \text{Meq. of CrSO}_4 \]
\[ 0.0509 \times n \times 12.53 = 0.1 \times 1 \times 25.52 \]
\[ \therefore \quad n = 4 \]
\[ a - 4 = 4 \]
\[ \text{or} \quad a = 0 \]

Thus redox change is: \[ \text{Se}^{4+} + 4e \rightarrow \text{Se}^{2+} \]

Question:

In a reaction, \( \text{Cr}_2\text{O}_7^{2-} \) is reduced to \( \text{Cr}^{3+} \). What is concentration of 0.1 \( M \text{K}_2\text{Cr}_2\text{O}_7 \) in equivalent per litre?

\[ \text{Cr}_2\text{O}_7^{2-} + 14\text{H}^+ + 6e \rightarrow 2\text{Cr}^{3+} + 7\text{H}_2\text{O} \]

Solution:

\[ N = M \times \text{valence factor} \]
\[ \therefore \quad N = 0.1 \times 6 \quad \text{ (Cr}_2^{6+} + 6e \rightarrow 2\text{Cr}^{3+}) \]
\[ \text{or} \quad N_{\text{K}_2\text{Cr}_2\text{O}_7} = 0.6 \quad N \]

Question:

What is molarity and normality of a \( \text{MnO}_4^- \) solution if 32.00 mL of the solution is required to titrate 40.00 mL of 0.400 \( N \text{Fe}^{2+} \)?

\[ \text{MnO}_4^- + 5\text{Fe}^{2+} + 8\text{H}^+ \rightarrow \text{Mn}^{2+} + 5\text{Fe}^{3+} + 4\text{H}_2\text{O} \]
Solution:

\[
\text{Meq. of } \text{MnO}_4^- = \text{Meq. of } \text{Fe}^{2+} \quad (\text{Mn}^{7+} + 5e \rightarrow \text{Mn}^{2+})
\]

\[
N \times 32 = 40 \times 0.400 \quad (\text{Fe}^{2+} \rightarrow \text{Fe}^{3+} + e)
\]

\[
N = 0.5
\]

or

\[
M_{\text{KMnO}_4} = \frac{N_{\text{KMnO}_4}}{\text{valence factor}}
\]

Also,

\[
M_{\text{KMnO}_4} = \frac{0.5}{5} = 0.1 \ M
\]

Question:

\[\text{Mn}^{2+}(aq) \text{ can be determined by titration with } \text{MnO}_4^-(aq)\]

\[
3\text{Mn}^{2+} + 2\text{MnO}_4^- \rightarrow 6\text{MnO}_2 + 2\text{H}_2\text{O}
\]

A 25.00 mL sample of Mn\(^{2+}\) (aq) requires 34.77 mL of 0.05876 \(M\) KMnO\(_4\) (aq) for its titration. What is the molarity of the Mn\(^{2+}\)(aq)?

Solution:

\[
\text{Mn}^{2+} \rightarrow \text{Mn}^{4+} + 2e
\]

\[
\text{Mn}^{7+} + 3e \rightarrow \text{Mn}^{4+}
\]

Also,

\[
\text{Meq. of } \text{MnO}_4^- = \text{Meq. of } \text{Mn}^{2+}
\]

\[
0.05876 \times 3 \times 34.77 = M \times 2 \times 25
\]

\[
M_{\text{Mn}^{2+}} = 0.1226 \ M
\]
Question:

A 1.100 g sample of copper ore is dissolved and the Cu\(^{2+}\)(aq) is treated with excess KI. The liberated I\(_2\) requires 12.12 mL of 0.10 M Na\(_2\)S\(_2\)O\(_3\) solution for titration. What is % copper by mass in the ore?

Solution:

\[
\text{Cu}^{2+} + e \rightarrow \text{Cu}^{1+} \\
2\text{I}^- \rightarrow \text{I}_2 + 2e \\
2\text{S}_2\text{O}_3^{2-} \rightarrow \text{S}_4\text{O}_6^{2-} + 2e
\]

\[
\text{Meq. of Cu}^{2+} = \text{Meq. of liberated I}_2 = \text{Meq. of Na}_2\text{S}_2\text{O}_3
\]

\[
= 12.12 \times 0.1 \times 1 = 1.212
\]

\[
\therefore \quad \frac{w_{\text{Cu}^{2+}}}{M/1} \times 1000 = 1.212
\]

\[
\therefore \quad w_{\text{Cu}^{2+}} = \frac{1.212 \times M}{1000} = \frac{1.212 \times 63.6}{1000} = 0.077 \text{ g}
\]

\[
\therefore \quad w_{\text{Cu}} = w_{\text{Cu}^{2+}} = 0.077 \text{ g}
\]

\[
\therefore \quad \% \text{ Cu} = \frac{0.077}{1.100} \times 100 = 7.00\%
\]
Question:

What mass of $\text{N}_2\text{H}_4$ can be oxidised to $\text{N}_2$ by 24.0 g $\text{K}_2\text{CrO}_4$, which is reduced to $\text{Cr(OH)}_4^{-}$.

Solution:

\[ \text{N}_2^{2-} \rightarrow \text{N}_2^{0} + 4e \]
\[ \text{Cr}^{6+} + 3e \rightarrow \text{Cr}^{3+} \]

Meq. of $\text{N}_2\text{H}_4 = \text{Meq. of K}_2\text{CrO}_4$

\[ \frac{w}{32/4} \times 1000 = \frac{24}{194.2/3} \times 1000 \quad \text{(E. wt. = M. wt.}\ \frac{\text{Valence factor}}{\text{Valence factor}}) \]

\[ w_{\text{N}_2\text{H}_4} = 2.97 \text{ g} \]

Question:

It requires 40.0 mL of 0.50 $M \text{Ce}^{4+}$ to titrate 10.0 mL of 1.0 $M \text{Sn}^{2+}$ to $\text{Sn}^{4+}$. What is the oxidation state of cerium in the reduced product.

Solution:

\[ \text{Sn}^{2+} \rightarrow \text{Sn}^{4+} + 2e \]
\[ ne + \text{Ce}^{4+} \rightarrow \text{Ce}^{(4-n)^+} \]

Meq. of $\text{Ce}^{4+} = \text{Meq. of Sn}^{2+}$

\[ 40 \times 0.5 \times n = 1 \times 2 \times 10.0 \]
\[ n = 1 \]

\[ \therefore \quad \text{Ce}^{4+} + e \rightarrow \text{Ce}^{3+} \]

Thus oxidation state of Ce in reduced state is +3.

Question:

Calculate the mass of oxalic acid ($\text{H}_2\text{C}_2\text{O}_4$) which can be oxidised to $\text{CO}_2$ by 100.0 mL of $\text{MnO}_4^{-}$ solution, 10 mL of which is capable of oxidising 50.0 mL of 1.0 $M \text{I}^{-}$ to $\text{I}_2$.
Solution:

\[
\begin{align*}
\text{Mn}^{7+} + 5e^- & \rightarrow \text{Mn}^{2+} \\
2I^- & \rightarrow I_2 + 2e^- \\
\text{Meq. of } I^- & = \text{Meq. of } \text{MnO}_4^- \\
50 \times 1.0 & = 10 \times N \\
N_{\text{MnO}_4^-} & = 5 \\
\text{Now } \text{Mn}^{7+} + 5e^- & \rightarrow \text{Mn}^{2+} \\
C_2^{3+} & \rightarrow 2C^{4+} + 2e^- \\
\text{Meq. of } H_2C_2O_4 & = \text{Meq. of } \text{MnO}_4^- \\
\frac{w}{90/2} \times 1000 & = 5 \times 100 \\
w & = 22.5 \text{ g}
\end{align*}
\]

Question:

A KMnO\textsubscript{4} solution can be standardised by titration against As\textsubscript{2}O\textsubscript{3} (s). A 0.1156 g sample of As\textsubscript{2}O\textsubscript{3} requires 27.08 mL of the KMnO\textsubscript{4} (aq) for its titration. What is the molarity of the KMnO\textsubscript{4} (aq) [As = 75].

\[5\text{As}_2\text{O}_3 + 4\text{MnO}_4^- + 9\text{H}_2\text{O} + 12\text{H}^+ \rightarrow 10\text{H}_2\text{AsO}_4^- + 4\text{Mn}^{2+}\]

Solution:

\[
\begin{align*}
\text{Meq. of As}_2\text{O}_3 & = \text{Meq. of KMnO}_4 & \text{(As}_2^{3+} & \rightarrow 2\text{As}^{5+} + 4e) \\
\frac{0.1156}{198/4} \times 1000 & = M \times 5 \times 27.08 & \text{(Mn}^{7+} + 5e^- & \rightarrow \text{Mn}^{2+}) \\
\therefore M & = 0.0172 \\
M_{\text{MnO}_4^-} & = 0.0172 M
\end{align*}
\]
Question:

A particular acid-rain water has $\text{SO}_3^{2-}$. If a 25.00 mL sample of this water requires 34.08 mL of 0.01964 $M \text{KMnO}_4$ for its titration, what is the molarity of $\text{SO}_3^{2-}$ in acid-rain?

$2\text{MnO}_4^- + 5\text{SO}_3^{2-} + 6\text{H}^+ \rightarrow 5\text{SO}_4^{2-} + 2\text{Mn}^{2+} + 3\text{H}_2\text{O}$

Solution:

\[
\text{Meq. of } \text{SO}_3^{2-} = \text{Meq. of } \text{KMnO}_4 \\
N \times 25 = 34.08 \times 0.01964 \times 5
\]

\[
N = 0.1339
\]

\[
M = \frac{0.1339}{2} = 0.0669
\]

$\text{S}^{4+} \rightarrow \text{S}^{6+} + 2e$

\[
N_{\text{SO}_3^{2-}} = 0.1339N
\]

\[
M_{\text{SO}_3^{2-}} = \frac{0.1339}{2} = 0.0669M
\]
Question:

A solution containing 1.984 g of crystalline \( \text{Na}_2\text{S}_2\text{O}_3 \cdot x\text{H}_2\text{O} \) in water required 40 mL of \( \text{N/5} \) iodine solution for complete reaction. Calculate the value of \( x \).

Solution:

\[
2\text{S}_2\text{O}_3^{2-} \rightarrow \text{S}_4\text{O}_6^{2-} + 2e
\]

\[
2e + I_2 \rightarrow 2I^-
\]

Also

\[
\text{Meq. of } I_2 = \text{Meq. of } \text{Na}_2\text{S}_2\text{O}_3 \cdot x\text{H}_2\text{O}
\]

\[
40 \times \frac{1}{5} = \frac{1.984}{M/1} \times 1000
\]

\[
M_{\text{Na}_2\text{S}_2\text{O}_3 \cdot x\text{H}_2\text{O}} = 248
\]

\[
2 \times 23 + 2 \times 32 + 16 \times 3 + 18x = 248
\]

or

\[
x = 5
\]

\[
\text{Na}_2\text{S}_2\text{O}_3 \cdot 5\text{H}_2\text{O}
\]

Question:

If 10.00 g \( \text{V}_2\text{O}_5 \) is dissolved in acid and reduced to \( \text{V}^{2+} \) by treatment with \( \text{Zn} \) metal, how many mole \( I_2 \) could be reduced by the resulting \( \text{V}^{2+} \) solution as it is oxidised to \( \text{V}^{4+} \). At. weight of \( \text{V} = 51 \).

Solution:

\[
\text{V}_2^{5+} + 6e \rightarrow 2\text{V}^{2+}
\]

\[
\text{Zn} \rightarrow \text{Zn}^{2+} + 2e
\]

\[
\text{V}^{2+} \rightarrow \text{V}^{4+} + 2e
\]

\[
l_2 + 2e \rightarrow 2I^-
\]

and

\[
\text{Meq. of } \text{V}^{2+} \text{ (v.f. } = 3) = \text{Meq. of } \text{V}_2\text{O}_5 \text{ (v.f. } = 6)
\]

\[
= \frac{10}{182/6} \times 1000
\]
Meq. of V^{2+} (v.f. = 2) = \frac{10}{182} \times 1000 \times \frac{2}{3}

Meq. of V^{2+} (v.f. = 2) = \text{Meq. of I}_2

\frac{10 \times 6}{182} \times 1000 \times \frac{2}{3} = \text{Meq. of I}_2

\text{or} \quad \text{Meq. of I}_2 = 219.78

m \ \text{Mole of I}_2 = \frac{219.78}{2} = 109.89

\text{Mole of I}_2 = \frac{109.89}{1000}

\text{Mole of I}_2 = 0.1098

\textbf{Question:}

A 0.56 g sample of limestone is dissolved in acid and the calcium is precipitated as calcium oxalate. The precipitate is filtered, washed with water and dissolved in dil. H\textsubscript{2}SO\textsubscript{4}. The solution required 40 mL of 0.25N KMnO\textsubscript{4} solution for titration. Calculate percentage of CaO in limestone sample.

\textbf{Solution:}

\[
\text{CaCO}_3 + 2\text{H}^+ \rightarrow \text{Ca}^{2+} + \text{CO}_2 + \text{H}_2\text{O}
\]

\[
\text{Ca}^{2+} + \text{C}_2\text{O}_4^{2-} \rightarrow \text{CaC}_2\text{O}_4
\]

\[
\text{CaC}_2\text{O}_4 + \text{KMnO}_4 \text{ in presence of } \text{H}_2\text{SO}_4 \text{ show the following redox changes:}
\]

\[
\text{C}_2\text{O}_4^{3+} \rightarrow 2\text{C}^{4+} + 2e
\]

\[
\text{Mn}^{7+} + 5e \rightarrow \text{Mn}^{2+}
\]

The above set of reaction shows:

\[
\text{Meq. of CaO} = \text{Meq. of CaC}_2\text{O}_4 = \text{Meq. of KMnO}_4
\]

(Valence factor is two for Ca throughout the changes)

\[
\frac{\nu_{\text{CaO}}}{56} \times 1000 = 40 \times 0.25
\]

\[
\therefore \quad \nu_{\text{CaO}} = 0.28
\]

\[
\therefore \quad \% \text{ of CaO} = \frac{0.28}{0.56} \times 100 = 50\%
\]
Question:

How many mL of aqueous solution of KMnO₄ containing 158 g/L must be used to complete the conversion of 75.0 g of KI to iodine by the reaction:

\[ \text{KMnO}_4 + \text{KI} + \text{H}_2\text{SO}_4 \rightarrow \text{K}_2\text{SO}_4 + \text{MnSO}_4 + \text{I}_2 + 6\text{H}_2\text{O} \]

Solution:

\[ N_{\text{KMnO}_4} = \frac{158}{31.6 \times 1} = 5.0 \]

\[ (\text{Mn}^{7+} + 5e \rightarrow \text{Mn}^{2+} : E = \frac{M}{5} = \frac{158}{5} = 31.6) \]

\[ \text{Meq. of KI} = \frac{75 \times 1000}{166} = 451.8 \quad (2\text{I}^{-} \rightarrow \text{I}_2 + 2e : E = \frac{M}{1} = \frac{166}{1} ) \]

\[ \text{Meq. of KMnO}_4 = \text{Meq. of KI} \\
\quad 5 \times \nu = 451.8 \\
\quad \nu = 90.36 \text{ mL} \]

Question:

What is the maximum weight of Cl₂ obtained by the action of 1 g HCl on 1 g MnO₂.

Solution:

\[ \text{MnO}_2 + 4\text{HCl} \rightarrow \text{MnCl}_2 + \text{Cl}_2 + 2\text{H}_2\text{O} \]

Initial mole

\[
\begin{array}{c|c|c|c|c|c}
\text{Mole} & \frac{1}{87} & \frac{1}{36.5} \\
\hline
\text{Initial} & 0.011 & 0.027
\end{array}
\]

Final mole

\[
\begin{array}{c|c|c|c|c|c}
\text{Mole} & 0.027 & 0.027 & 0.027 \\
\hline
\text{Initial} & 0.011 & \frac{1}{4} & \frac{1}{4} & \frac{1}{4}
\end{array}
\]

\[ \therefore \text{Mole of Cl}_2 \text{ formed} = \frac{0.027}{4} \]

\[ \therefore \text{wt. of Cl}_2 \text{ formed} = \frac{0.027}{4} \times 71 = 0.4792 \text{ g} \]
Question:

25 mL of 0.017 H₂SO₃ in strongly acidic medium required 16.9 mL of 0.01 M KMnO₄ and in neutral medium required 28.6 mL of 0.01M KMnO₄ for complete conversion of SO₃²⁻ to SO₄²⁻. Assign the oxidation no. of Mn in the product formed in each case.

Solution:

\[ \text{SO}_3^{2-} \rightarrow \text{SO}_4^{2-} \]
\[ S^{4+} \rightarrow S^{6+} + 2e \]
\[ \therefore \text{Valence factor of SO}_3 = 2 \]

In acid medium:

\[ \text{Meq. of SO}_3^{2-} = \text{Meq. of MnO}_4^- \]
\[ 25 \times 0.017 \times 2 = 16.9 \times 0.01 \times n_1 \]
\[ \therefore n_1 = 5 \]
\[ \therefore \text{Mn}^{7+} + 5e \rightarrow \text{Mn}^{2+} \]

In neutral medium:

\[ \text{Meq. of SO}_3^{2-} = \text{Meq. of MnO}_4^- \]
\[ 25 \times 0.017 \times 2 = 28.6 \times 0.01 \times n_2 \]
\[ \therefore n_2 = 3 \]
\[ \therefore \text{Mn}^{7+} + 3e \rightarrow \text{Mn}^{4+} \]
Topic wise grouping of information kind of Dictionary of General Chemistry

Graphite and Diamond comparison

With respect to graphite and diamond, which of the following statement(s) given below is(are) correct?
(a) Graphite is harder than diamond.
(b) Graphite has higher electrical conductivity than diamond.
(c) Graphite has higher thermal conductivity than diamond.
(d) Graphite has higher C—C bond order than diamond.

Solution:
The given facts about graphite and diamond are as follows:
Graphite is less hard than diamond. Graphite has higher electrical and thermal conductivities than diamond. This is due to highly delocalized nature of π electrons. Graphite has higher C—C bond order (due to σ and π bonds) than that of diamond (which has only σ bonds).
Therefore, the choices (b), (c) and (d) are correct.

Stability of lyophobic colloidal particles

Choose the correct reason(s) for the stability of lyophobic colloidal particles.
(a) Preferential adsorption of ions on their surface from the solution.
(b) Preferential adsorption of solvent on their surface from the solution.
(c) Attraction between different particles having opposite charges on their surface.
(d) Potential difference between the fixed layer and the diffused layer of opposite charges around the colloidal particles.

Solution:
The stability of lyophobic colloidal particles is due to the preferential adsorption of common ion from the solution. Each colloidal particle has the same charge and are thus kept away from each other due to electronic repulsion (choice a).
There exists potential difference between the fixed and diffused layer of opposite charge (choice d).
Therefore, the choice (a) and (d) are correct.

Physiorption and Chemisorption comparison

The given graph/data I, II, III and IV represent general trends observed for different physiorption and chemisorption processes under mild conditions of temperature and pressure. Which of the following choice(s) about I, II, III and IV is(are) correct.

![Graph](image)
(a) 1 is physisorption and II is chemisorption  
(c) IV is chemisorption and II is chemisorption
(b) 1 is physisorption and III is chemisorption  
(d) IV is chemisorption and III is chemisorption

**Solution:**

We have the following facts:
- Adsorption decreases with increase in temperature at constant pressure in case of physisorption.
- Adsorption increases initially with increase in temperature at constant pressure in case of chemisorption as more molecules satisfy the energy of activation.
- The enthalpy change during chemisorptions is high due to the bond formation between adsorbent and adsorbate.

With these guidelines, we will have:

- Graph I represents physisorption
- Graph III represents physisorption

Therefore, the choices (a) and (c) are correct.

A compound $M_yX_n$ has cubic close packing (CCP) arrangement of $X$. Its unit cell structure is shown below.

The empirical formula of the compound is:
(a) $MX$  
(b) $MX_2$  
(c) $M_2X$  
(d) $M_2X_3$

**Solution:**

The atom at the corner of the cube is shared amongst 8 unit cells; the atom at the centre of a face is shared between 2 unit cells, the atom at the centre of edges is shared amongst 4 unit cells and that at the centre of the cube belongs wholly to the unit cell.

There are eight $X$ atoms at the corners and six $X$ atoms at the centre of faces. Thus, the number of $X$ atoms per
unit cell is

\[ 8 \left( \frac{1}{8} \right) + 6 \left( \frac{1}{2} \right) = 1 + 3 = 4 \]

There are four M atoms at the centre of edges and one M atom at the centre of the cube. Thus, the number of M atoms per unit cell is

\[ 4 \left( \frac{1}{4} \right) + 1(1) = 1 + 1 = 2 \]

The formula of the compound will be \( M_2X_4 \) and the empirical formula will be \( MX_2 \). Therefore, the choice (b) is correct.

HCl, HBr and HI react with AgNO3 to give a precipitate that dissolves in Na2S2O3

Which of the following halides react(s) with AgNO3(aq) to give a precipitate that dissolves in Na2S2O3(aq)?

(a) HCl  
(b) HF  
(c) HBr  
(d) HI

**Solution:**

The precipitates of AgCl, AgBr and AgI are soluble in Na2S2O3(aq) forming the complex Na2[Ag(S2O3)2]. For example,

\[ Na_2S_2O_3 + 2AgBr \rightarrow Ag_2S_2O_3 + 2NaBr \]

\[ Ag_2S_2O_3 + 3Na_2S_2O_3 \rightarrow 2Na[Ag(S_2O_3)_2] \]

AgF is soluble in water.

Therefore, the choices (a), (c) and (d) are correct.

Reversible expansion of an ideal gas

The reversible expansion of an ideal gas under adiabatic and isothermal conditions is shown in the figure. Which of the following statement(s) is(are) correct?

(a) \( T_1 = T_2 \)

(b) \( T_2 > T_1 \)

(c) \( w_{\text{isothermal}} > w_{\text{adiabatic}} \)

(d) \( \Delta U_{\text{isothermal}} > \Delta U_{\text{adiabatic}} \)

**Solution:**

We will have

\( T_1 = T_2 \) as the expansion is isothermal.

\( T_2 > T_1 \) as the adiabatic expansion involves cooling.

\( w_{\text{isothermal}} > w_{\text{adiabatic}} \) as the area under the isothermal curve is greater than that under adiabatic curve.

\( \Delta U_{\text{isothermal}} = 0 \) as temperature remains constant while \( \Delta U_{\text{adiabatic}} < 0 \) as the expansion occurs at the expance of internal energy.

Therefore, the choices (a) and (d) are correct.

**Note:** The choice (c) is correct if the magnitude of the work is considered. Since expansion carries negative sign as per IUPAC recommendations, the choice (c) will not be correct if the negative sign is also considered.
Acid Ionization Constant

\[ K_a = \frac{[H^+][A^-]}{[HA]} \]

Adiabatic change

\[ PV = k \]

Base Ionization Constant

\[ K_b = \frac{[OH^-][HB^+]}{[B]} \]

Bohr Radius

\[ a_0 = \frac{\hbar^2}{m_e k c^2} \]

Boiling Point Elevation

\[ \Delta T_b = i K_b \times molality \]

Buffer Design Equation

\[ pH \approx pK_a - \log \frac{[HA]_0}{[A^-]_0} \]

Cell Voltage

\[ E_{cell} = E_{cell}^\circ - \frac{RT}{nF} \ln Q = E_{cell}^\circ - \frac{0.0592}{n} \log Q \]

Charles' Law

\[ \frac{V}{t} = k \]

Atomic Structure

<table>
<thead>
<tr>
<th>DESCRIPTION</th>
<th>EQUATION</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bohr Radius</td>
<td>[ a_0 = \frac{\hbar^2}{m_e k c^2} ]</td>
</tr>
</tbody>
</table>
| De Broglie Wavelength | \[ \lambda = \frac{\hbar}{mv} \]
| Linear Momentum | \[ p = mv \]
| Planck's Quantized (Quantum) Energy Equation | \[ \Delta E = h \nu \]
| Radii of stable orbits in the Bohr model | \[ r = n^2 \frac{\hbar^2}{m_e k Z c^2} = n^2 \frac{a_0}{Z} \]
| Relationship between Energy and Principal Quantum Number | \[ E_n = -R_H \left( \frac{1}{n^2} \right) = -2.178 \times 10^{-18} \text{ joule} \]
Rydberg Equation

$$\Delta E = R_H \left( \frac{1}{n_i^2} - \frac{1}{n_f^2} \right)$$

Speed of Light to Wavelength and Frequency Relationship

$$c = \lambda \nu$$

**Electrochemistry**

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<tr>
<td>Cell Voltage</td>
<td>( E_{cell} = E_{cell}^0 - \frac{RT}{nF} \ln Q = E_{cell}^0 - \frac{0.0592}{n} \log Q )</td>
</tr>
<tr>
<td>Electric Current</td>
<td>( I = \frac{q}{t} )</td>
</tr>
<tr>
<td>Reaction Quotient</td>
<td>( Q = \frac{[C]^c[D]^d}{[A]^a[B]^b} )</td>
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<tr>
<td>Relationship between Equilibrium Constant and Cell Voltage</td>
<td>( \log K = \frac{nE^0}{0.0592} )</td>
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**Equilibrium**

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</tr>
<tr>
<td>Base Ionization Constant</td>
<td>( K_b = \frac{[OH^-][HB^+]}{[B]} )</td>
</tr>
<tr>
<td>Buffer Design Equation</td>
<td>( pH \approx pK_a - \log \frac{[HA]}{[A^-]}_0 )</td>
</tr>
<tr>
<td>Gas Pressure and Concentration Relationship</td>
<td>( K_p = K_c (RT)^\Delta n )</td>
</tr>
<tr>
<td>Ion Product Constant for Water</td>
<td>( K_w = [OH^-][H^+] = K_a \times K_b = 1.0 \times 10^{-14} \text{ at } 25^\circ C )</td>
</tr>
<tr>
<td>pH and pOH Relationship</td>
<td>( 14 = pH + pOH )</td>
</tr>
<tr>
<td>pH Defined</td>
<td>( pH = - \log [H^+] )</td>
</tr>
<tr>
<td>pK_a Definition</td>
<td>( pK_a = - \log K_a )</td>
</tr>
</tbody>
</table>
### Definition

\[ pK_b = -\log K_b \]

### pOH and Base Ionization Equilibrium Constant Relationship

\[ pOH = pK_b + \log \left( \frac{[HB^+]}{[B]} \right) \]

### Gases, Liquids, and Solutions

<table>
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<td>( PV = k )</td>
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<tr>
<td>Boiling Point Elevation</td>
<td>( \Delta T_b = iK_b \times \text{molality} )</td>
</tr>
<tr>
<td>Charles' Law</td>
<td>( \frac{V}{t} = k )</td>
</tr>
<tr>
<td>Combined Gas Law</td>
<td>( \frac{P_1V_1}{n_1T_1} = \frac{P_2V_2}{n_2T_2} )</td>
</tr>
<tr>
<td>Density of a Material</td>
<td>( D = \frac{m}{V} )</td>
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<tr>
<td>Freezing Point Depression</td>
<td>( \Delta T_f = iK_f \times \text{molality} )</td>
</tr>
<tr>
<td>Graham's Law of Effusion</td>
<td>( \frac{r_1}{r_2} = \sqrt{\frac{M_2}{M_1}} )</td>
</tr>
<tr>
<td>Ideal gas equation</td>
<td>( PV = nRT )</td>
</tr>
<tr>
<td>Kinetic Energy per Mole</td>
<td>( KE_{\text{mole}} = \frac{3}{2}RTn )</td>
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Vander waal's force

For one mole of a van der Waals gas when $b = 0$ and $T = 300$ K, the plot of $pV$ versus $1/V$ is shown below.

The value of the van der Waals constant $a$ is
(a) 1.0 L$^2$ atm mol$^{-2}$    (b) 4.5 L$^2$ atm mol$^{-2}$    (c) 1.5 L$^2$ atm mol$^{-2}$    (d) 3.0 L$^2$ atm mol$^{-2}$

**Solution:**

For one mole of a gas, the van der Waals equation is

$\left(p + \frac{a}{V^2}\right)(V - b) = RT$

When $b = 0$, we have

$\left(p + \frac{a}{V^2}\right)V = RT$  i.e.  $pV = RT - \frac{a}{V}$
The plot of \(pV\) versus \(1/V\) will be a straight line with slope equal to \(-\alpha\). Hence

Slope of the given straight line is
\[
\frac{(20.1-21.6)\text{L atm mol}^{-1}}{(3.0-2.0)\text{mol L}^{-1}} = -1.5 \text{ L}^2 \text{ atm mol}^{-2}
\]

Equating this to \(-\alpha\), we get
\[
\alpha = 1.5 \text{ L}^2 \text{ atm mol}^{-2}
\]

Therefore, the choice (c) is correct.

Dilute solution containing 2.5 gm of nonvolatile

For a dilute solution containing 2.5 g of a non-volatile non-electrolyte solute in 100 g of water, the elevation of boiling point at 1 atm pressure is 2 °C. Assuming concentration of solute is much lower than the concentration of solvent, the vapour pressure of the solution (take \(K_b = 0.76 \text{ K kg mol}^{-1}\)) is

(a) 724 mmHg  
(b) 740 mmHg  
(c) 736 mmHg  
(d) 718 mmHg

**Solution:**

From the expression \(\Delta T_b = K_b m\), we find that the molality of solute \((m)\) in the solution is
\[
m = \frac{\Delta T_b}{K_b} = \frac{2 \text{ K}}{0.76 \text{ K kg mol}^{-1}} = \frac{2}{0.76} \text{ mol kg}^{-1}
\]

Since \(m = n_2/n_1\) (where \(n_2\) is the amount of solute and \(n_1\) is the mass of solvent expressed in kg), we get
\[
n_2 = mn_1 = \left(\frac{2}{0.76} \text{ mol kg}^{-1}\right) (0.1 \text{ kg}) = \frac{2}{7.6} \text{ mol}
\]

From the expression \(-\Delta p = x_2 p_1\) of the relative lowering of vapour pressure of solvent (whose vapour pressure is 1 atm as the solute is non-volatile), we find that
\[
x_2 = \frac{n_2}{n_1 + n_2} = \frac{n_2}{n_1} = \frac{2/7.6}{100/18} = \frac{36}{760} = 0.047
\]

Hence 
\[
\Delta p = \left(\frac{36}{760}\right) (760 \text{ mmHg}) = 36 \text{ mmHg}
\]

\[p = p_1 + \Delta p = 760 \text{ mmHg} - 36 \text{ mmHg} = 724 \text{ mmHg}
\]

Therefore, the choice (a) is correct.

Shape of XeO2F2 molecule is seesaw

The shape of XeO2F2 molecule is

(a) trigonal bipyramidal  (b) square planar  (c) tetrahedral  (d) see-saw

**Solution:**

The number of valence electrons in XeO2F2 is \(8 + 2 \times 6 + 2 \times 7 = 34\). These are distributed as shown in the following.
There are five pairs of electrons around Xe. To accommodate these, Xe undergoes dsp³ hybridization leading to the trigonal bipyramidal arrangement of five pairs of electrons. The structure of XeO₂F₂ is

The shape of XeO₂F₂ is sea-saw.
Therefore, the choice (d) is correct.

Various increasing and Decreasing properties

Decreasing order of reactivity towards S₂N₂ displacement:
- n-Butyl bromide, isobutyl bromide, sec-butyl bromide, tert-butyl bromide

Increasing order of reactivity towards S₂N₁ displacement:
- 1-bromopentane, 2-bromopentane, 2-bromo-2-methylbutane

Increasing reactivity sequence of S₂N₁ displacement of halogen: CH₃X, 1° X, 2° X, 3° X.

Increasing reactivity sequence of S₂N₂ displacement of halogen: CH₃X, 1° X, 2° X, 3° X.

Increasing reactivity of alcohols towards gaseous HBr:
- 2-butanol, 2-methyl-1-propanol, 2-methyl-2-propanol

Increasing order or reactivity towards E₂ dehydrohalogenation of the bromides:
- ethyl bromide, n-propyl bromide, isobutyl bromide, neopentyl bromide

Increasing stability of alkenes: R₂C=CR₂, R₂C=CHR, R₂C=CH₂, RCH=CH₂, CH₂=CH₂

Increasing dehydration of alcohol in the presence of H₂SO₄:
- ethyl alcohol, isopropyl alcohol, tert-butyl alcohol

Decreasing order of reactivity towards E₂ dehydrohalogenation:
- 2-bromo-2-methylbutane, 1-bromopentane, 2-bromopentane, 3-bromopentane

Increasing order of dehydrohalogenation:
- 1-bromo-3-methylbutane, 2-bromo-2-methylbutane, 3-bromo-2-methylbutane
**Calculate multiple bond energy of a C triple bond**

Using the data provided, calculate the multiple bond energy of a $\text{C}≡\text{C}$ bond in $\text{C}_2\text{H}_2$:

\[
\begin{align*}
2\text{C}(s) + \text{H}_2(g) & \rightarrow \text{C}_2\text{H}_2(g) & \Delta H_1 &= 225 \text{ kJ mol}^{-1} \\
2\text{C}(s) & \rightarrow 2\text{C}(g) & \Delta H_2 &= 1410 \text{ kJ mol}^{-1} \\
\text{H}_2(g) & \rightarrow 2\text{H}(g) & \Delta H_3 &= 330 \text{ kJ mol}^{-1}
\end{align*}
\]

Take the bond energy of $\text{C}–\text{H}$ bond equal to $350 \text{ kJ mol}^{-1}$.

(a) $1165 \text{ kJ mol}^{-1}$  
(b) $837 \text{ kJ mol}^{-1}$  
(c) $865 \text{ kJ mol}^{-1}$  
(d) $815 \text{ kJ mol}^{-1}$

**Solution:**

Consider the following transformations.

\[
\begin{align*}
2\text{C}(s) + \text{H}_2(g) & \xrightarrow{\Delta H_1} \text{C}≡\text{C} & \text{H} & \xrightarrow{-2\epsilon_{\text{C}–\text{H}} - \epsilon_{\text{C}≡\text{C}}} \text{C}≡\text{C} & \text{H} \\
2\text{C}(g) + 2\text{H}(g) & \xrightarrow{\Delta H_2 + \Delta H_3} \text{C}≡\text{C} & \text{H} & \xrightarrow{-2\epsilon_{\text{C}–\text{H}}} \text{C}≡\text{C} & \text{H}
\end{align*}
\]

According to Hess’s law

\[
\Delta H_1 = \Delta H_2 + \Delta H_3 - 2\epsilon_{\text{C}–\text{H}} - \epsilon_{\text{C}≡\text{C}}
\]

This gives

\[
\epsilon_{\text{C}≡\text{C}} = \Delta H_2 + \Delta H_3 - \Delta H_1 - 2\epsilon_{\text{C}–\text{H}} = (1410 + 330 - 225 - 2 \times 350) \text{ kJ mol}^{-1} = 815 \text{ kJ mol}^{-1}
\]

Therefore, the choice (d) is correct.

**Some facts student must know**

- **Largest Anion** - At⁻¹
- **Metals having highest b.pt and m.pt.** = Mercury and Tungsten
- **Most reactive solid element** - Li
- **Most reactive liquid element** - Cs
- **total no. of gaseous element in periodic table** - 11 (H, He, N, O, F, Ne, Cl, Ar, Kr, Xe, Ra)
- **TOTAL NO. OF LIQUID elements** - 6 (Ga, Br, Cs, Hg, Fr, EKa)
- **Liquid radioactive element** - Francium
- **N.metal wid highest M.Pt** - Carbon
- **Metal wid highest valency** = Plutonium
- **highest tensile strength** - Boron
- **Most ionic compound** = CsF
Strongest base = Cs (OH)

Strongest basic oxide = Cs₂O

Most conducting metal = Ag

**Zinc and Magnesium Oxide Battery**

\[
\text{Anode (oxidation): } \quad \text{Zn}(s) + 2\text{OH}^-_\text{(aq)} \rightarrow \text{ZnO}(s) + \text{H}_2\text{O}(l) + 2\text{e}^-
\]

\[
\text{Cathode (reduction): } \quad \text{MnO}_2(s) + 2\text{H}_2\text{O}(l) + 2\text{e}^- \rightarrow \text{Mn(OH)}_2(s) + 2\text{OH}^-_\text{(aq)}
\]

**Overall (cell) reaction:**

\[
\text{Zn}(s) + \text{MnO}_2(s) + \text{H}_2\text{O}(l) \rightarrow \text{ZnO}(s) + \text{Mn(OH)}_2(s) \quad E_{\text{cell}} = 1.5 \text{ V}
\]

**Zinc and Magnesium Oxide Battery**

\[
\text{Anode (oxidation): } \quad \text{Zn}(s) + 2\text{OH}^-_\text{(aq)} \rightarrow \text{ZnO}(s) + \text{H}_2\text{O}(l) + 2\text{e}^-
\]

\[
\text{Cathode (reduction): } \quad \text{Ag}_2\text{O}(s) + \text{H}_2\text{O}(l) + 2\text{e}^- \rightarrow 2\text{Ag}(s) + 2\text{OH}^-_\text{(aq)}
\]

**Overall (cell) reaction:**

\[
\text{Zn}(s) + \text{Ag}_2\text{O}(s) \rightarrow \text{ZnO}(s) + 2\text{Ag}(s) \quad E_{\text{cell}} = 1.6 \text{ V}
\]

The mercury battery uses HgO as the oxidizing agent instead of Ag₂O and has cell potential of 1.3 V.

**The Lithium Batteries use Silver Vanadium Pentoxide as cathode**

\[
\text{Anode (oxidation): } \quad 3.5\text{Li}(s) \rightarrow 3.5\text{Li}^+ + 3.5\text{e}^-
\]

\[
\text{Cathode (reduction): } \quad \text{AgV}_2\text{O}_5(s) + 3.5\text{Li}^+ + 3.5\text{e}^- \rightarrow \text{Li}_{3.5}\text{V}_2\text{O}_{5.5}
\]

**Overall (cell) reaction:**

\[
\text{AgV}_2\text{O}_5(s) + 3.5\text{Li}(s) \rightarrow \text{Li}_{3.5}\text{V}_2\text{O}_{5.5}
\]

**The most commonly used Battery is Lead-Acid Battery**

\[
\text{Anode (oxidation): } \quad \text{Pb}(s) + \text{HSO}_4^-_\text{(aq)} \rightarrow \text{PbSO}_4(s) + \text{H}^+_\text{(aq)} + 2\text{e}^-
\]

\[
\text{Cathode (reduction): } \quad \text{PbO}_2(s) + 3\text{H}^+_\text{(aq)} + \text{HSO}_4^-_\text{(aq)} + 2\text{e}^- \rightarrow \text{PbSO}_4(s) + 2\text{H}_2\text{O}(l)
\]

**Overall (cell) reaction (discharge):**

\[
\text{PbO}_2(s) + \text{Pb}(s) + \text{H}_2\text{SO}_4_\text{(aq)} \rightarrow 2\text{PbSO}_4(s) + 2\text{H}_2\text{O}(l) \quad E_{\text{cell}} = 2.1 \text{ V}
\]

**Overall (cell) reaction (recharge):**

\[
2\text{PbSO}_4(s) + 2\text{H}_2\text{O}(l) \rightarrow \text{PbO}_2(s) + \text{Pb}(s) + \text{H}_2\text{SO}_4_\text{(aq)}
\]
Nickel Metal Hydride Battery

Anode (oxidation):
\[ \text{MH(s)} + \text{OH}^- (aq) \rightarrow \text{M(s)} + \text{H}_2\text{O(l)} + \text{e}^- \]

Cathode (reduction):
\[ \text{NiO(OH)}(s) + \text{H}_2\text{O(l)} + \text{e}^- \rightarrow \text{Ni(OH)}_2(s) + \text{OH}^- (aq) \]

Overall (cell) reaction:
\[ \text{MH(s)} + \text{NiO(OH)}(s) \rightarrow \text{M(s)} + \text{Ni(OH)}_2(s) \quad E_{\text{cell}} = 1.4 \text{ V} \]

- 

Laptops, Cell-phones most commonly use Lithium ion Batteries

Anode (oxidation):
\[ \text{Li}_x\text{C}_y(s) \rightarrow x\text{Li}^+ + x\text{e}^- + \text{C}_y(s) \]

Cathode (reduction):
\[ \text{Li}_x\text{Mn}_2\text{O}_4(s) + x\text{Li}^+ + x\text{e}^- \rightarrow \text{LiMn}_2\text{O}_4(s) \]

Overall (cell) reaction:
\[ \text{Li}_x\text{C}_y(s) + \text{Li}_x\text{Mn}_2\text{O}_4(s) \rightarrow \text{LiMn}_2\text{O}_4(s) \quad E_{\text{cell}} = 3.7 \text{ V} \]

- 

Hydrogen Fuel Cell

Anode (oxidation):
\[ 2\text{H}_2(g) \rightarrow 4\text{H}^+(aq) + 4\text{e}^- \]

Cathode (reduction):
\[ \text{O}_2(g) + 4\text{H}^+(aq) + 4\text{e}^- \rightarrow 2\text{H}_2\text{O}(g) \]

Overall (cell) reaction:
\[ 2\text{H}_2(g) + \text{O}_2(g) \rightarrow 2\text{H}_2\text{O}(g) \quad E_{\text{cell}} = 1.2 \text{ V} \]

What is the value of the solubility product constant for AgCl?

We could calculate this by breaking the overall reaction into a series of redox reactions as follows:

Anode
\[ \text{Ag(s)} \rightarrow \text{Ag}^+ + \text{e}^- \quad E_{\text{an}} = -0.80 \text{ V} \]

Cathode
\[ \text{AgCl} + \text{e}^- \rightarrow \text{Ag(s)} + \text{Cl}^- \quad E_{\text{red}} = 0.22 \text{ V} \]

Overall (º K_sp reaction)
\[ \text{AgCl} \rightleftharpoons \text{Ag}^+ + \text{Cl}^- \quad E_{\text{cell}} = -0.58 \text{ V (not very spontaneous)} \]

\[
\ln K = \frac{nFE_{\text{cell}}}{RT} = \frac{1\times 96485 \frac{\text{C} \cdot \text{mol}}{\text{e} \cdot \text{V}} \times (-0.58 \text{ V})}{8.3145 \frac{\text{J}}{\text{mol} \cdot \text{K}}} \times 298.15 \text{K} = 22.5744
\]

\[ K = 6.3 \times 10^{-9} \]
Ion Concentrations can also be calculated...

Consider the Platinum-Hydrogen electrode coupled with a copper/copper(II) electrode.

The overall reaction is \( \text{Cu}^{2+} + \text{H}_2 \rightarrow \text{Cu} + 2\text{H}^+ \).

The Nernst equation is

\[
E = E^\circ - \frac{0.0257 \text{ V}}{2} \ln \left( \frac{a(\text{H}^+)^2}{a(\text{Cu}^{2+}) \times a(\text{H}_2)} \right)
\]

\[
E = E^\circ - \frac{0.0257 \text{ V}}{2} \ln \left( \frac{a(\text{H}^+)^2}{a(\text{Cu}^{2+}) \times a(\text{H}_2)} \right)
\]

If \([\text{Cu}^{2+}] = 1 \text{ M and } P(\text{H}_2) = 1 \text{ bar (both their activities are 1)}\) then

\[
E = E^\circ - 0.0257 \text{ V ln } [\text{H}^+] \]

(Note: \( \ln Y = 2.303 \times \log Y \) and \( -0.257 \times 2.303 = 0.05917 \))

\[
E = E^\circ + 0.05917 \text{ V } (-\log [\text{H}^+])
\]

\[
E = E^\circ + 0.05917 \text{ V } \times \text{pH}
\]

We see here that the cell potential is a function of pH. The probes in pH meters are set up this way. A complete electrochemical cell is contained within the probe casing. All chemicals are at standard conditions and a porous glass membrane allows only H+ ions to pass through.
Electrochemical Cell

Let us always remember the basics. If something, say M, loses electrons, it is getting oxidized. Metals are basic. Metals loose electrons and gets oxidized. The metals which lose electron very easily, get oxidized very easily. So these are better reducing agent.

Adding of Oxygen, or Halogens (F, Cl, Br, I) or other p-Block elements which accept electrons, is oxidation. So if Potassium (K) gets Oxygen (attaches to O) to form K₂O or KO₂ (Superoxide), K is getting oxidized. If Cs attaches with F then Cs is getting oxidized, while F is being reduced. Cs is a reducing agent as it is loosing electrons. F is an oxidizing agent as it is gaining electrons.

Adding of Hydrogen is reduction. So if Ethene receives Hydrogen to become Ethane, then Ethene is being reduced. Raney Nickel is Nickel with molecular level pores. Raney Ni with HCl gives Hydrogen atoms, (some teachers say this as Nascent Hydrogen) locally, meaning near the molecule where the reaction is needed. So Raney Ni + HCl is a good reducing agent. Same for Zinc dust + HCl. But then what is Birch reduction?

As my experience of teaching more than 25 years, almost every student understands these. Knowing this is essential but does not ensure success in chemistry. There are many kinds of details of oxidizing agent or Reducing agent that vary from reactions / situations.
A simple binary ionic compound contains only two elements - a metal and a non-metal. When the ionic compound is in the molten state, the locked ions within the ionic structure will be free to move about (conduct electricity).

A typical setup for electrolysis of molten compounds is shown below:

![Electrolysis Setup Diagram]

The metallic ions (cations - $M^{n+}$) will be discharged at the cathode to form a metal atom. The metallic ions are REDUCED to metal at the cathode. (because they gain electrons)

$$M^{n+} + ne^- \rightarrow M$$

The non-metallic ions (anions - $N^{n-}$) will be discharged at the anode to form a non-metallic atom. The non-metallic ions are OXIDISED to non-metallic atom at the anode. (because they lose electrons)

$$N^{n-} \rightarrow N + ne^-$$

**Case Study: Electrolysis of molten sodium chloride, NaCl**

**Electrodes:** Carbon rods (Graphite)

**Electrolytes:** Molten sodium chloride

**Ions present in electrolytes:** Sodium ions ($Na^+$), Chloride ions ($Cl^-$)

**Reaction at the CATHODE:**

- $Na^+$ ions are attracted to the cathode.
- Each $Na^+$ ion gains one electron from the cathode to form one sodium atom.
- Molten sodium is formed at the anode.
- equation: $Na^+(l) + e^- \rightarrow Na(s)$ (Reduction)

**Reaction at the ANODE:**

- $Cl^-$ ions are attracted to the anode.
Each \( \text{Cl}^- \) ion loses one electron to the anode to form one chlorine atom.

Chlorine gas is liberated at anode.

Equation: \( 2\text{Cl}^-(l) \rightarrow \text{Cl}_2(g) + 2e^- \) (Oxidation)

**Overall reaction:**

- Every 2 moles of sodium chloride produce 2 moles of sodium metal and 1 mole of chlorine gas
- \( 2\text{NaCl}(l) \rightarrow 2\text{Na}(s) + \text{Cl}_2(g) \)

Case Study: Electrolysis of molten magnesium oxide, \( \text{MgO} \)

**Electrodes:** Carbon rods (Graphite)

**Electrolyte:** Molten magnesium oxide

**Ions present in electrolyte:** Magnesium ions (\( \text{Mg}^{2+} \)), Oxygen ions (\( \text{O}_2^- \))

**Reaction at the CATHODE:**

- \( \text{Mg}^{2+} \) ions are attracted to the cathode.
- Each \( \text{Mg}^{2+} \) ion gains two electrons from the cathode to form one magnesium atom.
- \( \text{Mg}^{2+}(l) + 2e^- \rightarrow \text{Mg}(s) \)

**Reaction at the ANODE:**

- \( \text{O}_2^- \) ions are attracted to the anode.
- Each \( \text{O}_2^- \) ion loses two electrons to the anode to form one oxygen atom.
- \( 2\text{O}_2^-(l) \rightarrow \text{O}_2(g) + 4e^- \)

**Overall reaction:**

- Every two moles of magnesium oxide produce two moles of magnesium metal and one mole of oxygen gas.
- \( 2\text{MgO}(l) \rightarrow 2\text{Mg}(s) + \text{O}_2(g) \)
Gyan Question:

Determine the standard potential of the Cu^{2+}/Cu^{+} cell from two other standard cell potentials.

\( E^\circ(Cu^{2+}/Cu) = +0.340 \text{V} \) and \( E^\circ(Cu^{+}/Cu) = +0.522 \text{V} \).

Since Hess' Law allows us to add Gibbs energies for the reactions to arrive at the Gibbs energy of the desired reaction, we should go via Gibbs energies. Convert our standard potentials into Gibbs energies, perform the addition and then convert back to a standard potential.

The two reactions that will occur are:

(a) \( Cu(s) \rightleftharpoons Cu^{2+}(aq) + 2 e^- \)

\[ B^\circ = -0.340 \text{V} \quad \Delta_\text{f}G = -2(-0.340 \text{V}) = 0.680 \text{kJ/mol} \]

(b) \( Cu^{+}(aq) + e^- \rightleftharpoons Cu(s) \)

\[ B^\circ = +0.522 \text{V} \quad \Delta_\text{f}G = -(+0.522 \text{V}) = -0.522 \text{kJ/mol} \]

(c) \( Cu^{2+}(aq) + e^- \rightleftharpoons Cu^{+}(aq) \)

\[ B^\circ = +0.158 \text{V} \quad \Delta_\text{f}G = +0.158 \text{V} = 15.2 \text{kJ/mol} \]

Now, since \( B^\circ = -n \Delta_\text{f}G/F = -0.158\text{V} \) (\( n = 1 \) in the final equation)

We cannot simply add the electrode potentials in this case because, the final reaction is still a half-reaction.

If the final reaction is a complete cell reaction, with no electrons remaining, then we can skip to just adding the cell potentials because the factor \( v \) would be the same in all equations and hence, cancel out.
(1) \( \text{Cu}^+ (aq) + e^- \rightarrow \text{Cu} (s) \quad E^\circ = 0.521 \text{ V} \)

(2) \( \text{Cu}^{2+} (aq) + e^- \rightarrow \text{Cu}^+ (aq) \quad E^\circ = 0.153 \text{ V} \)

(1) is reduction and drives (2) as oxidation.

\[ 2 \text{Cu}^+ (aq) \rightarrow \text{Cu}^{2+} (aq) + \text{Cu} (s) \quad E^\circ_{\text{cell}} = 0.521 - 0.153 = 0.368 \text{ V} (>0) \]

Which means \( E^\circ_{m_n} > 0 \) and \( \Delta G^\circ_{m_n} < 0 \) spontaneous. Yes, it disproportionates.

\[
\begin{align*}
\text{Cu}^+ (aq) + e^- &= \text{Cu} (s) & E^\circ &= 0.521 \text{ V} \\
\text{Cu}^+ (aq) &= \text{Cu}^{2+} (aq) + e^- & E^\circ &= -0.153 \text{ V} \\
2\text{Cu}^+ (aq) &= \text{Cu}^{2+} (aq) + \text{Cu} (s) & E^\circ &= 0.368 \text{ V}
\end{align*}
\]

We have value of \( K \)

\[
\ln K = \frac{nFE^\circ}{RT}
\]

\[
= \frac{(1)(96485)(0.368)}{(8.314)(298.15)} = 14.3.
\]

\[ K = \frac{a_{\text{Cu}^{2+}}}{a_{\text{Cu}^+}^2} = 1.62 \times 10^6. \]

The large \( K \) indicates that the point of equilibrium in the reaction is far toward products, and disproportionation will occur until \( a_{\text{Cu}^+} \) becomes very small.
Discuss Disproportion of Iron

Decide whether $\text{Fe}^{2+} (aq)$ in its standard state at $25^\circ\text{C}$ is stable without disproportionation.

In Appendix E we find:

1. $\text{Fe}^{3+} (aq) + e^- \rightarrow \text{Fe}^{2+} (aq) \quad E^\circ = 0.771 \text{V}$
2. $\text{Fe}^{2+} (aq) + 2e^- \rightarrow \text{Fe} (s) \quad E^\circ = -0.447 \text{V}$

In order for $\text{Fe}^{2+} (aq)$ to disproportionate reaction (1) would have to be driven backwards (oxidation): $3\text{Fe}^{2+} \rightarrow \text{Fe} (s) + 2\text{Fe}^{3+} (aq)$

And $E^\circ = -0.447 - (0.771) = -1.218 \text{V} (< 0)$ and $\Delta G^\circ > 0$ (non-spontaneous) and therefore stable against disproportionation.

0.01 M Solution of HCN has a specific resistance of $x$ ohm cm its equivalent conduction would be ?

(a) $10x$ (b) $100x$ (c) $10^5/x$ (d) $10^6/x$

\[
\begin{align*}
C &= 0.01 \text{ M} = 0.01 \text{ N (for HCN)} \\
\therefore V &= \frac{1}{0.01} = 100 \text{ litres/eq} \\
\text{or,} \quad V &= 100 \times 1000 \text{ cm}^3 \\
kV &= \frac{1}{\text{Specific resistance}} \\
&= \frac{1}{x} \text{ ohm}^{-1} \text{ cm}^{-1} \\
\text{Equivalent conductance} \quad \lambda V &= kV \times V \\
&= \frac{1}{x} \times 100 \times 1000 \\
&= \frac{10^5}{x} \text{ ohm}^{-1} \text{ cm}^2
\end{align*}
\]
Recall again

### Difference Between Electrolytic and Galvanic Cells

<table>
<thead>
<tr>
<th>Electrolytic Cell</th>
<th>Galvanic Cell</th>
</tr>
</thead>
<tbody>
<tr>
<td>It requires a source of external energy.</td>
<td>It is a source of energy.</td>
</tr>
<tr>
<td>It converts electrical energy into chemical energy.</td>
<td>Converts chemical energy into electrical energy.</td>
</tr>
<tr>
<td>Has cathode as the negative electrode.</td>
<td>Has cathode as positive electrode.</td>
</tr>
<tr>
<td>Has anode as the positive electrode.</td>
<td>Has anode as negative electrode.</td>
</tr>
<tr>
<td>Involves oxidation at anode and reduction at cathode.</td>
<td>Involves oxidation at anode and reduction at cathode.</td>
</tr>
</tbody>
</table>

**Faraday’s second law of electrolysis**—It states that when same quantity of electricity is passed through different electrolytes, the amount of different substances deposited at the electrodes is directly proportional to their equivalent weights.

For example, if same quantity of electricity is passed through three voltameters or coulometers containing a dilute solution of $\text{H}_2\text{SO}_4$, a solution of $\text{CuSO}_4$ and a solution of $\text{AgNO}_3$, it has been found that one coulomb of electricity liberates $1.036 \times 10^{-3}$ g. of hydrogen, $3.292 \times 10^{-4}$ g. of copper and $1.118 \times 10^{-3}$ g. of silver. The weight of element obtained in each case is proportional to the equivalent weight. Thus,

\[
\begin{align*}
\text{Weight of } \text{H}_2 &= \frac{\text{Eq. Wt. of } \text{H}_2}{\text{Weight of } \text{Cu}} = \frac{1.008}{31.78} = 0.0318 \text{ g.} \\
\text{Weight of } \text{Cu} &= \frac{\text{Eq. Wt. of } \text{Cu}}{\text{Weight of } \text{Ag}} = \frac{31.78}{107.88} = 0.0294 \text{ g.}
\end{align*}
\]

The quantity of electricity required to liberate one gm equivalent of each element is calculated as,

- **Hydrogen**: $[1.008/1.036 \times 10^{-3}] = 96515 \text{ Coulombs}$
- **Copper**: $[31.78/3.292 \times 10^{-4}] = 96567 \text{ Coulombs}$
- **Silver**: $[107.88/1.118 \times 10^{-3}] = 96494 \text{ Coulombs}$

**Faraday**—We know that an $\text{Ag}^+$ ion takes up one electron from the cathode and gets converted into $\text{Ag}$ atom.

\[\text{Ag}^+ + e^- \rightarrow \text{Ag}\]

The equivalent weight of $\text{Ag}$ (107.88) is equal to its atomic weight. There are $6.023 \times 10^{23}$ (Avogadro’s number) atoms in one atomic weight or one mole. Thus 1 mole of $\text{Ag}$ will be deposited when 1 mole of $\text{Ag}^+$ ion takes up one mole of electrons. The charge on each electron is $1.603 \times 10^{-19}$ Coulombs. The total charge of 1 mole of electrons is therefore $1.603 \times 10^{-19} \times 6.023 \times 10^{23} = 96500$ Coulombs. Hence 96,500 Coulombs of electricity is required to deposit one gm equivalent of silver. 96,500 Coulombs of electricity is called one Faraday. The Faraday (F) is also the quantity of charge carried by one mole of electrons, because

\[F = e \times N = 1.603 \times 10^{-19} \times 6.023 \times 10^{23} = 96500 \text{ Coulombs.}\]

This fact can be used in calculating the electrochemical equivalents. For example,

\[\text{Electrochemical equivalent of silver} = \frac{\text{Eq. Wt. of Ag}}{96500} = \frac{107.88}{96500} = 1.118 \times 10^{-3} \text{ g.}\]

\[\text{Electrochemical equivalent of copper} = \frac{\text{Eq. Wt. of Cu}}{96500} = \frac{31.78}{96500} = 3.292 \times 10^{-4} \text{ g.}\]

Hence Faraday (F) is the quantity of current in Coulombs required to deposit one gm equivalent of any substance. Passage of one Faraday of electricity means flow of $6.023 \times 10^{23}$ electrons.

If one Faraday of electricity is passed through fused $\text{NaCl}$, 23.0g. or one mole or one gm. equivalent of sodium is deposited at the cathode and 35.5 g. or 0.5 mole or one gm equivalent of chlorine is liberated at the anode.

\[\text{Na}^+ + e^- \rightarrow \text{Na} \quad \text{(One electron is taken up)}\]
If one Faraday of electricity is passed through CuCl₂ solution, one gm. equivalent weight of copper ions is discharged at the cathode. This is only 1/2 mole of copper, since copper is bivalent.

\[ \text{Cu}^{2+} + 2e^- \rightarrow \text{Cu} \quad \text{(Two electrons are taken up)} \]

If one Faraday of electricity is passed through AlCl₃ solution, one gm. equivalent weight of Al³⁺ ions is discharged at the cathode. This is only 1/3 mole of Al³⁺ ions, because Al is trivalent.

\[ \text{Al}^{3+} + 3e^- \rightarrow \text{Al} \quad \text{(Three electrons are taken up)} \]

In other words, for depositing one mole of Na atoms, one mole of copper atoms and one mole of aluminium atoms, we need one, two and three moles of electrons respectively. In general, to deposit one mole of atoms of any metal of valency Z, we need Z moles of electrons. Thus,

\[ M = eZF \]

where \( M \) is the mass of one mole. Now we know \( W = eQ \) where \( Q = \text{coul} \times t \). Thus,

\[ \frac{W}{M} = \frac{eQ}{eZF} = \frac{Q}{ZF} \]

\[ \text{Thus the equivalent weight (E) of copper is 31.8 g. In general, equivalent weight (E) of an element is,} \]

\[ E = \frac{\text{Gram Atomic Weight of Element}}{Z} \]

\[ \text{Applications of electrolysis} – \text{There are various industrial applications of electrolysis. For example, (a) Electrometallurgy – The metals like Na, K, Mg, Al etc. are obtained by the electrolysis of fused electrolytes. (b) Electrefining of metals – A large number of metals, such as Cu, Ag, Au, Al, Sn etc. are purified by electrolysis. (c) Manufacture of compounds – Electrolysis has also been used in the large scale production of various compounds such as NaOH, KOH, KCl, Na₂CO₃, H₂O etc. (d) Manufacture of non-metals – Electrolysis has been employed also in the manufacture of H₂, Cl₂, F₂ etc. (e) Electroplating – Deposition of precious metals like Ag, Au, Pt etc. on less expensive metals like Cu is known as electroplating. This is also carried out by electrolysis. (f) Preparation of alloys and electrotyping – Electrolysis is very useful in the preparation of alloys and also in electrotyping.} \]
**Electrical conductivity** – Electrical conductivity is the conductance in mhos of a solution containing one gm equivalent of solute when placed between two sufficiently large electrodes which are one cm apart. It is denoted by \( \lambda \). The equivalent conductivity is related to the specific conductivity by the expression

\[
\lambda = K \times V
\]

where \( V \) is the volume in c.c. of the solution containing one gm equivalent of an electrolyte.

**Molecular conductivity** – It is the conductance in mhos of a solution containing one gm mole of the solute when placed between two sufficiently large electrodes which are exactly one cm apart. It is denoted by \( \lambda_m \) and is numerically equivalent to the product of sp. conductivity and the volume \( V \) in c.c. of the solution containing one gm. mole of the electrolyte. Thus

\[
\lambda_m = K \times V
\]

**Cell constant** – The conductance of a solution is measured by a cell, known as conductance cell. The electrodes in this cell may not be exactly one cm apart and may not have an area of one sq. cm. Thus the value of observed conductance will not be equal to the specific conductance, but a value proportional to it. It is therefore essential to calculate a factor for the conductance cell, called the cell constant, which when multiplied by the observed conductance gives the value of specific conductance. The value of the conductance will, therefore, have to be multiplied by a certain factor in order to get specific conductance. This factor is known as cell constant. We know

\[
R = \frac{p}{x/a} = \frac{p}{x/a} \text{ cell constant}
\]

Thus \( x \) (cell constant) = \( \frac{R}{p} = \frac{V}{\text{observed conductivity}} \)

So, Specific Conductivity = Cell constant \( x \) × Obs. Conductivity

In order to determine the cell constant, the conductance cell is filled with exact N/50 solution of chemically pure KCl. The specific conductance of this solution at 25°C has been found to be 0.002765 mhos. The cell is placed in a thermostat at 25°C and the resistance and hence the conductance of the solution is determined. Thus

\[
\text{Cell Constant} = \frac{0.002765}{\text{Observed conductivity}}
\]

The cell constant is expressed in cm\(^{-1}\).

**The important conditions for the cell to be reversible are:**

(a) If the external e.m.f. or potential is equal to the e.m.f. of the cell, no current flows through the cell and no chemical reaction occurs at all.

(b) If the external e.m.f. or potential is slightly smaller than the e.m.f. of the cell, a small current flows out of the cell and a small amount of chemical reaction takes place in the cell.

(c) If the external e.m.f. or potential is slightly more than the e.m.f. of the cell, current flows into the cell and cell reaction is reversed.

(d) A cell which does not fulfil the above conditions is said to be irreversible.
Effect of dilution on conductivity – (a) The conductivity of the solution increases with dilution, because it depends upon the number of ions present in the solution. On dilution, the degree of ionisation of electrolyte increases and more ions are produced in the solution. The greater the number of ions, the greater is the conductivity.

(b) The specific conductivity decreases on dilution. This is due to the fact that although the total number of ions increases with dilution owing to increased dissociation, the number of current carrying particles or ions per c.c. decreases. This causes a decrease in specific conductivity.

(c) Equivalent and molecular conductivity both increase with dilution. These are the products of specific conductivity ($k_\nu$) and volume $V$ of the solution containing one gm equivalent and one gm mole of the solute dissolved respectively. On dilution, more and more electrolyte ions (degree of ionisation increases). When whole of the substance has ionised further dilution produces little or no change in the equivalent and molecular conductivities. This limiting value of equivalent conductivity (or molecular conductivity) is called the equivalent conductivity at infinite dilution. The ratio of equivalent conductivity at any dilution to that at infinite dilution ($\lambda_\infty$) is called the conductivity ratio ($\alpha$). Thus $\alpha = \lambda_\nu / \lambda_\infty$.

It should be noted that increase in equivalent conductivity in the case of strong electrolytes like KCl, NaCl, HCl etc. is not so large (because these are already highly ionised) as in case of weak electrolytes such as CH$_3$COOH, NH$_3$OH etc. (because they are slightly ionised and their ionisation increases with dilution).

Migration of ions – Electricity is carried through the solution of an electrolyte by migration of ions. The ions are present independently in solution and when a current of electricity is passed through the solution, the ions present in it move towards the oppositely charged electrodes. Thus,

(a) The ions move towards the oppositely charged electrodes at different speeds.

(b) During electrolysis, the ions are discharged or liberated in equivalent amounts at the two electrodes, no matter what their relative speed is.

(c) Concentration of the electrolyte changes around the electrodes as a result of difference in the speed of the ions.

(d) Loss of concentration around any electrode is proportional to the speed of the ion that moves away from the electrode. Thus,

\[
\text{Fall Round Anode} = \text{Speed of Cation} \\
\text{Fall Round Cathode} = \text{Speed of Anion}
\]

This relation is valid on the assumption that the discharged ions do not react with the atoms or material of the electrodes. But in many cases, they do combine with the electrode and in such cases, the concentration around such an electrode shows an increase, rather than a decrease. Electrolysis of AgNO$_3$ solution using silver electrodes is an example of such a case. On electrolysis, it has been observed that instead of fall, the concentration of AgNO$_3$ around the anode increases, because every NO$_3$ ion that reaches at the anode dissolves from it one Ag$^+$ ion to form AgNO$_3$.

Transport number or transference number – The fraction of the current carried by an ion is termed as its transport number or transference number. If $n_a$ and $n_c$ are the transport numbers of anion and cation respectively then

\[
n_a = \frac{\text{Current Carried by Anion}}{\text{Total Current Carried}} \quad \text{and} \quad n_c = \frac{\text{Current Carried by Cation}}{\text{Total Current Carried}}
\]

Concentration has marked effect on the transport number, especially in the case of some partially dissociated electrolytes such as ZnI$_2$, CdI$_2$ etc. For example, concentration has a marked effect on the transport number of cadmium ion in CdI$_2$. The value decreases from 0.49 at low concentration to almost zero at higher concentrations and the values become negative at still higher concentrations. The transport numbers of cadmium ions in 0.01N, 0.05N, 0.02N and 0.50N CdI$_2$ at 25°C are 0.449, 0.402, 0.131, and 0.005 respectively. This abnormal behaviour can be explained by assuming:

(a) That in very dilute solution, CdI$_2$ ionises to Cd$^{2+}$ ions and I$^-$ ions and thus Cd$^{2+}$ shows the usual transport number. CdI$_2$ $\rightarrow$ Cd$^{2+}$ + 2I$^-$. (b) That with increase in concentration, CdI$_2$ takes on I$^-$ ions and forms complex, CdI$_2$ + 2I$^-$ $\rightarrow$ [CdI$_4$]$^{2-}$.
Thus the concentrated solution contains both very charged Cd\(^{2+}\) ions and negatively charged [CdI\(_4\)]\(^{-}\) ions. Hence some of the cadmium forms a part of the anion and migrates towards the anode. Therefore, fall of concentration round the anode is not as much as it should be. Since relative amounts of two ions depend upon the concentration of electrolyte CdI\(_2\), higher the concentration of CdI\(_2\) greater would be the proportion of the complex [CdI\(_4\)]\(^{-}\) ion. Thus, if anion [CdI\(_4\)]\(^{-}\) moves faster than the cation Cd\(^{2+}\), there will be an increase in the concentration round the anode, instead of decrease and thus explains the negative value for the transport number of Cd\(^{2+}\) ion at much higher concentration.

There is a slight variation of transport number with temperature. For 1:1 electrolytes, it has been observed that if transport number is greater than 0.5, it decreases with an increase in temperature, but if it is less than 0.5, it indicates an increase with an increase in temperature. Thus transport numbers tend to approach a value of 0.5 as the temperature is increased. This is probably due to the fact that at higher temperature the ionic conductance as well as the speed tend to approach the same limiting value, resulting in equal contribution from the two ions.

(A) Consider the reaction, (Hg\(^{2+}\) + Ag → Ag\(^{+}\) + Hg)
Comment on the feasibility of the reaction under the different concentrations of ions given below
(i) \([\text{Hg}^{2+}] = 10^{-4} \text{ M}; [\text{Ag}^{+}] = 10^{-4} \text{ M}\)
(ii) \([\text{Ag}^{+}] = 10^{-4} \text{ M}; [\text{Hg}^{2+}] = 10^{-4} \text{ M}\)

Given: \(E^0_{\text{Ag}^{+}/\text{Ag}} = 0.80 \text{ V}; E^0_{\text{Hg}^{2+}/\text{Hg}} = 0.79 \text{ V}\).

(B) For a process, Water \(\rightleftharpoons\) ice, the enthalpy change \(\Delta H = -500 \text{ J/mol}; T = 100 \text{ K}; P = 1 \text{ atm}\).
What will be the values of
(i) Entropy change \(\Delta S\)
(ii) Free energy change \(\Delta G\) ?

Sol. (A) \(E^0_{\text{cell}} = E^0_{\text{C}} - E^0_{\text{A}} = 0.79 - 0.80 = -0.01 \text{ V}\)
The net cell reaction is
\(\text{Hg}^{2+} + 2\text{Ag} \rightarrow 2\text{Ag}^{+} + \text{Hg}\)
Anode is: \(\text{2Ag} \rightarrow 2\text{Ag}^{+} + 2e^-\)
Cathode: \(\text{Hg}^{2+} + 2e^- \rightarrow \text{Hg}\)

Case-1: \(E_{\text{cell}} = -0.01 + \frac{0.059}{2} \log \frac{10^{-4}}{10^{-2}} = -0.069 \text{ V}\)

So Reaction not feasible as \(E_{\text{cell}}\) is negative
Case-II: \( E_{cell} = -0.01 + \frac{0.059}{2} \log \frac{10^{-2}}{10^{-4}} = 0.049 \, \text{V} \)

\[ \therefore \text{Feasible because } E_{cell} \text{ is } + \text{ve.} \]

(B) The given process is

\[ \text{H}_2\text{O}(l) \iff \text{H}_2\text{O}(s) \]

(i) \( \Delta S = \frac{\Delta H}{T} = \frac{-500}{100} = -5 \text{J mol}^{-1} \text{K}^{-1} \)

(ii) At equilibrium \( \Delta G = 0 \).

**Kohlrausch law** – This law states that the value of equivalent conductivity at infinite dilution (\( \lambda_{\infty} \)) may be regarded as made up of the sum of two independent factors, one characteristic of cation and other to anion. Thus,

\[ \lambda_{\infty} = \lambda_+ + \lambda_- \]  \hspace{1cm} \text{...(12)}

If we consider a salt \( \text{NaCl} \), its equivalent conductivity at infinite dilution is the sum of two terms, one due to \( \text{Na}^+ \) ion (\( \lambda_{\text{Na}^+} \)) and other due to \( \text{Cl}^- \) ion (\( \lambda_{\text{Cl}^-} \)). These terms are known as ionic mobilities of anion and cation. The ratio of ionic mobilities is equal to their transport numbers. Thus,

\[ \frac{\lambda_+}{\lambda_-} = \frac{n_+}{n_-} = \frac{n_+}{(1 - n_+)} \]  \hspace{1cm} \text{(Since } n_+ + n_- = 1 \text{)} \hspace{1cm} \text{...(13)}

The **absolute velocity** of an ion may be defined as the velocity of an ion in \( \text{cm} \) per sec. when it moves under an electric field due to two electrodes placed one \( \text{cm.} \) apart and having a potential difference of one volt between them (i.e., under unit potential gradient).

Hence unit of absolute velocity is \( \text{cm per sec. (volts per cm)}, \text{ i.e., } \text{cm}^2 \text{ per sec. per volt.} \) The absolute velocity is numerically equal to the ionic mobility divided by one Faraday (96450 Coulombs).

Thus,

\[ U = \frac{\lambda_+}{96500} = \frac{\lambda_-}{96500} \]

From equation (13) We have

\[ \frac{\lambda_+}{\lambda_-} = \frac{n_+}{1 - n_+} \text{ or } \lambda_+ - n_+ \lambda_- = n_+ \lambda_+ \]

\[ \lambda_+ = n_+ (\lambda_+ + \lambda_-) = n_+ \times \lambda_{\infty} \]  \hspace{1cm} \text{(since } \lambda_+ + \lambda_- = \lambda_{\infty} \text{)} \hspace{1cm} \text{...(14)}

Hence conductance of an ion is obtained by multiplying the equivalent conductivity at infinite dilution of any strong electrolyte containing that ion by its transport number. Similarly,

\[ \lambda_+ = n_+ \times \lambda_{\infty} \]

**Applications of Kohlrausch law** – (1) In the determination of ionic mobilities and absolute velocity of ions, (2) Calculation of equivalent conductivities of weak electrolytes at infinite dilution. (3) Determination of degree of dissociation, since, \( \alpha, \) the degree of dissociation of an electrolyte at the dilution \( V \) is given by \( \alpha = \lambda V / \lambda_{\infty} \) (where \( \lambda_{\infty} = \lambda + \lambda_- \))
Gyan Question:

Q: What is carborundum? Does it conduct electricity?

Ans:

Silicon carbide, also known as carborundum, is a compound of silicon and carbon with chemical formula SiC. It occurs in nature as the extremely rare mineral moissanite. Silicon carbide powder has been mass-produced since 1893 for use as an abrasive.

Grains of silicon carbide can be bonded together by sintering to form very hard ceramics that are widely used in applications requiring high endurance, such as car brakes, car clutches and ceramic plates in bulletproof vests. Electronic applications of silicon carbide as light-emitting diodes (LEDs) and detectors in early radios were first demonstrated around 1907, and today SiC is widely used in high-temperature/high-voltage semiconductor electronics.
Relative tendencies of electrodes to liberate electrons – When two electrodes are connected to from a cell, a competition sets in between the two electrodes to liberate electrons. For example, if the cell consists of zinc and copper electrodes, as in Daniell cell, the electrons flow from negative zinc electrode to positive copper electrode, because zinc has a greater tendency to liberate electrons than copper. If the cell is constituted with copper and silver electrodes, then copper has a greater tendency to liberate electrons than silver. This indicates that copper is negative electrode and silver is positive electrode and electrons flow from copper to silver electrode. If a cell is set up with Zn and H₂(g) electrodes, then zinc has a greater tendency to liberate electrons than the H₂(g) electrode. This indicates that electrons will flow from negative zinc electrode to positive H₂(g) electrode. If the cell is constituted with copper and H₂(g) electrodes, then H₂(g) electrode has a greater tendency to liberate electrons than the copper electrode. This indicates that electrons will flow from negative H₂(g) electrode to positive copper electrode. Thus in the above cases, the tendency of the electrodes to release electrons decreases in the order Zn > H₂(g) > Cu > Ag.

It should be noted that electrode at which electrons are released is written on the left, while that at which electrons are taken up is written on the right. For example,

Cu || Cu²⁺ || Ag⁺ || Ag  Zn || Zn²⁺ || H⁺, H₂(g)

AgNO₃ solution turns blue when a piece of copper is placed in it, because copper metal can be oxidised by Ag⁺ ions. Ag cannot be oxidised by Cu²⁺ ions.

Also remember that the metal which goes into the solution in the form of its salt is undergoing oxidation and thus will act as negative electrode (anode), while the other metal acts as positive electrode (cathode). For example, in the reaction Zn + H₂SO₄ → ZnSO₄ + H₂, the Zn is oxidised to ZnSO₄, hence zinc will be negative electrode or anode and the cell will be constructed as:

Zn || ZnSO₄ || H₂SO₄ || H₂, Pt

Role of salt bridge – Most often salt bridges, which are U-shaped tubes are made with agar-agar gel containing KCl or KNO₃. Sometimes NH₄NO₃ is also used. The reason of using these salts is that the ions of these salts, e.g., KCl (K⁺ ion and Cl⁻) have almost the same speed as well as almost same transport numbers. If a salt bridge is not used, say, in the Daniell cell, the electrons released by the zinc electrode (Zn → Zn²⁺ + 2e⁻) can flow to the copper electrode through the external circuit. These electrons neutralise or reduce some Cu²⁺ ions of the CuSO₄ solution (Cu²⁺ + 2e⁻ → Cu). As a result, copper sulphate solution acquires a negative charge for a short while, due to discharge of positively charged Cu²⁺ ions. In the mean time zinc sulphate solution becomes positively charged because of the formation of Zn²⁺ ions in the zinc electrode. Hence there will be accumulation of charges near the two electrodes, which prevents further flow of current. Hence the electrolychemical change stops and current drops to zero.

The salt bridge provides a passage for the flow of charges in the internal circuit and thus prevents the accumulation of charges. In the presence of salt bridge, the charges present flow towards the oppositely charged electrodes. The –ve ions move from the copper electrode end through the CuSO₄ solution, salt bridge and zinc sulphate solution to the zinc rod. The positive ions move in the reverse direction, i.e., zinc rod to copper rod. Hence flow of electricity is maintained when the salt bridge is used in the Galvanic cell.

To calculate the \( \mathbf{E}^\mathbf{\text{CELL}} \) of Daniell Cell:

\[
\text{Zn || Zn}^{2+} || \text{Cu}^{2+} || \text{Cu}
\]

We know \( \mathbf{E}^\mathbf{\text{CELL}} = \mathbf{E}^\mathbf{\text{RHE}} - \mathbf{E}^\mathbf{\text{LHE}} \) (Both Reduction potentials)

Reaction at LHE

\[
\text{Zn} \rightleftharpoons \text{Zn}^{2+} + 2e^- \quad \text{(Oxidation)}
\]

Reaction at RHE

\[
\text{Cu}^{2+} + 2e^- \rightarrow \text{Cu} \quad \text{(Reduction)}
\]

Now \( \mathbf{E}^\mathbf{\text{LHE}} = \mathbf{E}^\mathbf{\text{Zn}^{2+}, \text{Zn} = -0.76V} \) and \( \mathbf{E}^\mathbf{\text{RHE}} = \mathbf{E}^\mathbf{\text{Cu}^{2+}, \text{Cu} = +0.34V} \). Thus

\[
\mathbf{E}^\mathbf{\text{CELL}} = \mathbf{E}^\mathbf{\text{RHE}} - \mathbf{E}^\mathbf{\text{LHE}} = \mathbf{E}^\mathbf{\text{Cu}^{2+}, \text{Cu} - E^\text{Zn}^{2+}, \text{Zn} = 0.34 V - (-0.76V) = 1.10\text{Volts}}.
\]

Since \( \mathbf{E}^\mathbf{\text{CELL}} \) is positive, the cell reaction, Zn + Cu²⁺ \( \rightarrow \) Zn²⁺ + Cu is spontaneous.
To calculate $E^{\circ}_{\text{CELL}}$ of the cell $\text{Zn} | \text{Zn}^{2+} \ || \text{Pb}^{2+} | \text{Pb}$

Reaction at LHE: $\text{Zn} \rightarrow \text{Zn}^{2+} + 2e^-$ (Oxidation)

Reaction at RHE: $\text{Pb}^{2+} + 2e^- \rightarrow \text{Pb}$ (Reduction)

Cell Reaction: $\text{Zn} + \text{Pb}^{2+} \rightarrow \text{Zn}^{2+} + \text{Pb}$

$E^{\circ}_{\text{CELL}} = E^{\circ}_{\text{LHE}} - E^{\circ}_{\text{RHE}} = E^o_{(\text{Zn}^{2+} | \text{Zn})} - E^o_{(\text{Cu}^{2+} | \text{Cu})}$

$E^{\circ}_{\text{CELL}} = -0.130 - (-0.76) = 0.640 \text{ V}.$

As the $E^{\circ}_{\text{CELL}}$ is positive, the above reaction is spontaneous and thus cell is a source of electricity.

To calculate $E^{\circ}_{\text{CELL}}$ of the cell $\text{Cu} | \text{Cu}^{2+} \ || \text{Zn}^{2+} | \text{Zn}$

Reaction at LHE: $\text{Cu} \rightarrow \text{Cu}^{2+} + 2e^-$ (Oxidation)

Reaction at RHE: $\text{Zn}^{2+} + 2e^- \rightarrow \text{Zn}$ (Reduction)

$E^{\circ}_{\text{CELL}} = E^{\circ}_{\text{RHE}} - E^{\circ}_{\text{LHE}} = E^o_{(\text{Zn}^{2+} | \text{Zn})} - E^o_{(\text{Cu}^{2+} | \text{Cu})} = 0.8 - 0.34 = 0.46 \text{ V}.$

Since the $E^{\circ}_{\text{CELL}}$ is positive, the cell reaction is spontaneous.

Displacement of metals – The reduction potentials of electrodes given in table 1 are very useful in predicting whether a metal can displace another from its salt solution. When a metal $M_1$ displaces another metal $M_2$ from its salt solution, $M_1$ gets oxidised as $M_1 \rightarrow M_1^{2+} + 2e^-$ and $M_2$ is reduced as $M_2^{2+} + 2e^- \rightarrow M_2$. If $M_1$ has the higher oxidation potential than that of $M_2$, then $M_1$ has a greater tendency to get oxidised and hence $M_1$ can displace $M_2$ from its salt solution. For example, zinc displaces silver from AgNO$_3$ solution, because standard oxidation potential of zinc (+0.76V) is greater than that of silver (−0.80V). Similarly aluminium can displace nickel from nickel salt solution because oxidation potential of Al (Al$^{3+}$ (+1.66V) is greater than Ni (Ni$^{2+}$ (0.25 V). Copper can not displace zinc from copper sulphate solution when zinc piece is added to copper sulphate solution, because standard oxidation potential of copper (−0.34V) is less than the standard oxidation potential (+0.76V) of zinc. This also indicates that more electropositive metals displace less electropositive metals from their salt solutions. Copper is thus less electropositive than zinc.
the strongest reducing agent and the one at the bottom is the weakest reducing agent. Hence lower the reduction potential stronger the reducing agent. The decreasing order of the reducing property of some of the metals is Na > Mg > Zn > Fe > Cu > Ag (More the reduction potential lesser is the reducing power). Note that this is also the arrangement of metals (as given in the electrochemical series) when arranged in the order of their electrode potentials. The arrangement of metals in the order of decreasing tendency of their atoms to give electrons (i.e. undergoing oxidation) is also known as electrochemical series.

**Reaction of metals with acids** – By making use of standard potentials it is also possible to predict whether a metal would react with an acid to liberate hydrogen or not. For example, consider the reaction between zinc and dilute sulphuric acid. The reaction may be represented as,

\[ \text{Zn} + \text{H}_2\text{SO}_4 \rightarrow \text{ZnSO}_4 + \text{H}_2 \]

This reaction is a combination of two half cells.

\[ \text{Zn(s)} \rightarrow \text{Zn}^{2+} + 2e^- \quad \text{(Oxidation)} \]
\[ 2\text{H}^+ + 2e^- \rightarrow \text{H}_2(\text{g}) \quad \text{(Reduction)} \]

In a galvanic cell oxidation occurs at the left hand electrode and reduction at the right hand electrode. Thus the cell can be represented as: \( \text{Zn, Zn}^{2+} \mid \text{H}^+, \text{H}_2(\text{g}), \text{Pt} \)

Thus \( \Delta E_{\text{cell}} = E_{\text{HE}} - E_{\text{LE}} = E_{\text{H}_2(\text{g}), \text{H}^+} - E_{\text{Zn}^{2+}, \text{Zn}} = 0 - (-0.76\text{V}) = +0.76\text{V} \)

Since \( \Delta E_{\text{cell}} \) is positive, the reaction of zinc with hydrogen is spontaneous or feasible. Now consider the reaction of silver with dilute sulphuric acid.

\[ 2\text{Ag} + \text{H}_2\text{SO}_4 \rightarrow \text{Ag}_2\text{SO}_4 + \text{H}_2 \]

The cell may be represented as \( \text{Ag, Ag}^{+} \mid \text{H}^+, \text{H}_2(\text{g}), \text{Pt} \)

\[ \Delta E_{\text{cell}} = E_{\text{H}_2(\text{g}), \text{H}^+} - E_{\text{Ag}^{+}, \text{Ag}} = 0 -(-0.80\text{V}) = +0.80\text{V} \]

Since \( \Delta E_{\text{cell}} \) is negative, the reaction of silver with dilute \( \text{H}_2\text{SO}_4 \) is non-spontaneous or not feasible. In general, any metal lying above hydrogen electrode will reduce \( H^+ \) ions to give \( H_2 \) gas, while any metal lying below hydrogen electrode will not reduce \( H^+ \) ions to give \( H_2 \) gas. Thus metals like Zn, Cd, Fe etc., lying above hydrogen are capable of reacting with dilute acids such as \( \text{H}_2\text{SO}_4, \text{HCl} \) etc. While metals like Ag, Cu etc., which lie below hydrogen electrode, can not evolve hydrogen with dilute acids.

**Secondary cells** – A secondary cell once used can be recharged by passing current through it and it may, therefore, be used over and over again. Certain chemical changes occur when the cell is charged with electricity and these changes are reversed during discharging.

The production of electricity or electric energy at the cost of chemical energy inside is called discharging of the cell. When the external electrical energy is used into the cell to be transformed into chemical energy, the process is called charging of the cell.

**The Leclanche or dry cell** – This is a primary cell used in torches, radio, transistors, hearing aids etc and consists of a negative zinc electrode, a carbon electrode acting as positive electrode, surrounded by a paste of \( \text{MnO}_2 \) and \( \text{ZnCl}_2 \) as an electrolyte. The zinc plate usually forms the outer covering of the cell. The oxidation taking place at the negative zinc electrode (anode) is

\[ \text{Zn} + 2\text{MnO}_2 + \text{H}_2\text{O} \rightarrow \text{Zn}^{2+} + 2\text{MnO}_3 + 2\text{OH}^- \]

The reduction taking place at positive electrode (cathode) is

\[ 2\text{MnO}_2 + \text{H}_2\text{O} + 2e^- \rightarrow 2\text{MnO}_3 + 2\text{OH}^- \]

The net cell reaction is:

\[ \text{Zn} + 2\text{MnO}_2(\text{s}) + \text{H}_2\text{O} \rightarrow \text{Zn}^{2+} + 2\text{MnO}_3 + 2\text{OH}^- \]
During the electrolysis of fused NaCl, chloride ions are oxidised at anode. If current is passed into the solution of an electrolyte, anions move towards anode and cations move towards cathode. In the electrolysis of dilute H₂SO₄ using platinum electrodes H₂ is evolved at cathode and O₂ is evolved at anode. Electrolytes when dissolved in water dissociate into ions because the forces of electrostatic attraction are broken down by water. Magnesium can liberate H₂ gas from hydrochloric acid.

When zinc piece is added to CuSO₄ solution, copper gets precipitated because standard reduction potential of zinc is less than that of copper. In the construction of a cell, the electrode with high reduction potential acts as cathode and the electrode with low reduction potential acts as anode. The electrode at which oxidation occurs is called negative electrode and at which reduction occurs is known as positive electrode. In a galvanic cell cathode acts as positive electrode and anode as negative electrode. In electrolytic cell cathode acts as negative electrode and anode as positive electrode. The more negative the standard reduction potential of a metal, the greater is its ability to displace hydrogen from acids. More the reduction potential less is the reducing power.

By convention, if the oxidation potential of an electrode is positive, the natural tendency of the electrode reaction is oxidation. If it is negative, the natural tendency of the electrode reaction is reduction. Similar is the case with reduction potential. If the reduction potential is positive the natural tendency for the electrode reaction is reduction. If it is negative, the tendency for the electrode reaction is oxidation. The reduction potential is positive for strong oxidising agents, which have great tendency to accept electrons and be reduced. The oxidising power of some species follows the order F₂(g) > MnO₂ > Br₂ (aq) > Ag⁺ > Cu²⁺ > H⁺ > Pb²⁺ > Zn²⁺ > Na⁺.

Fluorine gas (F₂) is the strongest oxidising agent and oxidises almost every other element of the periodic table and is reduced to fluoride ion, F⁻. The more negative the reduction potential, the stronger is the reducing agent. The reducing powers of some species decreases in the order Na⁺ > Zn⁺ > Pb⁺ > H₂ > Cu > Ag⁺ > Br⁻ > Mn²⁺ > F⁻. Thus sodium metal is the strongest reducing agent and F⁻ is the weakest reducing agent. Of all the oxidising agents given above, the sodium ion, Na⁺ is the weakest oxidising agent. Stronger the reducing agent, the weaker is its corresponding oxidising agent. Thus fluorine gas is a strong oxidising agent and F⁻ is a weak reducing agent. We know that weaker an acid, the stronger is its conjugate base. Similarly, the weaker an oxidising agent, the stronger is its corresponding reducing agent.

F₂(g) + 2e⁻ → 2F⁻  Na⁺ + e⁻ → Na

Strong oxidising agent  Weakest reducing agent  Weakest oxidising agent  Strongest reducing agent

A reducing agent will react spontaneously with any oxidising agent stronger than the oxidising agent with which it is coupled (that is, any oxidising agent above its corresponding oxidising agent in the
table of standard reduction potentials). Hence Pb will reduce H\(^{+}\)(aq), Cu\(^{2+}\), Ag\(^{+}\), Br\(_2\) etc but not react with Zn\(^{2+}\) or Na\(^{+}\). An oxidising agent will react with any reducing agent stronger than the reducing agent with which it is coupled (that is, any reducing agent below its corresponding reducing agent in the table of standard reduction potentials). Hence Br\(_2\) will oxidise Ag, Cu, H\(_2\), Pb, Zn and Na, but it will not react with Mn\(^{2+}\) or F\(^{-}\) ions. For example Cu will reduce Ag\(^{+}\) ions because Ag\(^{+}\) is stronger oxidising agent than Cu\(^{2+}\) because Ag\(^{+}\) is above Cu\(^{2+}\) in the table). Similarly Cu will not reduce Pb\(^{2+}\) ions, because Pb\(^{2+}\) is below Cu\(^{2+}\) in the table.

In general, a strong oxidising agent is a species which is readily reduced and hence has a large positive standard reduction potential. Such species are Fe\(_2\) > Co\(^{3+}\) > H\(_2\)O. The halogens above fluorine decrease in oxidising power in the order Cl\(_2\) > Br\(_2\) > I\(_2\). Strong reducing agents are those which are easily oxidised. They have large positive standard oxidation potentials. They include Li > K > Na > ; Ba > Ca > Mg ; Al and some of the transition metals, Zn > Fe > Cd. Hydrogen gas, (SOP) \(_0\) = 0 is moderately good reducing agent. Species below H\(_2\) (which all have negative standard oxidation potentials) are relatively weak reducing agents.

According to preferential discharge theory, out of a number of ions the one which requires least energy will be the first to be discharged at a given electrode.

Greater the standard oxidation potential, stronger the reducing agent will it be or greater would be its reducing power. Greater the reduction potential, greater is the tendency to gain electrons and hence greater is the tendency to be reduced. For example, the reduction potentials of H\(^{+}\), Na\(^{+}\), K\(^{+}\) and Mg\(^{2+}\) ions are 0.000, –2.715, –2.924 and –2.370 respectively. Among these the reduction potential of H\(^{+}\) ion is more. Thus H\(^{+}\) ion has maximum tendency to gain electrons.

Lesser the standard reduction potential of a metal more readily it loses electron and hence greater is its reducing power. Hence more difficult is to reduce it.

The metal having lesser value of standard reduction potential can displace the other from its aqueous salt solution. Thus Sn can displace lead from aqueous lead bromide solution because standard reduction potential of Sn is less than that of Pb. When Zn piece is kept in CuSO\(_4\) solution, copper gets precipitated because standard reduction potential of zinc is less than copper.

The metal having maximum positive value of standard reduction potential is deposited in maximum amount on the cathode. For example, the standard reduction potentials of Ag | Ag\(^{+}\) = 0.8 V, 2H\(_2\) + O\(_2\) = 0.79 V, Cu | Cu\(^{2+}\) = 0.34 V and Mg | Mg\(^{2+}\) = –2.37 V. Thus the sequence of deposition of metals at the cathode is Ag, Hg, Cu, Mg. Hence greater the positive standard reduction potential of a metal, greater would be the amount deposited at the cathode.

In general, a metal can displace another metal, lying below it in the electrochemical series. For example, zinc displaces copper from copper sulphate solution because copper lies below zinc in the electrochemical series.

**One coulomb** = 6.28 \times 10^{28} electrons. **One electric charge** = 1.6 \times 10^{-19} coulombs. **In galvanic cell electrical energy is generated at the expense of chemical energy.** **In an electrolytic cell the chemical energy is generated at the expense of electrical energy.** In **an electrolytic cell current flows from anode to cathode inside the cell.** The electrode which pushes the electrons into the external circuit is called anode. The electrodes which do not participate in the chemical changes are called inert electrodes. The electrode at which electrons flow into an electrolyte is called cathode. **Cathode is negatively charged electrode while anode is positively charged.**  **Unless oxidising and reducing agents are kept separate, electric current will not be produced.**
Abscisic acid (ABA) - A PLANT HORMONE once thought to be responsible for the shedding (abscission) of flowers and fruit and for the onset of dormancy in buds (hence its early name, dormin). The compound is associated with the closing of pores (stoma) in the leaves of plants deprived of water.

Absolute configuration - A particular molecular configuration of a CHIRAL molecule, as denoted by comparison with a reference molecule or by some sequence rule. There are two systems for expressing absolute configuration in common use: the D-L convention and the R-S convention.

Absorption
1. A process in which a gas is taken up by a liquid or solid, or in which a liquid is taken up by a solid. In absorption, the substance absorbed goes into the bulk of the material. Solids that absorb gases or liquids often have a porous structure. The absorption of gases in solids is sometimes called sorption. There is a distinction between absorption (in which one substance is assimilated into the bulk of another) and ADSORPTION (which involves attachment to the surface). Sometimes it is not obvious which process is occurring. For example, a porous solid, such as activated CHARCOAL, may be said to
absorb a large volume of gas, but the process may actually be adsorption on the high surface area of internal pores in the material.

2. The process in which electromagnetic radiation, particles, or sound waves lose energy in passing through a medium. Absorption involves conversion of one form of energy into another.

**Accelerator** - A substance that increases the rate of a chemical reaction. In this sense the term is synonymous with CATALYST. It is common to refer to catalysts as ‘accelerators’ in certain industrial applications. For example, accelerators are used in the VULCANIZATION of rubber and in the polymerization of adhesives. Also, in the production of composite materials using polyester resins a distinction is sometimes made between the catalyst (which initiates the polymerization reaction) and the accelerator (which is an additional substance making the catalyst more effective). The terms promoter and activator are used in a similar way.

**Acceptor** - The atom or group to which a pair of electrons is donated in forming a COORDINATE BOND.

**Acenaphthene** - (C\(_{12}\)H\(_{10}\)) A colorless crystalline derivative of naphthalene, used in producing some dyes.

**Acetal** - A type of compound formed by reaction of an alcohol with either an aldehyde or a ketone. The first step in formation of an acetal is the formation of an intermediate, known as a hemiacetal. For example, ethanal (acetaldehyde; CH\(_3\)CHO) reacts with ethanol (C\(_2\)H\(_5\)OH) as follows: CH\(_3\)CHO + C\(_2\)H\(_5\)OH \xrightleftharpoons{} CH(OH)(CH\(_3\))(C\(_2\)H\(_5\)O) The hemiacetal has a central carbon atom (from the aldehyde) attached to a hydrogen, a hydroxyl group, a hydrocarbon group (methyl - CH\(_3\)), and an alkoxy group (C\(_2\)H\(_5\)O). If a ketone is used rather than an aldehyde, the resulting hemiacetal contains two hydrocarbon groups. For example, reaction of the ketone R\(_1\)COR\(_2\) with the alcohol R\(_3\)OH is: R\(_1\)COR\(_2\) + R\(_3\)OH \xrightleftharpoons{} >CR\(_1\)R\(_2\)(OH)(OR\(_3\))
The formation of a hemiacetal is an example of NUCLEOPHILIC ADDITION to the carbonyl group of the aldehyde or ketone. The first step is attack of the lone pair on the O of the alcohol on the (positively charged) C of the carbonyl group. This is catalyzed by both acids and bases. Acid catalysis occurs by protonation of the O on the carbonyl, making the C more negative and more susceptible to nucleophilic attack. In base catalysis the OH⁻ ions from the base affect the -OH group of the alcohol, making it a more effective nucleophile.

In general, hemiacetals exist only in solution and cannot be isolated because they easily decompose back to the component alcohol and aldehyde or ketone. However, some cyclic hemiacetals are more stable. For example, cyclic forms of SUGAR molecules are hemiacetals.

Further reaction of hemiacetals with another molecule of alcohol leads to a full acetal. For example:

\[
\text{CH(OH)(CH₃)(OC₂H₅)} + \text{C₅H₅OH} \rightarrow \text{CH(CH₃)(OC₂H₅)}
\]

The overall reaction of an aldehyde or ketone with an alcohol to give an acetal can be written: R₁COR₂ + R₃OH \rightarrow CR₁R₂(OR₃)₂

It is also possible to have ‘mixed’ acetals with the general formula CR₁R₂(OR₃)-(OR₄). Note that if the acetal is derived from an aldehyde, then R₁ and/or R₂ may be a hydrogen atom. The mechanism of formation of an acetal from a hemiacetal is acid catalyzed. It involves protonation of the -OH group of the hemiacetal followed by loss of water to form an oxonium ion, which is attacked by the alcohol molecule.

Formerly it was conventional to use the terms ‘hemiacetal’ and ‘acetal’ for compounds formed by reaction between aldehydes and alcohols. Similar reactions between ketones and alcohols gave rise to compounds called hemiketals and ketals. Current nomenclature uses ‘hemiacetal’ and ‘acetal’ for compounds derived from either an aldehyde or a ketone, but reserves ‘hemiketal’ and ‘ketal’ for those derived from ketones. In other words, the ketals are a subclass of the acetals and the hemiketals are a subclass of the hemiacetals.
Achiral - Describing a molecule that does not have chiral properties; i.e. one that does not exhibit OPTICAL ACTIVITY.

**Bond fission**

A covalent bond is formed when electrons are shared between two atoms in the classical sense. A single bond (sigma bond) is thus made up of two electrons. Now a chemical reaction takes place when old bonds are broken and new ones are created. So how can one break a single bond—there are plainly two ways to go about breaking a bond as shown below.

**Homolytic fission**

\[
\text{H}_3\text{C} - \text{CH}_3 \rightarrow \overset{\cdot}{\text{CH}}_3 + \overset{\cdot}{\text{CH}}_3
\]

**Homolysis**

Homolytic fission is where each atom of the bond keeps an electron each resulting in species called free radicals. Radicals are important intermediates in organic chemistry and we will talk about them later. As the bond breaks to give two similar species each keeping an electron this form of bond breaking is called *Homolytic Fission*.

**Heterolytic fission**

\[
\text{H}_3\text{C} - \text{OH} \rightarrow \overset{+}{\text{CH}}_3 + \overset{-}{\text{OH}}
\]

**Heterolysis**

In this case we can see that one of the atoms carry a negative charge after bond cleavage indicating that it has both the electrons of the bond and the other has no electrons at all. Hence it is electron
deficient thus positively charged. As the electrons are not divided equally after bond cleavage this is called **Heterolytic Fission**. In a case the C atom carries a positive charge it is called a carbocation and in the case it carries both the electrons of the broken bond and is negatively charged, it is called a Carbanion. Carbocation and Carbanions are the most important carbon intermediates in organic chemistry. Please note that both types of fissions are applicable to both homoatomic and heteroatomic bonds (bonds between two different atoms say C-N or C-O).

**Free radicals**

These are neutral intermediates, formed due to homolytic cleavage of a single bond. Some common bonds which cleave to give free radicals in organic chemistry are shown: C-O, C-Cl, C-Br, C-I, C-C, C-H. Carbon free radicals are mainly generated by:

i. Photolysis (action of light) like acetone alpha cleavage
ii. Other radical initiator like allylic bromination by N-Bromosuccinimide (NBS)

There has been a certain degree of debate as to what the shape and geometry of a free radical is like. Revisiting the theory of hybridization, there can be two basic shapes of these radicals.

If the centre carbon atom of the radical is sp\(^3\) hybridized (remember the one which was made of one s and three orbitals as in CH\(_4\)), the geometry will be tetrahedral.\(^2\) But in the case of a radical there are only three groups attached to the sp\(^3\) hybridized carbon atom so they will have a shape of what resembles a pyramid—it’s a tetrahedron with its head cut off. So sp\(^3\) hybridized radicals are pyramidal in shape. The single electron of the radical would then be housed in a sp\(^3\) orbital. The other option is sp\(^2\) hybridization. In that case the C atom is sp\(^2\) hybridized, so as discussed previously the shape would be planar with the single electron in the unhybridized p-orbital with the three substituents having sp\(^2\) hybridized bonds.

![Two different geometries of free radicals. The single electrons are shown as black dots.](image)

So to summarize free radicals:

- Formed under activation by light or use of additional compounds called Radical Initiators.
- They are very reactive, because they have an unpaired electron which wants to get paired up.
- They are either pyramidal or planar with the lone electron in their sp\(^3\) or p orbitals respectively.
- Because of their high reactivity, they tend to be less selective. In simple terms it means that it sometimes difficult to predict what products are formed in reactions which involve free radicals and we actually get several products from a single reaction.
This reaction shows the formation of two products with the Br atom attached to different carbons.

**Carbocations**

Carbocations are formed from the heterolytic cleavage of a carbon-heteroatom (meaning a non carbon atom in general) bond where the other atom is more electronegative than carbon like a C-O, C-N, C-X (X can be Cl, Br, I, etc) bond. This is quite logical as after the cleavage if a carbocation is to be formed the two electrons of the bond must go to the other atom. And this is favoured if that other atom is electronegative. Carbocations can be made in difficult conditions by using so-called superacids, developed by George Olah (Nobel Prize, 1994), which helps stabilize these intermediates substantially to be analyzed. Formation of carbocations can be assisted by using cations like Ag⁺, with alkyl halides as substrates.

The precipitating out of the silver salt forces the equilibrium to shift towards the forwards reaction.

A similar reaction is Finkelstein reaction is particularly useful for preparing iodoalkanes. The iodoalkanes are obtained by heating chloro or bromoalkanes with a concentrated solution of sodium iodide in dry acetone.

In Acetone after Reflux (Heating in enclosed container) for a long time [Refluxing is a process of heating a liquid in a flask provided with a condenser. The vapours of the liquid which rise up get condensed back into the flask.]

RX + NaI ------ > RI + NaX where X = Cl, Br, and R is an Alkyl group

Sodium chloride and sodium bromide being less soluble in acetone get precipitated from the solution and can be removed by filtration. This also prevents the backward reaction to occur according to Le Chatelier's principle. The reaction gives the best results with primary halides.

With iodide, the solvate NaI · 3(CH₃)₂CO can be formed, in which each Na⁺ is coordinated by 6 acetone ligands via oxygen lone pairs, and the I⁻ ions fill in the gaps between these octahedral units, being surrounded by the methyl groups of the acetones. I⁻ has a larger ionic radius compared to Cl⁻ and Br⁻ and thus fills the gap completely, thereby minimizing empty space in the lattice and its distance to Na⁺, which is, however, still larger than in solvent-free sodium iodide. This larger distance between the ions decreases the lattice energy, which in turn makes the compound readily soluble in acetone. Cl⁻ and Br⁻ are too small to fit in these gaps, and as the cation-anion distance becomes too large, the only way to minimize the lattice energy is the formation of a solvent-free structure NaX (X =
Cl\(^-\), Br\(^-\)). When the lattice energy is larger than the energy released by solvation of the ions, namely that of Na\(^+\) by complexation with acetone, the compound becomes insoluble and precipitates from the solution.

The positively charged carbon atom in carbocations is sp\(^3\) hybridized, which means it’s planar as we know by now. The three substituents of the carbocation lie in a plane leaving the unhybridized empty p orbital perpendicular to them.

These intermediates react with species which are electron rich (quite obvious) and being charged are stabilized in polar solvents. (Just as Na\(^+\) is soluble and stable in polar water). Carbocations are important intermediates in most mechanisms along with carbanions as we shall see later.

To summarize carbocations:

- Formed due to heterolysis of a C-X bond (where X is more electronegative) and thus has a positive charge.
- Planar in shape (sp\(^2\) hybridized carbon), with empty p orbital perpendicular to the plane of the molecule.
- Reactive towards electron rich species.

Carbanions

These are intermediates also formed as a result of heterolysis, but here the electron pair from the bond is kept by the carbon atom. From what we saw earlier the more electronegative atom keeps the electrons, so in this case carbon must the more electronegative of the two atoms making up the bond. Now there are only a few atoms (non-metals; metals are not usually part of organic chemistry) which are less electronegative, so the most common bond cleavage which yields carbanions is the C-H bond. The ease of breaking this bond and creating a carbanion is also a measure of the compound’s acidity, because a H\(^+\) is also generated with the carbanion, which makes the molecule an acid in the Bronsted sense.

Carbanions have three groups attached to each other and a lone pair of electrons which gives it its negative charge (similar to the ammonia molecule where the central N has 3 Hs and a lone pair of electrons). So its geometry is pyramidal (tetrahedral but since there is no fourth group again it’s like a tetrahedral with head cut off) and the carbon atom is sp\(^3\) hybridized.

Carbanions are also stable in polar solution (electrostatic stabilization).

To summarize carbanions:

- Formed due to heterolysis of a C-X bond (where X is less electronegative) and thus has a negative charge.
- Pyramidal is shape (sp\(^3\) hybridized) with the excess electrons placed in one sp\(^3\) hybrid orbital.
- Reactive towards positively charged (electron deficient species).

Stability of intermediates

Most organic reactions take place via formation of intermediates. So the study of different intermediates would help us predict the course of the reaction and the main aspect to look at would be their stability. No organic mechanism has been conclusively ‘PROVEN’, all the mechanism we see are the most plausible ones derived from many experiments, a major component of which is isolating and
studying the intermediates. It is difficult to say that a certain mechanism is absolutely correct, but it is quite simple to point out an incorrect mechanism. One of the ways a chemist would confirm an incorrect mechanism is if it involves a very unstable intermediate. The good thing about this is that with a few empirical rules and principles in mind, it is quite simple to assign relative stability of intermediates like radicals, carbocations and carbanions. And what is even better is that we have already discussed these principles.

For carbocations and free radicals (both electron poor species), any group which donates electron density to the carbon centre would stabilize it and inversely electron withdrawing groups would increase electron deficiency on the carbon centre leading to destabilization. (Remember charge is not desirable, the most stable species are usually neutral). So following the same logic the effect should just be opposite in the case of carbanions as they are electron rich (negatively charged) instead of being electron deficient like the above two. So groups which pull away electrons from the charged carbon atom would have a stabilizing effect whereas electron donation would destabilize the intermediate as it loads more negative charge on an already negatively charged atom.

The name of a complex radical is considered to start with the first letter of its complete name.

In cases where names of complex radicals are composed of identical words, preference for citation is given to that radical which contains the lowest locant at the first cited point of difference in the radical.

In cases where names of complex radicals are composed of identical words, preference for citation is given to that radical which contains the lowest locant at the first cited point of difference in the radical.
If two or more side chains are in equivalent positions the one to be assigned the lower number is that cited first in the name.

The lowest sum rule.

2,3,5-Trimethyl-4-propylheptane (not 3,5,6-Trimethyl-4-propylheptane)

4-Isobutyl-2,5-dimethylheptane (not 4-Isobutyl-1,2,6-dimethylheptane)
Alphabetical order of numbering

Cyclopentane  1-Methyl-3-propylcyclohexane

More branched carbon gets lower number

3-Ethyl-1,1-dimethylcyclohexane  
(not 1-ethyl-3,3-dimethylcyclohexane)

CLASSIFICATION OF ORGANIC COMPOUNDS

Acyclic or Open chain compounds (I)

Organic compounds

Cyclic or Closed chain or Ring compounds (II)

Homocyclic or Carbocyclic compounds

Alicyclic compounds

Aromatic compounds

Benzenoid compounds

Non-benzenoid compounds

Heterocyclic compounds
**Alicyclic or closed chain or ring compounds**

Alicyclic (aliphatic cyclic) compounds contain carbon atoms joined in the form of a ring (homocyclic). Sometimes atoms other than carbon are also present in the ring (heterocyclic).

Some examples are

- Cyclopropane
- Cyclohexane
- Cyclohexene
- Tetrahydrofuran

**Nomenclature of Substituted Benzene Compounds**

For IUPAC nomenclature of substituted benzene compounds, the substituent is placed as prefix to the word benzene as shown in the following examples.

- Methylbenzene (Toluene)
- Chlorobenzene
- Aminobenzene (Aniline)
- Nitrobenzene
- Bromobenzene

How Isomers are made?

(a) Normal to branched chain in alkane: $n$-Alkane is heated with $\text{AlCl}_3$ at 570 K to give branched chain

$$\text{CH}_3\text{CH}_2\text{CH}_2\text{CH}_2\text{CH}_3 \xrightarrow{\text{AlCl}_3 / 570 \text{K}} \text{CH}_3-\text{CH}-\text{CH}_2-\text{CH}_3$$

(b) Shift of double bond

In the presence of catalyst like $\text{Al}_2(\text{SO}_4)_3$, alkene undergo isomerization at high temperature ($770 - 970 \text{ K}$).

$$\text{CH}_3\text{CH}=\text{CH}_2 \rightarrow \text{CH}_3\text{CH}_2\text{CH}=\text{CH}_3$$

(c) Shift of triple bond

$$\text{CH}_3\text{C}≡\text{CH} \xrightarrow{\text{KOH / ethanol}} \text{[CH}_3\text{C}≡\text{CH}] \rightarrow \text{CH}_3-\text{C}≡\text{C}-\text{CH}_3$$

$$\xrightarrow{\text{2-butyne}} \text{CH}_3\text{CH}_2\text{C}≡\text{CNa} \xrightarrow{\text{NaNH}_3 \text{in inert solvent}}$$
Heating with ethanoic potassium hydroxide shifts the triple bond towards the centre of the chain whereas heating with sodamide in an inert solvent shifts the bond towards the end of chain.

The choice of principal functional group is made on the basis of order of preference. The order of decreasing priority for some functional groups is: -COOH, -SO3H, -COOR (R=alkyl group), COCl, -CONH2, -CN, -HC=O, >C=O, -OH, -NH2, >C=C<, -C=C-.

Epimers

Epimer Anomer comparison
Epimers are monosaccharides that differ in chirality at only one carbon. In the straight-chain format, epimers will have H and OH-substituents switched at one backbone carbon, but not at any others.

Anomers are special epimers; in cyclic forms of one single monosaccharide, anomers differ in chirality at the anomeric (hemiacetal) carbon only. In the straight-chain format, anomers will have the same configuration.
Various Methods of Reduction used in Organic Chemistry

(a) \( \text{CH}_2\text{CH} = \text{CHCHO} \xrightarrow{\text{H}_2, \text{Ni}} \text{CH}_3\text{CH}_2\text{CH}_2\text{CH}_2\text{OH} \)

(b) \( \text{CH}_2\text{CH} = \text{CHCHO} \xrightarrow{\text{Zn} + \text{conc. HCl}} \text{CH}_3\text{CH} = \text{CHCH}_2\text{OH} \)

(c) \( \begin{align*} &\text{CH}_2\text{CH}_2\text{CHO} \xrightarrow{\text{Ni} + \text{NH}_3 + \text{KOH}} \text{CH}_3\text{CH}_2\text{CH}_2\text{CH}_2\text{OH} \\ &\text{CH}_2\text{CH}_2\text{CHO} \xrightarrow{\text{Zn} + \text{conc. HCl}} \text{CH}_3\text{CH} = \text{CHCH}_2\text{OH} \end{align*} \)

(d) \( \text{CH}_3\text{COCH}_3 \xrightarrow{\text{LiAlH}_4} \text{CH}_3\text{CHCH}_3\text{OH} \)

(e) \( \text{RCOOH} \xrightarrow{1. \text{LiAlH}_4 + 2. \text{H}_2\text{O}} \text{RCH}_2\text{OH} \)

(f) \( \begin{align*} &\text{NO}_2 \xrightarrow{\text{H}_2, \text{Pt} \text{ or } \text{Sn, HCl} \text{ heat} \text{ or } \text{Fe, HCl}} \text{NH}_2 \\ &\text{NO}_2 \xrightarrow{(\text{NH}_4)_2\text{S}} \text{NH}_2 \text{NO}_2 \end{align*} \)

(g) \( \text{CH}_2\text{CN} \xrightarrow{\text{H}_2, \text{Ni} \text{ at } 140 \degree \text{C}} \text{CH}_3\text{CH}_2\text{NH}_2 \)

(h) \( \begin{align*} &\text{NO}_2 \xrightarrow{\text{H}_2, \text{Pt} \text{ or } \text{Sn, HCl} \text{ heat} \text{ or } \text{Fe, HCl}} \text{NH}_2 \\ &\text{NO}_2 \xrightarrow{(\text{NH}_4)_2\text{S}} \text{NH}_2 \text{NO}_2 \end{align*} \)

(i) \( \text{RC} \xrightarrow{\text{Mg}} \text{R}^\prime \xrightarrow{\text{MgCl}_2} \text{LiAlH}_4 \xrightarrow{\text{H}_2\text{O}} \text{RH} + \text{MgCl} \)

(j) \( \text{Rosenmund's reduction} \)

Various methods of Oxidation

**Oxidation**

(a) \( \text{RCH}_2\text{OH} \xrightarrow{\text{pyridinium chlorochromate}} \text{RCO} \)

(b) \( \text{R}^\prime \text{CHOH} \xrightarrow{\text{K}_2\text{Cr}_2\text{O}_7} \text{R}^\prime \text{C} = \text{O} \)

(c) \( \text{RCH}_2\text{OH} \xrightarrow{\text{KMnO}_4} \text{RCOOH} \)

(d) \( \text{CH}_3 \xrightarrow{\text{CrO}_3} \xrightarrow{\text{H}_2\text{O}, \text{Ac}_2\text{O}, \text{H}^+} \text{CHO} \)

(e) \( \text{CH}_3 \xrightarrow{\text{KMnO}_4} \text{COOH} \)

(f) \( \text{RCH} = \text{CHCHO} \xrightarrow{\text{Tollens reagent}} \text{RCH} = \text{CHCOOH} \)
Synthesis of p-bromonitrobenzene in 2 steps


a synthesis of p-bromonitrobenzene from benzene in two steps.

![Chemical reaction diagram]

\[ \text{Secondary alcohol to tertiary alcohol} \]
\[ \begin{align*}
\text{CH}_3\text{CHCH}_3 & \xrightarrow{K_2\text{Cr}_2\text{O}_7} \text{CH}_3\text{CCH}_3 \\
\text{OH} & \xrightarrow{1. \text{CH}_3\text{MgBr}, \text{Br}, \text{H}^+} \text{CH}_3\text{C}\text{-OH}
\end{align*} \]

Prepare m-bromiodobenzene from benzene

prepare m-bromiodobenzene from benzene (in not more than 5-7 steps)?

![Chemical reaction diagram]

Alternatively, we may proceed as follows:

![Chemical reaction diagram]
Secondary alcohol to Primary alcohol

\[
\begin{align*}
\text{CH}_3\text{CHCH}_2\text{OH} & \xrightarrow{\text{conc. H}_2\text{SO}_4} \text{CH}_3\text{CH} \equiv \text{CH}_2 & \xrightarrow{\text{HBr}} \text{CH}_3\text{CHCH}_2\text{Br} \\
& \xrightarrow{\text{oq. KOH}} \text{CH}_3\text{CH}_2\text{CH}_2\text{OH}
\end{align*}
\]

To convert Ethylebenzene to Benzene and Ethylebenzene to phenylpropionic acid

Show the steps to carry out the following transformations.
(i) Ethylbenzene → benzene

(ii) Ethylbenzene → 2-phenylpropionic acid.

Alcohol to alcohol with one carbon more

\[
\begin{align*}
\text{CH}_3\text{OCH}_3 & \xrightarrow{\text{PBr}_3} \text{CH}_3\text{Br} & \xrightarrow{\text{KCN}} & \text{CH}_3\text{CN} & \xrightarrow{[\text{H}]^+} & \text{CH}_3\text{CH}_2\text{NH}_2 & \xrightarrow{\text{HNO}_2} & \text{CH}_3\text{CH}_2\text{OH}
\end{align*}
\]

Carry out the following transformation in not more than three steps.

Alcohol to alcohol with one carbon less

\[
\begin{align*}
\text{CH}_3\text{CH}_2\text{OH} & \xrightarrow{\text{[O]}^\cdot} \text{CH}_3\text{COOH} & \xrightarrow{\text{soda lime}} & \text{CH}_4\text{Cl}_2 & \xrightarrow{\text{aq. KOH}} & \text{CH}_3\text{OH}
\end{align*}
\]
Conversion of Aniline to Benzylamine

How would you bring about the following conversion (in 3 steps)?
Aniline $\rightarrow$ Benzylamine

\[
\text{NH}_2 \quad \text{NaNO}_2/\text{dil. HCl} \quad 273-283 \text{ K} \quad \text{CuCN} \quad \text{H}_2\text{Ni} \quad \text{CH}_2\text{NH}_2
\]

\[
\text{O} \quad \text{O} \quad \text{O} \\
\text{H-C-\text{OH}} \quad \text{C-C-\text{OH}} \quad \text{C-C-\text{COH}}
\]

Methanoic acid Ethanoic acid Propanoic acid

(i) Carbon-oxygen bond lengths in formic acid are different but are the same in sodium formate. Justify.

(ii) Phenate ion has more number of contributing structures than benzoate ion: but still benzoic acid is a stronger acid. Explain.

Solution. (i) In sodium formate, the contributing structures for the anion are equivalent while these are not the same in formic acid.

\[
\text{H-C=O} \quad \text{H-C=O} \\
\text{Formate ion} \quad \text{Formate ion}
\]

Thus, carbon-oxygen bond length in formate ion is the same for both the bonds while these are different in formic acid.

(ii) In phenate ion, the negative charge is dispersed only one electronegative oxygen atoms while there are two available oxygen atoms in benzoate ion to disperse the negative charge.
Amine to amine containing one carbon more

\[ \text{RCH}_2\text{NH}_2 \xrightarrow{\text{HNO}_2, 0 \degree \text{C}} \text{RCH}_2\text{OH} \xrightarrow{\text{PBr}_3} \text{RCH}_2\text{Br} \xrightarrow{\text{NaCN}} \text{RCH}_2\text{CN} \xrightarrow{\text{H}_2/\text{Ni}} \text{RCH}_2\text{CH}_2\text{NH}_2 \]

Synthesis of 4-methoxyphenol from bromobenzene

How would you synthesise 4-methoxyphenol from bromobenzene is NOT more than five steps? State clearly the reagents used in each step and show the structure of the intermediate compounds in your synthetic scheme.

The introduction of \(-\text{NO}_2\) group at \textit{para} position in bromobenzene facilitates the nucleophilic replacement of \(-\text{Br}\) by \(-\text{OCH}_3\).

The negative charge is strongly stabilised by resonance involving \textit{para}-nitro group.

\textbf{Write chemical tests to distinguish}

\textbf{Solution.} Formic acid \( \left( \begin{array}{c} \text{O} \\ \text{H} - \text{C} - \text{OH} \end{array} \right) \) contains both an aldehyde \( \left( \begin{array}{c} \text{H} \\ -\text{C} = \text{O} \end{array} \right) \) as well as carboxyl.
group \(\text{C--O--H}\) but acetic acid contain only a carboxyl group. Formic acid behaves as reducing agent whereas acetic acid does not.

(a) Formic acid reduces Tollen’s reagent to metallic silver but acetic acid does not.

\[
\text{HCOOH} + 2\text{[Ag(NH}_3\text{)_2]}^+ + 2\text{(OH)}^- \\
\text{Tollen’s reagent} \\
\quad \longrightarrow 2\text{Ag} \downarrow + \text{CO}_2 \uparrow + 2\text{H}_2\text{O} + 4\text{NH}_3 \\
\text{silver mirror}
\]

No silver mirror is formed with acetic acid.

(b) Formic acid reduced Fehling solution to red ppt. of Cu\(_2\)O but acetic acid does not.

\[
\text{HCOOH} + 2\text{Cu}^{2+} + 4\text{(OH)}^- \longrightarrow \text{Cu}_2\text{O} \downarrow + \text{CO}_2 \uparrow + 3\text{H}_2\text{O} \\
\text{Fehling solution} \quad \text{red ppt. (cuprous oxide)}
\]

Amine to amine containing one carbon less

\[
\text{RCH}_2\text{NH}_2 \xrightarrow{\text{HNO}_3, 0^\circ\text{C}} \text{RCH}_2\text{OH} \xrightarrow{\text[K_mO}_4\text{]} \text{RCOOH} \xrightarrow{\text{SOCl}_2} \text{RCOCl} \xrightarrow{\text{NH}_3} \text{RCO\text{NH}_2} \xrightarrow{\text{OBr}} \text{RNH}_2
\]

Convert Benzoic acid to meta fluorobenzoic acid (2-fluorobenzoic acid)

Convert

\[
\begin{array}{cc}
\text{COOH} & \xrightarrow{\text{not more than 3 steps}} \text{COOH} \\
\text{benzoic acid} & \text{metafluoro benzoic acid}
\end{array}
\]
Convert Nitrobenzene to meta hydroxynitrobenzene (or 4-nitrophenol)

Conditions and intermediates in the above conversion. The given conversion can be carried out as follows.

Nitrobenzene resonance structure

4 nitrophenol or meta hydroxynitrobenzene
Many conversions or preparations

Write down the reactions involved in the preparation of the following using the reagents indicated against each in parenthesis.

(i) Ethylbenzene from benzene \([C_7H_8OH, \text{PCl}_3, \text{anhydrous AlCl}_3]\)
(ii) Propanoic anhydride from propanol \([\text{AgNO}_3/\text{NH}_4\text{OH}, \text{P}_2\text{O}_5]\)
(iii) Acetoxime from acetaldehyde \([\text{K}_2\text{Cr}_2\text{O}_7/\text{H}^+, \text{Ca(OH)}_2 \text{and NH}_2\text{OH.HCl}]\)

\[
\begin{align*}
\text{C}_7\text{H}_8&+\text{PCl}_3 \rightarrow \text{C}_7\text{H}_8\text{Cl} + \text{POCl}_3 + \text{HCl} \\
\text{C}_3\text{H}_6 + \text{C}_2\text{H}_2\text{Cl} & \xrightarrow{\text{AlCl}_3} \text{C}_6\text{H}_5\text{C}_2\text{H}_3 + \text{HCl}
\end{align*}
\]

\[
\begin{align*}
\text{C}_3\text{H}_8\text{OH} \xrightarrow{\text{AgNO}_3/\text{NH}_4\text{OH}} & \text{C}_2\text{H}_5\text{COOH} \\
2\text{C}_2\text{H}_5\text{COOH} & \xrightarrow{\text{P}_2\text{O}_5} (\text{C}_2\text{H}_5\text{CO})_2\text{O} \\
\text{CH}_3\text{CHO} \xrightarrow{\text{K}_2\text{Cr}_2\text{O}_7/\text{H}^+} & \text{CH}_3\text{COOH} \\
\text{CH}_3\text{COOH} & \xrightarrow{\text{Ca(OH)}_2} (\text{CH}_3\text{COO})_2\text{Ca} \\
(\text{CH}_3\text{COO})_2\text{Ca} & \xrightarrow{\text{heat}} \text{CH}_3\text{COCH}_3 + \text{CaCO}_3 \\
\text{CH}_3\text{COCH} & \xrightarrow{\text{NH}_2\text{OH}} \text{CH}_3\text{C} = \text{N} - \text{OH}
\end{align*}
\]

Benzene to p-toluic acid

Benzene to p-toluic acid

Benzene to m-chlorobenzoic acid

Benzene to p-chlorobenzoic acid
Benzene to p-chlorobenzoic acid

Acetic acid to Malonic acid

Ethyne to propanoic acid

Isopropyl alcohol to β-hydroxybutyric acid

Acetic acid to Propanoic acid
Ethanoic acid to a mixture of methanoic acid and diphenyl ketone

\[
\text{CH}_2\text{COOH} \xrightarrow{\text{SOCl}_2} \text{CH}_3\text{COCl} \quad \text{(acetic acid)} \\
\text{C}_6\text{H}_5\text{COC}_6\text{H}_5 \xrightarrow{\text{KMnO}_4} \text{C}_6\text{H}_5\text{C}=\text{C} \quad \text{(formic acid)}
\]

Alternatively, we may proceed as follows:

\[
\text{CH}_3\text{COOH} \xrightarrow{\text{SOCl}_2} \text{CH}_3\text{COCl} \quad \text{(acetyl chloride)} \\
\text{C}_6\text{H}_5\text{C}=\text{C} \xrightarrow{\text{C}_6\text{H}_5\text{MgBr excess}} \text{C}_6\text{H}_5\text{CH}_3 \\
\text{HCOOH} \xrightarrow{[\text{O}]} \text{HCHO} \quad \text{(formaldehyde)}
\]

Propanoic acid to acetic acid

\[
\text{CH}_3\text{CH}_2\text{COOH} \xrightarrow{\text{NH}_3, \text{heat}} \text{CH}_3\text{CH}_2\text{CONH}_2 \xrightarrow{\text{Br}_2, \text{KOH}} \text{CH}_3\text{CH}_2\text{NH}_2 \xrightarrow{\text{HNO}_3} \text{CH}_3\text{CH}_2\text{OH} \xrightarrow{\text{Cr}_2\text{O}_7^{2-}, \text{H}^+} \text{CH}_3\text{COOH}
\]
Aniline to 1, 2, 3-tribromobenzene

Ethanal to 2-hydroxy-3-butenoic acid

Benzene to m-bromophenol

Ethyl alcohol to Vinyl alcohol

Methylamine to ethylamine
Benzene to m-bromotoluene

Benzaldehyde to cyanobenzene

Aniline to Chlorobenzene

Ethylamine to Methylamine
Benzene to m-dibromobenzene

Aniline to p-nitroaniline

Toluene to m-nitrotoluene
Convert Benzene + Alcohol to α-methylphenylacetic acid

Benzene + alcohol to α-methylphenylacetic acid

\[
\begin{align*}
C_2H_5OH & \xrightarrow{C_6H_5NH^+CrO_4Cl^-} CH_3CHO \\
\text{Br} & \xrightarrow{\text{Fe}} \text{Mg} \\
\text{MgBr} & + \text{CH}_3\text{CHO} \rightarrow \text{CH}_3\text{CH(OH)CH}_3 \\
\text{CHCOOH} & \xrightarrow{\text{CO}_2} \text{CHMgCl} \\
\end{align*}
\]

Toluene to 3,5 dibromotoluene

Toluene to 3, 5-dibromotoluene

\[
\begin{align*}
\text{CH}_3 & \xrightarrow{\text{NO}_2} \text{CH}_3 \\
\text{Fe} & \xrightarrow{\text{H}^+} \text{NH}_2 \\
2\text{Br} & \rightarrow \text{Br} \text{Br} \\
\text{H}_3\text{PO}_2 & \rightarrow \text{N}^+ \text{Br} \\
\text{HNO}_2 & \rightarrow \text{H}^+ \\
\end{align*}
\]
Benzene to p-chloronitrobenzene

\[
\text{Benzene to } p\text{-chloronitrobenzene}
\]

\[
\text{Cl}_2 \xrightarrow{\text{Fe}} \text{Cl} \quad \text{HNO}_3 \xrightarrow{\text{H}_2\text{SO}_4} \text{NO}_2
\]

Toluene to 2,6 dibromotoluene

\[
\text{Toluene to } 2,6\text{-dibromotoluene}
\]

\[
\text{CH}_3 \xrightarrow{\text{HNO}_3 \xrightarrow{\text{H}_2\text{SO}_4}} \text{NO}_2 \quad 2\text{Br}_2 \xrightarrow{\text{Fe}} \text{Br} \quad \text{CH}_3 \xrightarrow{\text{Fe} \xrightarrow{\text{H}^+}} \text{Br} \quad \text{NO}_2
\]

Benzene to 3,4-dibromonitrobenzene

\[
\text{Benzene to } 3,4\text{-dibromonitrobenzene}
\]

\[
\text{Br}_2 \xrightarrow{\text{Fe}} \text{Br} \quad \text{HNO}_3 \xrightarrow{\text{H}_2\text{SO}_4} \text{NO}_2 \quad \text{Br}_2 \xrightarrow{\text{Fe}} \text{Br}
\]
Benzene to m-chloronitrobenzene

Benzene to m-chloronitrobenzene

Nitrobenzene to benzamide

Nitrobenzene to benzamide

Benzene to 4-nitrobenzaldehyde

Benzene to 4-nitrobenzaldehyde

Benzene to 4-amino-2-bromotoluene

Benzene to 4-amino-2-bromotoluene

2,4-Dinitrophenol (DNP)
Convert a mixture of cis-2-pentene and trans-2-pentene into cis-2-pentene or trans-2-pentene

A mixture of cis-2-pentene and trans-2-pentene into cis-2-pentene or trans-2-pentene

Benzene to p-chlorobenzylalcohol

Benzene to p-chlorobenzylalcohol
Convert Propanol to $\alpha$-hydroxypropanoic acid

\[
\text{CH}_3\text{CH}_2\text{CH}_2\text{OH} \xrightarrow{\text{KmnO}_4} \text{CH}_3\text{CH}_2\text{COOH} \xrightarrow{\text{Br}_2/P} \text{CH}_3\text{CH}_2\text{COOH} \xrightarrow{\text{OH}^-} \text{CH}_3\text{CHCOOH}
\]

4-chloropentane-2-one

3-Bromo-4-methylpentanoic acid

$p,p'$-dihydroxybenzophenone

Hex-2-ene-4-ynoic acid

Heptan-2-one
Benzene to Mandelic Acid

4-Bromo-2-methylhexane

Heptanal

1-(4-Methoxyphenyl)-3-phenylprop-2-en-1-one

4-phenylprop-3-ene-2-one
3-phenylprop-2-ene-1-al

3-phenyl-prop-2-ene-1-ol

cyclopentancarbaldehyde

Benzophenone also diphenylmethanone
Tollen’s Reagent

$$RCHO(aq) + 2Ag(NH_3)_2^+(aq) + 3OH^-(aq) \rightarrow$$

An aldehyde

$$RCOO^-(aq) + 2Ag(s) + 4NH_3(aq) + 2H_2O$$

Free silver

An aldose

Tollens’ reagent (clear solution)

Carboxylate anion

Silver

An aldose

Benedict’s reagent (blue solution)

Carboxylate anion

Brick-red precipitate

Benzene to 2,4-dinitrobenzaldehyde

Acetaldehyde to Acetylene
Identify the compounds

compound contains C, H and O unreactive towards Na (Sodium)

A compound (X) containing only C, H and O is unreactive towards sodium. It does not add bromine. It also does not react with Schiff’s reagent. On refluxing with an excess of hydroiodic acid, X yields only one organic product Y. On hydrolysis Y yields a new compound Z which can be converted to Y by reaction with red phosphorus and iodine. The compound Z on oxidation with potassium permanganate gives a carboxylic acid. The equivalent weight of this acid is 60. What are the compounds X, Y and Z? Write chemical equations leading to the conversion of X to Y. (1981)

Solution Since the compound is unreactive towards sodium, it is neither alcohol (including phenol) nor carboxylic acid. It is also not an aldehyde. It may be a symmetrical ether because on refluxing with excess of HI, it gives only one product. Thus, we have

\[ \text{ROR} \xrightarrow{\text{HI}} \text{2RI} \xrightarrow{\text{H}_2\text{O}} \text{2ROH} \xrightarrow{\text{IO}_3^-} \text{2R’COOH} \]

Now the equivalent weight of R’COOH is 60. From this, it follows that R’ is \(-\text{CH}_3\) group and hence R is \(-\text{CH}_2\text{CH}_3\) group. Thus, the compounds X, Y and Z are

X: \text{CH}_3\text{CH}_2\text{OCH}_2\text{CH}_3; \quad Y: \text{CH}_3\text{CH}_2\text{I}; \quad Z: \text{CH}_3\text{CH}_2\text{OH}
Identify the compound

A hydrocarbon reacts with Bromine to give dibromo compound

One mole of a hydrocarbon A reacts with one mole of bromine giving a dibromo compound $\text{C}_4\text{H}_8\text{Br}_2$. Substance A on treatment with cold, dilute alkaline potassium permanganate solution forms a compound $\text{C}_3\text{H}_6\text{O}_2$. On ozonolysis, A gives equimolar quantities of propanone and ethanol. Deduce the structural formula of A.

Solution From the products of ozonolysis, we conclude that

\[
\begin{align*}
\text{propanone} & : & \begin{array}{c}
\text{CH}_3
\end{array} & \begin{array}{c}
\text{C} = \text{O}
\end{array} \quad \begin{array}{c}
\text{H}
\end{array} \\
\text{ethanol} & : & \begin{array}{c}
\text{CH}_3
\end{array} & \begin{array}{c}
\text{C} = \text{C}
\end{array} & \begin{array}{c}
\text{H}
\end{array}
\end{align*}
\]

The compound A contains a double bond. This is also confirmed from the fact that the compound A reacts with one mole of bromine. Thus, the given reactions are

\[
\begin{align*}
\text{OH} & \quad \text{OH} \\
\text{CH}_3 & \quad \text{C} \quad \text{C} \quad \text{CH}_3 & \text{dil alkaline KMnO}_4 & \text{cold} & \text{Br}_2 & \text{Br} & \text{Br} \\
\text{CH}_3 & \quad \text{H} & \text{CH}_3 & \quad \text{Br} & \text{CH}_3 & \quad \text{C} \quad \text{C} \quad \text{CH}_3 & \text{CH}_3 & \quad \text{H}
\end{align*}
\]

Identify the compound

An alkene A on ozonolysis yields acetone and an aldehyde

An alkene A on ozonolysis yields acetone and an aldehyde. The aldehyde is easily oxidized to an acid B. When B is treated with bromine in presence of phosphorus, it yields a compound C which on hydrolysis gives a hydroxy acid D. This acid can also be obtained from acetone by the reaction with hydrogen cyanide followed by hydrolysis. Identify the compounds A, B, C and D.

Solution The structure of compound D is easily deduced from the fact that it is obtained by the combination of acetone with hydrogen cyanide followed by hydrolysis.

\[
\text{CH}_3 & \quad \text{C} = \text{O} & \text{HCN} & \text{H}^+ & \text{H}_2\text{O} & \text{CH}_3 & \quad \text{C} \quad \text{COOH}
\]

The conversion B to C and then C to D indicates that the reaction B to C is Hell-Volhard-Zelinsky reaction. Thus, we have

\[
\begin{align*}
\text{CH}_3 & \quad \text{C} \quad \text{H} & \text{Br}_2 & \text{P} & \text{Br} & \text{CH}_3 & \quad \text{C} \quad \text{COOH} & \text{H}_2\text{O} & \text{CH}_3 & \quad \text{C} \quad \text{COOH}
\end{align*}
\]

Hence, we conclude that the aldehyde from which compound B is obtained by oxidation has the structure

\[
\begin{align*}
\text{CH}_3 & \quad \text{H} & \text{2-methylpropanal}
\end{align*}
\]

Now, the ozonolysis of the compound A gives acetone and 2-methyl-propanal, i.e.
Identify the compound

An aromatic compound contains 69.4% carbon, 5.8% hydrogen

An aromatic compound contains 69.4% carbon and 5.8% hydrogen. A sample of 0.303 g of this compound was analysed for nitrogen by Kjeldahl’s method. The ammonia evolved was absorbed in 50 mL of 0.05 M sulphuric acid. The excess acid required 25 mL of 0.1 M sodium hydroxide for neutralization. Determine the molecular formula of the compound if its molar mass is 121 g mol$^{-1}$. Draw two possible structures for this compound.

**Solution**

The reactions involved are

$$2\text{NH}_4 + \text{H}_2\text{SO}_4 \rightarrow (\text{NH}_3)_2\text{SO}_4$$
$$\text{H}_2\text{SO}_4 + 2\text{NaOH} \rightarrow \text{Na}_2\text{SO}_4 + 2\text{H}_2\text{O}$$

Amount of sodium hydroxide to neutralize excess of sulphuric acid = (25 mL) $\frac{0.1 \text{ mol}}{1000 \text{ mL}} = \frac{1}{400} \text{ mol}$

Amount of sulphuric acid neutralized with sodium hydroxide = $\frac{1}{2} \left( \frac{1}{400} \text{ mol} \right) = \frac{1}{800} \text{ mol}$

Amount of sulphuric acid taken to start with = (50 mL) $\frac{0.05 \text{ mol}}{1000 \text{ mL}} = \frac{1}{400} \text{ mol}$
An Organic compound CxH2yOy was burnt with twice the amount of Oxygen

An organic compound CxH2yO2 was burnt with twice the amount of oxygen needed for complete combustion to CO2 and H2O. The hot gases, when cooled to 0 °C and 1 atm pressure, measured 2.24 L. The water collected during cooling weighed 0.9 g. The vapour pressure of pure water at 20 °C is 17.5 mmHg and is lowered by 0.104 mmHg when 50 g of the organic compound is dissolved in 1000 g of water. Give the molecular formula of the organic compound.

Solution: The combustion reaction is \( \text{C}_x\text{H}_y\text{O}_2 + x \text{O}_2 \rightarrow x \text{CO}_2 + y \text{H}_2\text{O} \)

To start with, the amount of O2 taken is 2x. Hence, after the combustion reaction, we will be left with the following amounts.

- Amount of oxygen left unreacted = x
- Amount of carbon dioxide = x
- Amount of water = y

When this mixture is cooled to 0 °C and 1 atm, we will be left with oxygen and carbon dioxide. Hence, the amount 2x occupies the given volume of 2.24 L at STP. Hence,

\[
22.4 \text{ L} \text{ mol}^{-1} \times \frac{2.24 \text{ L}}{x} = 0.05 \text{ mol}
\]

Now,

\[
\Delta p = \frac{m_i}{(50 \text{ g/M})} = \frac{0.104 \text{ mmHg}}{17.5 \text{ mmHg}} = \frac{0.05 \text{ mol}}{18 \text{ g mol}^{-1}}
\]

Thus, the empirical formula of the compound is \( \text{C}_x\text{H}_y\text{O}_2 \), i.e., CH2O. Now, according to Raoult’s law,

\[
\boxed{\text{Empirical molar mass} = 121 \text{ g mol}^{-1}}
\]
Identify the Compound

Alcohol when heated with concentrated H₂SO₄

An alcohol A, when heated with concentrated H₂SO₄ gives an alkene B. When B is bubbled through bromine water and the product obtained is dehydrohalogenated with excess of sodiumide, a new compound C is obtained. The compound C gives D when treated with warm dilute H₂SO₄ in presence of HgSO₄. D can also be obtained either by oxidizing A with KMnO₄ or from acetic acid through its calcium salt. Identify A, B, C and D.

Solution
The given reactions are as follows.

\[
\text{A (alcohol)} \xrightarrow{\text{conc. } \text{H}_2\text{SO}_4} \text{B (alkene)} \xrightarrow{1. \text{Br}_2, 2. -2\text{HBr}} \text{C} \xrightarrow{\text{HgSO}_4/\text{H}_2\text{SO}_4} \text{D}
\]

The reaction of obtaining D from calcium acetate is

\[(\text{CH}_3\text{COCO})_2\text{Ca} \rightarrow \text{CH}_3\text{COCH}_3 + \text{CaCO}_3
\]

Hence, D is acetone. Since the latter is also obtained from the oxidation of alcohol A, compound A will be CH₃CH₂CH₃.

Therefore, the reactions depicted above are as follows.

\[
\text{CH}_3\text{CHCH}_3 \xrightarrow{\text{HCl, } \text{H}_2\text{O}} \text{CH}_3\text{CH}═\text{CH}_2 \xrightarrow{\text{Br}_2} \text{CH}_3\text{CH}═\text{CH}_2
\]

Hence, the reactions depicted above are as follows.

\[
\text{CH}_3\text{CHCH}_3 \xrightarrow{\text{HCl, } \text{H}_2\text{O}} \text{CH}_3\text{CH}═\text{CH}_2 \xrightarrow{\text{Br}_2} \text{CH}_3\text{CH}═\text{CH}_2
\]

Hence, the reactions depicted above are as follows.

\[
\text{CH}_3\text{CHCH}_3 \xrightarrow{\text{HCl, } \text{H}_2\text{O}} \text{CH}_3\text{CH}═\text{CH}_2 \xrightarrow{\text{Br}_2} \text{CH}_3\text{CH}═\text{CH}_2
\]
Certain hydrocarbon was found to contain 85.7% carbon

A certain hydrocarbon A was found to contain 85.7 per cent carbon and 14.3 per cent hydrogen. This compound consumes 1 molar equivalent of hydrogen to give a saturated hydrocarbon B. 1.0 g of hydrocarbon A just decolorized 38.05 g of a 5 per cent solution (by mass) of Br₂ in CCl₄. Compound A, on oxidation with concentrated KMnO₄, gave compound C (molecular formula C₆H₄O₂) and compound C could easily be prepared by the action of acidic aqueous mercuric sulphate on 2-butyne. Determine the molecular formula of A and deduce the structures of A, B and C.

**Solution**

The ratio of atoms in the compound A is \( \frac{85.7}{12} : \frac{14.3}{1} : \frac{7.14}{14.3} : \frac{14.3}{1} : \frac{1}{2} \)

Thus, Empirical formula of A is CH₂.

Since the compound A consumes 1 mol of hydrogen, the molecule of A contains only one carbon-carbon double bond. From the data on the absorption of bromine, we can calculate the molar mass of A as shown in the following.

Mass of bromine absorbed by 1.0 g of hydrocarbon = \( \frac{5}{100} \times 38.05 \) g

Mass of hydrocarbon absorbing 160 g (= 1 mol) of Br₂ = \( \frac{1.0}{(5 \times 38.05/100)} \) × 160 g = 84.1 g.

Hence, Molar mass of A is 84.1 g mol⁻¹.

The number of repeating CH₂ group in one molecule of A will be \( \frac{6}{(84.1/14)} \). Hence, Molecular formula of A is C₆H₁₂₂₄. Now, it is given that

\[ \text{C}_6\text{H}_{12} \xrightarrow{\text{conc. KMnO}_4} \text{C}_4\text{H}_6\text{O} + \text{CH}_3\text{COOH} \]

The compound C is obtained by the hydration of 2-butyne. Hence, its structure obtained from the reaction is

<table>
<thead>
<tr>
<th>C₄H₆O₂</th>
<th>CH₂C=CHCH₂OH</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>H₂O</td>
</tr>
<tr>
<td></td>
<td>HgSO₄/H₂SO₄</td>
</tr>
<tr>
<td></td>
<td>CH₃C(=CHCH₃) OH</td>
</tr>
<tr>
<td></td>
<td>CH₃CCH₂CH₃</td>
</tr>
<tr>
<td></td>
<td>2-butylene</td>
</tr>
<tr>
<td></td>
<td>2-butane</td>
</tr>
</tbody>
</table>

Identify the compound

A liquid X having a molecular formula C₆H₁₂O₂ is hydrolyzed with water

A liquid X, having a molecular formula C₆H₁₂O₂, is hydrolysed with water in the presence of an acid to give a carboxylic acid, Y, and an alcohol Z. Oxidation of Z with chromic acid gives Y. What are the structures of X, Y and Z?

**Solution**

Since the hydrolysis of X gives a carboxylic acid (Y) and an alcohol (Z), the compound X must be an ester. Let it be RCOOR'. The compounds Y and Z will be RCOOH and R'OH, respectively. Since the oxidation of Z gives Y, we will have

\[ \text{R'O} \xrightarrow{[O]} \text{RCOOH} \quad \text{or} \quad \text{RCO}_{2}\text{H} \xrightarrow{[O]} \text{RCOOH} \]

Hence, the given compound X may be written as RCOOCH₂R. From this it follows that

\[ 2\text{R}=\text{C}_6\text{H}_{12} \xrightarrow{\text{O}} \text{C}_2\text{H}_4 \text{O}=\text{C}_6\text{H}_{10} \quad \text{or} \quad \text{R}=\text{C}_2\text{H}_5 \]

\[ \text{X} : \text{CH}_3\text{CH}_2\text{COOCH}_2\text{CH}_3 \quad \text{Propylene glycol} \]

\[ \text{Y} : \text{CH}_2\text{CH}_2\text{COOH} \quad \text{Propanoic acid} \]

\[ \text{Z} : \text{CH}_3\text{CH}_2\text{CH}_2\text{OH} \quad \text{Propanol} \]
Identify the compound

An unknown compound of carbon, hydrogen, and oxygen contains 69.77% carbon and 11.63% hydrogen and has a molar mass of 86 g mol⁻¹. It does not reduce Fehling’s solution, but forms a bisulphite addition compound and gives a positive iodoform test. What are the possible structures?

Solution The given compound is a ketone as it does not reduce Fehling’s solution, but forms a bisulphite addition compound. It will also contain CH₃CO group as it shows positive iodoform test.

The ratios of atoms in the compound are:

\[ C : O : H :: \frac{69.77}{12} : \frac{11.63}{16} : \frac{5.81}{5} = 5.16 : 11.63 : 5 : 1 : 10 \]

Hence, empirical formula is C₈H₆O₃. Molar empirical mass = 86 g mol⁻¹; same as the given molar mass.

Thus, molecular formula is C₈H₁₀O₃. The structure of the compound will be

- CH₃C(CH₂)₂CH₃ or 2-pentanone
- CH₃C=CH(CH₃)O
- 3-methylbutan-2-one

Identify the compound

An organic compound A on treatment with acetic acid in the presence of sulphuric acid

An organic compound A on treatment with acetic acid in the presence of sulphuric acid produces an ester B. The compound A on mild oxidation gives C. The compound C with 50% potassium hydroxide followed by acidification with dilute hydrochloric acid generates A and D. The compound D with phosphorus pentachloride followed by reaction with ammonia gives E. The compound E on dehydration produces hydrocyanic acid. Identify the compounds A, B, C, D, and E.

Solution The given reactions are:

A + CH₃COOH \xrightarrow{H_2SO_4} ester; \quad A \xrightarrow{\text{mild oxidation}} C \xrightarrow{50\% \text{ KOH}} A + D

D \xrightarrow{\text{PCl}_3} \xrightarrow{\text{NH}_3} E \xrightarrow{-H_2O} \text{HCN}

The reaction \( C \xrightarrow{50\% \text{ KOH}} A + D \) is a Cannizaro reaction and thus the compound C is an aldehyde with no \( \alpha \)-hydrogen. C must be HCHO. From this it follows that A must be CH₃OH as its mild oxidation gives HCHO.

With this fact, we can write the given reactions as shown below.

\[ \text{CH}_3\text{OH} + \text{CH}_3\text{COOH} \rightarrow \text{CH}_3\text{COOCH}_2\text{CH}_3 \quad \text{methyl acetate} \quad \text{CH}_3\text{OH} \xrightarrow{\text{I}_2} \text{HCHO} \xrightarrow{50\% \text{ KOH}} \text{CH}_2\text{OH} + \text{HCOOH} \]

Hence

HCOOH \xrightarrow{\text{PCl}_5} \text{HOCI} \xrightarrow{\text{NH}_3} \text{HCONH}_2 \xrightarrow{-H_2O} \text{HCN}

A : CH₃OH \quad \text{methanol} \quad B : \text{CH}_3\text{COOCH}_3 \quad \text{methyl ethanoate}

C : HCHO \quad \text{methanal} \quad D : \text{HCOOH} \quad \text{methanoic acid}

E : \text{HCONH}_2 \quad \text{methanamide}
Iodoform Test

\[ \text{RCH}_2\text{CH}_3 + \frac{3}{2} \text{I}_2 + 4 \text{NaOH} \rightarrow \]

\[ \text{RCOO}^- \text{Na}^+ + 3 \text{NaI} + 3 \text{H}_2\text{O} + \text{CH}_3(\text{aq}) \]

\[ \text{HO}^- \]

\[ \text{RCH} \]

\[ \text{H} \]

\[ \text{I} \]

\[ \text{I} \]

\[ \text{I} \]

\[ \text{RCHOH} \rightarrow \text{RCOO}^- \text{Na}^+ \]

\[ \text{RCH}_2\text{CH}_3 + \frac{1}{2} \text{I}_2 + \text{NaOH} \rightarrow \text{RCOO}^- \text{Na}^+ + \text{CH}_3 \]

\[ \text{RCHCH}_3 + \frac{1}{2} \text{I}_2 + \text{NaOH} \rightarrow \text{RCOO}^- \text{Na}^+ + \text{CH}_3 \]

\[ \text{Cl}_3\text{C} + \text{OH}^- \rightarrow \text{CH}_3 + \text{Cl} + \text{RCOO}^- \]

This bond is broken.
Identify the compound

An organic compound X, on analysis gives 24.24 per cent carbon and 4.04 per cent hydrogen. Further, sodium extract of 1.0 g of X gives 2.90 g of silver chloride with acidified silver nitrate solution. The compound X may be represented by two isomeric structures, Y and Z. The compound Y on treatment with aqueous potassium hydroxide solution gives a dihydroxy compound, while Z on similar treatment gives ethanal. Find out the molecular formula of X and give the structures of Y and Z.

Solution  Mass per cent of chlorine in the organic compound X

\[
\text{Mass per cent of chlorine} = \frac{M_{\text{Cl}}}{M_{\text{AgCl}}} \times \frac{m_{\text{AgCl}}}{m_{\text{compound}}} \times 100 = \frac{35.5}{143.5} \times \frac{2.90}{1.0} \times 100 = 71.2\%
\]

The ratios of atoms in the molecule of X are

C : H : Cl = 24.24 : 4.04 : 71.72

\[\frac{12}{35.5} : \frac{1}{1} : \frac{2 : 4 : 2}{2 : 1} : \frac{1}{2} : 1 : 1 \]

Empirical formula of X is CH₂Cl₂

Since the isomer Y of the compound gives a dihydroxy compound on treating with aqueous KOH, it follows that the compound must contain two Cl atoms. Hence,

Molecular formula of X is C₃H₆Cl₂.

Its two isomers are \( \text{CH}_2\text{CHCl}_2 \) and \( \text{ClCH}_2\text{CH}_2\text{Cl} \).

The reaction are

\[
\text{CH}_2\text{CHCl}_2 \xrightarrow{\text{aq. KOH}} [\text{CH}_2\text{CH(OH)}_2] \rightarrow \text{CH}_3\text{CHO}
\]

\[
\text{ClCH}_2\text{CH}_2\text{Cl} \xrightarrow{\text{aq. KOH}} \text{HOCH}_2\text{CH}_2\text{OH}
\]

Match the entries given on the left with those given on the right.

<table>
<thead>
<tr>
<th>(a)</th>
<th>(b)</th>
<th>(c)</th>
<th>(d)</th>
<th>(e)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cyclopropane, chlorine and light</td>
<td>Propanone and sodium bisulphite</td>
<td>Ethanal and Methanal</td>
<td>Benzene, nitric acid and sulphuric acid</td>
<td>Propene, hydrogen bromide and a peroxide catalyst</td>
</tr>
<tr>
<td>(p)</td>
<td>(q)</td>
<td>(r)</td>
<td>(s)</td>
<td>(t)</td>
</tr>
<tr>
<td>Electrophilic substitution</td>
<td>Homologous pair</td>
<td>Homolytic addition</td>
<td>Free radical substitution</td>
<td>Nucleophilic addition</td>
</tr>
</tbody>
</table>

Ans :

(a) – (s);  (b) – (t);  (c) – (q);  (d) – (p);  (e) – (r)
Match the entries given on the left with those given on the right.
(a) Pyrolysis of alkanes
(b) Benzene + Chloroethane
(+ anhydrous AlCl₃)
(c) CH₂COOC₂H₅ + NaOH
(d) Preparation of alkanes
(e) Phenol + CHCl₃ (NaOH)
(f) C₂H₅Br + alcKOH
(p) Elimination reaction
(q) Saponification
(r) Wurtz reaction
(s) Friedel-craft reaction
(t) Reimer-Tiemann reaction
(u) Cracking
Ans :
(a) – (u);
(b) – (s);
(c) – (q);
(d) – (r);
(e) – (t);
(f) – (p)

Identify the compound

A mixture of two aromatic compounds A and B was separated by dissolving in chloroform followed by extraction with aqueous KOH solution. The organic layer containing compound A, when heated with alcoholic solution of KOH produced a compound C (C₆H₅N) associated with an unpleasant odour. The alkaline aqueous layer on the other hand, when heated with chloroform and then acidified gave a mixture of two isomeric compounds D and E of molecular formula C₆H₅O₂. Identify the compounds A, B, C, D, and E and write their structures.

Solution We have

Mixture of A and B

CHCl₃ + KOH

organic layer

Aqua layer

CHCl₃

H⁺

C₆H₅N (C)

unpleasant odour

C₆H₅O₂ (D)

C₆H₅O₂ (E)

The compound C is an isocyanide which is produced from a primary amine. Hence, the structures of C and A are

\[
\text{(C)} \quad \text{(A)}
\]

The compound B exhibits Reimer-Tiemann reaction. Thus, the compound B must be phenol. The reaction is

\[
\text{phenol} + \text{CHCl₃} \xrightarrow{1. \text{HCl}} \text{salicylaldehyde} + \text{H⁺}
\]
Rearrangement by migration of Bromine

\[
\text{Rearrangement, by migration of } \text{Br}, \text{ of initially formed } 1^\circ \text{ radical into more stable } 2^\circ \text{ or } 3^\circ \text{ radical.}
\]

Chlorination of either \(n\)-propyl or isopropyl bromide gives 1-bromo-2-chloropropane.

\[
\begin{align*}
\text{CH}_3\text{CH}_2\text{Br} & \quad \text{Cl}_2 \\
\text{CH}_3\text{Br}\text{CH}_3 & \quad \text{Cl}_2
\end{align*}
\]

Chlorination of either isobutyl or \textit{tert}-butyl bromide gives 1-bromo-2-chloro-2-methylpropane.

\[
\begin{align*}
\text{(CH}_3\text{)_2CBr} & \quad \text{Cl}^- \\
\text{(CH}_3\text{)_2CBr} & \quad \text{Cl}^-
\end{align*}
\]

Identify the Compound

An organic compound containing C, H and O exists in two isomeric forms A and B. A mass of 0.108 g of one of the isomers gives on combustion 0.308 g of CO\(_2\) and 0.072 g of H\(_2\)O. A is insoluble NaOH and NaHCO\(_3\) while B is soluble in NaOH. A reacts with concentrated H\(_2\)SO\(_4\) to give compounds C and D. C can be separated from D by the ethanolic AgNO\(_3\) solution and D is soluble in NaOH. B reacts readily with bromine to give compound E of molecular formula, C\(_x\)H\(_y\)O\(_z\). Identify A, B, C, D and E with justification and give their structures.

\textbf{Solution}

We have

\[
\text{Per cent of carbon in the compound} = \frac{M_C}{M_{\text{CO}_2}} \times \frac{m_{\text{CO}_2}}{m_{\text{compound}}} \times 100 = \left(\frac{12}{44}\right) \left(\frac{0.308}{0.108}\right) (100) = 77.78
\]

\[
\text{Per cent of hydrogen in the compound} = \frac{2M_H}{M_{\text{H}_2\text{O}}} \times \frac{m_{\text{H}_2\text{O}}}{m_{\text{compound}}} \times 100 = \left(\frac{2}{18}\right) \left(\frac{0.072}{0.108}\right) (100) = 7.41
\]

\[
\text{Per cent of oxygen in the compound} = 100 - (77.78 + 7.41) = 14.81.
\]

The ratios of atoms in the compound are

\[
\text{C : H : O} = \frac{77.78}{12} : \frac{7.41}{1} : \frac{14.81}{16} = 6.48 : 7.41 : 0.926 : 7 : 8 : 1.
\]

Hence, Empirical formula of the compound is C\(_6\)H\(_8\)O\(_2\).

Since the isomer B on reacting with bromine water gives compound E (C\(_x\)H\(_y\)O\(_z\)), the molecular formula of A and B will be the same as the empirical formula derived above, since both contain the same number of carbon atoms. As E is obtained from B by the substitution of hydrogen with bromine and since there is high carbon content in B, the compounds A and B must be aromatic. Now, since compound A is insoluble in NaOH and NaHCO\(_3\) and compound B is soluble in NaOH, it may be concluded that B is a phenolic compound and A is an ether. Hence, the structures of A and B are

\[
\begin{align*}
\text{(A) anisol} & \\
\text{(B) m-cresol}
\end{align*}
\]

The bromination of B gives
Identify the compound

Basic volatile Nitrogen compound

A basic volatile nitrogen compound gave a foul smelling gas when treated with chloroform and alcoholic potash. A 0.295 g sample of the substance, dissolved in aqueous HCl, and treated with NaNO₂ solution at 0 ℃ liberated a colourless, odourless gas whose volume corresponded to 112 mL at STP. After the evolution of the gas was complete, the aqueous solution was distilled to give an organic liquid which did not contain nitrogen and which on warming with alkali and iodine gave a yellow precipitate. Identify the original substance. Assume that it contains one N atom per molecule.

Solution: Since the compound gives a foul smelling gas on treating with CHCl₃ and alcoholic KOH, the compound must be a primary amine.

\[
\text{RNH}_2 + \text{CHCl}_3 + 3\text{KOH} \rightarrow \text{RNC} + 3\text{KCl} + 3\text{H}_2\text{O} \quad (1)
\]

Where RNC is the alkyl isocyanide (foul smelling gas).

Since the compound on treating with NaNO₂/HCl at 0 ℃ produces a colourless gas, the compound must be an aliphatic primary amine.

\[
\text{RNH}_2 + \text{HNO}_2 \rightarrow \text{ROH} + \text{N}_2 + \text{H}_2\text{O} \quad (2)
\]

Thus, the gas produced is nitrogen.

Amount of gas liberated = \(\frac{112 \text{ mL}}{22400 \text{ mL mol}^{-1}} = \frac{1}{200} \text{ mol}\)
From the above equation, it is obvious that

\[
\text{Amount of compound } RNH_2 = \frac{1}{200} \text{ mol}
\]

If \( M \) is the molar mass of \( RNH_2 \), then

\[
\frac{2.295 \text{ g}}{M} = \frac{1}{200} \text{ mol} \quad \text{or} \quad M = 0.295 \times 200 \text{ g mol}^{-1} = 59 \text{ g mol}^{-1}.
\]

Thus, the molar mass of alkyl group \( R \) is \((59 - 16) \text{ g mol}^{-1}, i.e. 43 \text{ g mol}^{-1}\). Hence, \( R \) must be \( \text{C}_3\text{H}_7 \).

From Eq. (2), it is obvious that the liquid obtained after distillation is ROH. Since this gives yellow precipitate with alkali and iodine (iodoform test), it must contain \( \text{CH}_3\text{C}==\text{CH}_3 \) group. Hence, it is concluded that ROH is

\[
\text{CH}_3\text{C}==\text{CH}==\text{CH}_3, \quad \text{Thus, the original compound is } \text{CH}_3\text{C}==\text{CH}==\text{CH}_3
\]

\[
\text{OH}
\]

\[
\text{OH}
\]

\[
\text{NH}_2
\]

Thus, \( \text{isopropylamine} \)

Match the entries given on the left with those given on the right.

(a) Lucas test (b) Neutral \( \text{FeCl}_3 \) (c) Dye test (d) Tollens test (p) Phenol (q) Glucose (r) Tertiary alcohol (s) Aniline

Ans:

(a) \( \rightarrow \) (r); \hspace{1cm} (b) \( \rightarrow \) (p); \hspace{1cm} (c) \( \rightarrow \) (s); \hspace{1cm} (d) \( \rightarrow \) (q)

Identify the compound

An organic compound \( A, \text{C}_6\text{H}_{10} \), on treatment with dilute sulphuric acid containing mercuric sulphate gives a compound \( B \), which can also be obtained from a reaction of benzene with an acid chloride in the presence of anhydrous aluminium chloride. The compound \( B \), when treated with iodine in aqueous \( \text{KOH} \), yields \( C \) and a yellow compound \( D \). Identify \( A, B, C \) and \( D \) with justification. Show how \( B \) is formed from \( A \).

**Solution**

The given reactions are

\[
\text{C}_6\text{H}_6 + \text{H}_2\text{SO}_4 + \text{C}_6\text{H}_5\text{Cl} + \text{acid chloride} \rightarrow \text{C}_6\text{H}_5\text{Cl} + \text{C}_6\text{H}_6
\]

The reaction of \( B \) with \( I \), in \( \text{KOH} \), is iodoform reaction. The compound \( D \) is iodoform, \( \text{CH}_3\text{I} \). The compound \( B \) must contain \( -\text{COCH}_3 \) group so as to exhibit iodoform reaction. Since \( D \) is obtained from benzene by Friedel-Crafts reaction, it is an aromatic ketone \( (\text{C}_6\text{H}_5\text{COCH}_3) \). The compound \( C \) must be an acid.

The compound \( A \) may be represented as \( \text{C}_6\text{H}_5\text{C}_2\text{H}_4 \). Since it gives \( \text{C}_6\text{H}_5\text{COCH}_3 \) on treating with dilute \( \text{H}_2\text{SO}_4 \) and \( \text{HgSO}_4 \), it must contain a triple bond \((-\text{C}==\text{CH})\) in the side chain. Hence, the given reactions may be represented as
Identify the compound

An organic compound E \((C_6H_8)\) on hydrogenation gives compound F\((C_6H_{12})\). Compound E on ozonolysis gives formaldehyde and 2-keto-propanal. Deduce the structure of compound E.

**Solution**

Since there are four hydrogen atoms added in the reaction

\[ C_6H_8 + 4H \rightarrow C_6H_{12} \]

the compound E may contain two double bonds. The ozonolysis reaction is

\[ C_6H_8 \rightarrow HCHO + CH_3C=CH \]

Since there is no loss of carbon atoms in an ozonolysis reaction, probably 2 molecules of formaldehyde are formed. In 2-ketopropanal, \(-C=CH_2\) and \(-CHO\) groups must have resulted due to the cleavage of double bonds. Hence, the net reaction can be explained as follows.

\[ H_2C=O + O=CH-CCH_3 \xrightarrow{\text{O}_3} CH_2=CH-CCH_3 \]

Hence, the compound E is 2-methyl-1, 3-butadiene.
Friedel Crafts Reaction

\[
\text{Friedel-Crafts Reaction}
\]

\[
\text{C}_6\text{H}_6 + \text{CH}_3\text{CH}_2\text{CH}_2\text{Cl} \xrightarrow{\text{AlCl}_3 \text{ at } -18^\circ \text{C to } 80^\circ \text{C}} \text{C}_6\text{H}_5\text{CH}_2\text{CH}_3 + \text{CH}_3\text{CH} = \text{CH}_2
\]

\[
\text{C}_6\text{H}_6 + \text{CH}_3\text{CHCH}_2\text{Cl} \xrightarrow{\text{AlCl}_3 \text{ at } -18^\circ \text{C to } 80^\circ \text{C}} \text{C}_6\text{H}_5 - \text{CH} = \text{CH}_3
\]

\[
\text{C}_6\text{H}_6 + \text{CH}_3\text{CH}_2\text{OH} \xrightarrow{\text{BF}_3 \text{ at } 60^\circ \text{C}} \text{C}_6\text{H}_5\text{CH} = \text{CH}_2\text{CH}_3
\]

Identify the compound

An Organic compound A has 76.6% C and 6.38% H (Aspirin)

An organic compound A has 76.6% C and 6.38% H. Its vapour density is 47. It gives characteristic colour with FeCl₃ solution. The compound A when treated with CO₂ and NaOH at 140 °C under pressure gives B which on being acidified gives C. The compound C reacts with acetyl chloride to give D which is a well known pain killer. Identify A, B, C and D and explain the reactions involved.

**Solution.** The ratios of atoms present in one molecule of A are

- C : H : O :: 76.6 : 6.38 : 17.02
- C : H : O :: 12 : 1 : 16

Hence, empirical formula of A is C₆H₆O. Molar empirical formula mass of A = 94 g mol⁻¹. Thus, molecular formula of A is C₆H₆O. Since, the compound A gives characteristic colour with FeCl₃, it must be a phenol.

The given reactions are

\[
\text{(A)} \quad \text{OH} + \text{CO}_2 \xrightarrow{\text{NaOH \ 140^\circ \text{C, high pressure}}} \text{(B)} \quad \text{OH} \quad \text{COONa} \xrightarrow{\text{H}^+} \text{(C)} \quad \text{OH} \quad \text{COOH}
\]

Aspirin (pain killer)
Identify the compound

An organic compound A of molar mass 140.5 gm per mole has 68.32% Hydrogen

An organic compound A of molar mass 140.5 g mol\(^{-1}\) has 68.32% C, 6.4% H and 25.26% Cl. Hydrolysis of A with dilute acid gives compound B, C\(_6\)H\(_5\)O. Compound B can be oxidized under mild conditions to compound C, C\(_6\)H\(_5\)O. Compound C forms a phenylhydrazone D with PhNHNH\(_2\) and gives a positive iodoform test. Deduce the structures of compounds A to D with proper reasoning.

**Solution**

The ratios of atoms in a molecule of A are

\[
\begin{align*}
C : H : Cl & = 68.32 : 6.4 : 25.26 \\
& = 5.69 : 0.71 : 0.355 \\
& \approx 8 : 9 : 1
\end{align*}
\]

Hence, the empirical formula of A is C\(_8\)H\(_9\)Cl. 

Molar empirical formula mass of A = 140.5 g mol\(^{-1}\)

Thus, molecular formula of A is C\(_{14}\)H\(_{18}\)Cl.

The high content of carbon indicates the presence of aromatic (or benzene) ring in the molecule.

Since Cl atom in the compound A is easily replaced during hydrolysis of A with dilute acid, the Cl atom must be attached to the aliphatic portion of the molecule.

Since the compound C gives a positive iodoform test, it must contain—COCH\(_3\) group.

Since the compound C is obtained by mild oxidation of B, the latter must be a secondary alcohol. Hence, the compound A also contains secondary chlorine atom. Consistent with this, the given reactions are as follows.

\[
\begin{align*}
\text{Cl} & \quad \text{OH} \\
\text{H} - \text{C} - \text{CH}_3 & \quad \text{H} - \text{C} - \text{CH}_3 \\
(A) & \quad \text{B} \\
\end{align*}
\]

\[
\begin{align*}
\text{B} & \quad \text{[O]} \\
\text{H} - \text{C} - \text{CH}_3 & \quad \text{O} - \text{C} - \text{CH}_3 \\
\text{C} & \quad \text{D} \\
\end{align*}
\]

\[
\begin{align*}
\text{D} & \quad \text{PhNHNH}_2 \\
\text{CH}_3 - \text{C} & \quad \text{NNHPh} \\
\end{align*}
\]
Question

Bleaching powder and bleach solution are produced on a large scale and used in several household products. The effectiveness of bleach solution is often measured by iodometry. 25 mL of household bleach solution was mixed with 30 mL of 0.50 M KI and 10 mL of 4 N acetic acid. In the titration of the liberated iodine, 48 mL of 0.25 N Na₂S₂O₃ was used to reach the end point. The molarity of the household bleach solution is

(a) 0.48 M  (b) 0.96 M  (c) 0.24 M  (d) 0.024 M

Bleaching powder contains a salt of an oxoacid as one of its components. The anhydride of that oxoacid is

(a) Cl₂O  (b) Cl₂O₃  (c) ClO₂  (d) Cl₂O₆

Solutions:

For Q 11, we may write

Household bleach + 2 KI → I₂ + products

I₂ + 2Na₂S₂O₃ → Na₂S₄O₆ + 2NaI

Amount of Na₂S₂O₃ used = \( \frac{1}{2} \) \( \frac{V}{M} \) = \( 48 \times 10^{-3} \text{ L} \) \( \times \frac{0.25 \text{ mol L}^{-1}}{1} \) = \( 12 \times 10^{-3} \text{ mol} \)

Amount of I₂ generated = \( \frac{1}{2} \) \( 12 \times 10^{-3} \text{ mol} \) = \( 6 \times 10^{-3} \text{ mol} \)

Assuming 1 mol of household bleach produces 1 mol I₂, we will have

Amount of household bleach in 25 mL solution = \( 6 \times 10^{-3} \text{ mol} \)

Molarity of household bleach = \( \frac{n}{V} \) = \( \frac{6 \times 10^{-3} \text{ mol}}{25 \times 10^{-3} \text{ L}} \) = 0.24 M

For Q 12, we have

Bleaching powder contains Ca(OCl)Cl.

The oxoacid of this salt is HOCl. The formation of this acid through its anhydride is \( \text{Cl}_2\text{O} + \text{H}_2\text{O} \rightarrow 2\text{HOCl} \)

Hence, we have

The choice (c) is correct.
The choice (a) is correct.

Match the compounds/ions in Column I with their properties/reactions in Column II.

<table>
<thead>
<tr>
<th>Column I</th>
<th>Column II</th>
</tr>
</thead>
<tbody>
<tr>
<td>(a) C₂H₄CHO</td>
<td>(p) gives precipitate with 2,4-dinitrophenylhydrazine</td>
</tr>
<tr>
<td>(b) CH₃C≡CH</td>
<td>(q) gives precipitate with AgNO₃</td>
</tr>
<tr>
<td>(c) CN⁻</td>
<td>(r) is a nucleophile</td>
</tr>
<tr>
<td>(d) I⁻</td>
<td>(s) is involved in cyanohydrin formation</td>
</tr>
</tbody>
</table>

Ans:

(a) \(-\) (p), (q), (s);
(b) \(-\) (q);
(c) \(-\) (q), (r), (s);
(d) \(-\) (q)
In decreasing order of Oxidation state of Nitrogen

Which ordering of compounds is according to the decreasing order of the oxidation state of nitrogen?
(a) HNO₃, NO, NH₄Cl, N₂  
(b) HNO₃, NO, N₂, NH₄Cl  
(c) HNO₃, NH₄Cl, NO, N₂  
(d) NO, HNO₃, NH₄Cl, N₂  

**Solution:**
The oxidation states of nitrogen in the given compounds are as follows.
HNO₃: +1 + x + 3 (−2) = 0  ⇒  x = +5
NO:  x + (−2) = 0  ⇒  x = +2  
NH₄Cl: x + 4 (+1) + (−1) = 0  ⇒  x = −3
N₂:  2x = 0  ⇒  x = 0  
Thus, the decreasing oxidation numbers of nitrogen is HNO₃, NO, N₂ and NH₄Cl.
Therefore, the **choice (b)** is correct.

Gyan Question

The Kinetic energy of an electron in the second Bohr orbit

The kinetic energy of an electron in the second Bohr orbit of a hydrogen atom (\(a₀\) is Bohr radius) is
(a) \(\frac{\hbar^2}{4\pi^2ma₀^2}\)  
(b) \(\frac{\hbar^2}{16\pi^2ma₀^2}\)  
(c) \(\frac{\hbar^2}{32\pi^2ma₀^2}\)  
(d) \(\frac{\hbar^2}{64\pi^2ma₀^2}\)  

**Solution:**
Bohr model of an atom satisfies the following two requirements.
• Equality of centripetal and centrifugal forces, i.e.

\[
\frac{(Ze)(c)}{(4\pi\varepsilon_0)r^2} = \frac{ma^2}{r}
\]

• Quantization of angular momentum, i.e.

\[
mvr = n\left(\frac{\hbar}{2\pi}\right)
\]

Eliminating \(v\) in these two expressions, we get

\[
r = n^2\left[\frac{\hbar^2}{4\pi^2m(Ze/4\pi\varepsilon_0)}\right] = \frac{n^2}{Z}a₀
\]

where \(a₀\) is the Bohr radius. The kinetic energy of the electron in Bohr orbit of an atom is

\[
KE = \frac{1}{2}mv^2 = \frac{1}{2}m\left[\frac{n}{mv}\left(\frac{\hbar}{2\pi}\right)\right]^2 = \frac{1}{2}m\left[\frac{n}{m(n^2a₀/Z)}\left(\frac{\hbar}{2\pi}\right)\right]^2 = \frac{Z^2}{n^2}\left(\frac{\hbar^2}{8\pi^2ma₀^2}\right)
\]

For the second Bohr orbit of hydrogen atom, \(Z = 1\), and \(n = 2\). Hence

\[
KE = \frac{1}{32}\left(\frac{\hbar^2}{\pi^2ma₀^2}\right)
\]

Therefore, the **choice (c)** is correct.
Question on Aldol reaction

The number of aldol reaction(s) that occurs in the given transformation

\[ \text{CH}_3\text{CHO} + 4\text{HCHO} \xrightarrow{\text{conc. aq. NaOH}} \]

is

(a) 1  (b) 2  (c) 3  (d) 4

**Solution:**

The given reaction may be formulated as follows.

- **First aldol condensation**

![First aldol condensation]

- **Second aldol condensation**

![Second aldol condensation]

- **Third aldol condensation**

![Third aldol condensation]

- **Cannizzaro reaction**

![Cannizzaro reaction]

There are three aldol condensations and one Cannizzaro reaction. Therefore, the choice (c) is correct.

An organic compound undergoes first order decomposition

An organic compound undergoes first-order decomposition. The time taken for its decomposition to 1/8 and 1/10 of its initial concentration are \(t_{1/8}\) and \(t_{1/10}\), respectively. What is the value of \((t_{1/8}/t_{1/10})\)? (log 2 = 0.3)

**Solution:**

For first-order decomposition of \(A\), the rate law is \(\ln \left(\frac{[A]}{[A]_0}\right) = -kt\)

Hence \[ \ln(1/8) = -kt_{1/8} \quad \text{and} \quad \ln(1/10) = -kt_{1/10} \]

Thus \[ \frac{\ln(1/8)}{\ln(1/10)} = \frac{t_{1/8}}{t_{1/10}} \]

Hence \( (t_{1/8}/t_{1/10}) = \log 8 = \log 2^3 = 3 \log 2 = 3 \times 0.3 = 0.9 \) or \( 10(t_{1/8}/t_{1/10}) = 10 \times 0.9 = 9 \)

Therefore, the answer is 9.
Allene has sp hybridization

In allene (C$_3$H$_4$), the type(s) of hybridization of the carbon atom is (are)
(a) sp and sp$^3$  (b) sp and sp$^2$  (c) sp$^3$ only  (d) sp$^3$ and sp$^3$

**Solution:**

The structure of allene is

In this molecule, carbon atoms 1 and 3 are sp$^2$ hybridized while the carbon atom 2 is sp hybridized. Therefore, the choice (b) is correct.

Number of chiral carbon and optically active products

The number of optically active products obtained from the complete ozonolysis of the given compound

CH$_3$-CH=CH-C=CH=CH-C=CH=CH-CH$_3$

(a) 0  (b) 1  (c) 2  (d) 4

**Solution:**

Ozonolysis

CH$_3$-CH$_2$=CH-C-CH$_2$-CH=CH$_2$-CH$_3$ → CH$_3$CHO + OCH$_3$ + OCH$_3$ + C=CHO

None of the products will be optically active. Therefore, the choice (a) is correct.
Carboxylic functional group is present in aspirin

The carboxylic functional group (—COOH) is present in:
(a) picric acid (b) barbituric acid (c) ascorbic acid (d) aspirin

Solution:
The structures of the given compounds are as follows.

Picric acid

Barbituric acid

Ascorbic acid

Aspirin

Therefore, the choice (d) is correct.

Identify the compound

In the following reaction sequence, the compound J is an intermediate.

\[ \text{I} \xrightarrow{(\text{CH}_2\text{CO})_2\text{O}} \xrightarrow{\text{CH}_2\text{COONa}} \text{J} \xrightarrow{\text{H}_2\text{Pd/C}} \xrightarrow{\text{(ii) SOCl}_2 \text{, (iii) anhyd. AlCl}_3} \text{K} \]

J(C_7H_8O_2) gives effervescence on treatment with NaHCO_3 and positive Baeyer’s test.

Q - The compound K is

(a) ![Compound A](image1)
(b) ![Compound B](image2)
(c) ![Compound C](image3)
(d) ![Compound D](image4)

Q - The compound I is

(a) ![Compound E](image5)
(b) ![Compound F](image6)
(c) ![Compound G](image7)
(d) ![Compound H](image8)
Perkin Condensation

_Solutions_

The compound J must contain a —COOH group as it gives effervescence with NaHCO₃. Also, it is an unsaturated compound as it gives positive Baeyer’s test (decolourization of pink colour of alkaline KMnO₄). Since the compound J is obtained by treating the compound I with (CH₃CO)₂O and CH₃COONa, and more over the compound J seems to be aromatic (high carbon content), the compound I amongst the choices given in Q.14 seems to be benzaldehyde as it shows Perkin condensation shown in the following

\[
\text{CHO} + (\text{CH₃CO})₂\text{O} \xrightarrow{\Delta} \text{CH₂COO} \xrightarrow{\text{Na}} \alpha, \beta-\text{unsaturated carboxylic acid}
\]

The conversion J to K is as follows.

- **CHO**
- **CH₂COOH**
- **CH₂CH₂COOH**
- **CH₃COOH**

Hence, we have

- The choice (c) is correct.
- The choice (a) is correct.

**Question on Isomers**

Which of the given statement(s) about N, O, P and Q with respect to M is(are) correct?

- (a) M and N are non-mirror image stereoisomers.
- (b) M and O are identical.
- (c) M and P are enantiomers.
- (d) M and Q are identical.

**Solution**

First of all, we may draw Fischer projections of the given compounds.
Aldohexose exists in D configuration

When the following aldohexose exists in D-configuration, the total number of stereoisomers in its pyranose form is

\[
\begin{align*}
&\text{CHO} \\
&\text{CH}_2 \\
&\text{CHOH} \\
&\text{CHOH} \\
&\text{CHOH} \\
&\text{CH}_2\text{OH}
\end{align*}
\]

**Solution:**

In D-configuration, the hydroxyl group at the asymmetric carbon placed at the bottom of Fischer projection lies to the right of the straight skeleton. Thus, we will have
There are four chiral carbon atoms in the pyranose form of the given aldohexose. Thus, there will be 16 (= 2⁴) stereoisomers, out of which 8 are of D-configurations (CH₂OH group above the ring) and 8 are of L-configurations (CH₂OH group below the ring).
Therefore, the correct answer is 8.

Peptides

The substituents R₁ and R₂ for nine peptides are listed in the table given below. How many these peptides are positively charged at pH = 7.0?

<table>
<thead>
<tr>
<th>Peptide</th>
<th>R₁</th>
<th>R₂</th>
</tr>
</thead>
<tbody>
<tr>
<td>I</td>
<td>H</td>
<td>H</td>
</tr>
<tr>
<td>II</td>
<td>H</td>
<td>CH₃</td>
</tr>
<tr>
<td>III</td>
<td>CH₂COOH</td>
<td>H</td>
</tr>
<tr>
<td>IV</td>
<td>CH₂CONH₂</td>
<td>(CH₂)₂NH₂</td>
</tr>
<tr>
<td>V</td>
<td>CH₂CONH₂</td>
<td>CH₂CONH₂</td>
</tr>
<tr>
<td>VI</td>
<td>(CH₂)₂NH₂</td>
<td>(CH₂)₂NH₂</td>
</tr>
<tr>
<td>VII</td>
<td>CH₂COOH</td>
<td>CH₂CONH₂</td>
</tr>
<tr>
<td>VIII</td>
<td>CH₂OH</td>
<td>(CH₂)₂NH₂</td>
</tr>
<tr>
<td>IX</td>
<td>(CH₂)₂NH₂</td>
<td>CH₃</td>
</tr>
</tbody>
</table>

Solution

In general, the pH of the solution at which amino acids exist as zwitterion follows the order:

Acidic side chain < neutral chain < basic side chain  
(pH = 3)  (pH = 5.5 - 6.0)  (pH = 9 - 10)

This pH is known as isoelectric point.

At pH = 7, the acidic side chain (for which isoelectric point < 7) exists as a negatively-charged species (e.g., —COO⁻) and the basic side chain (for which isoelectric point > 7) exists as a positively charged species (e.g., —NH₃⁺). Since peptides IV, VI, VIII and IX contain —NH₂ group in R₁ and/or R₂ group(s), these are expected to exist as positively-charged species.

Therefore, the correct answer is 4.
Try to figure out what sequence is depicted

neopentane < isopentane < n-pentane. A branched chain isomer has a lower boiling point than a straight chain isomer. The more numerous the branches, the lower the boiling point. The branching lowers the boiling point for all families of organic compounds. Branching causes the shape of molecule to approach towards a sphere and thus surface area decreases, with the result that the intermolecular forces become weaker and are overcome at a lower temperature.

n-butyl alcohol < isobutyl alcohol < sec-butyl alcohol < tert-butyl alcohol. Polarity of O—H bond increases in the same order.

(CH₃)₂CCOOH < (CH₃)₂CHCOOH < CH₃CH₂COOH < CH₃COOH < HCOOH

The presence of bulky groups near the site of reaction, whether in the alcohol or in the acid, slows down esterification as well as de-esterification (hydrolysis).

(CH₃)₂CCOOCH₃ < (CH₃)₂CHCOOCH₃ < CH₃COOCH₃ < HCOOCH₃
CH₂COOC(CH₃)₂ < CH₃COOCH(CH₃)₂ < CH₃COOC₂H₅ < CH₃COOCH₃

Try to figure out the sequences

adipic acid < succinic acid < malonic acid < oxalic acid
Acidity decreases with increase in the intervening CH₂ groups.

isobutane < n-butane < n-butyl chloride < n-butanol
chlorobenzene < benzene < toluene < methoxybenzene
Decreasing order of reactivity towards alcoholic silver nitrate:
2-bromo-1-phenylethene, α-phenylethyl bromide, β-phenylethyl bromide
Increasing order of reactivity towards aqueous NaOH:
chlorobenzene, m-chloronitrobenzene, o-chloronitrobenzene, 2,4-dinitrochlorobenzene,
2,4,6-trinitrochlorobenzene
Increasing reactivity towards HCN: CH₃CHO, CH₃COCH₃, HCHO, C₂H₅COCH₃
Increasing basicity: p-toluidine, N, N-dimethyl-p-toluidine, p-nitroaniline, aniline.
Increasing ease of hydrolysis: CH₃COOC₂H₅, CH₃COCl, (CH₃CO)₂O, CH₃CONH₂
Increasing order of acid strength:
CICH₂COOH, CH₃CH₂COOH, CICH₂CH₂COOH, (CH₃)₂CHCOOH, CH₃COOH
Increasing reactivity in nucleophilic substitution reactions: CH₃F, CH₃I, CH₃Br, CH₃Cl

major product H in the given reaction sequence

The major product H in the given reaction sequence

\[
\text{CH}_3\text{-CH}_2\text{-CO-CH}_3 + \text{HCN} \xrightarrow{95\% \text{ H}_2\text{SO}_4, \text{Heat}} \text{H}
\]

is

(a) CH₃-CH=C-COOH
(b) CH₃-CH=C-CN
(c) CH₃-CH₂-C-COOH
(d) CH₃-CH=C-CO-NH₂

Solution:
The first reaction is nucleophilic addition reaction across the -C=O bond.

In the second reaction, -CN is hydrolysed to -COOH. The resultant molecule undergoes dehydration on heating.

Therefore, the choice (a) is correct.
Decreasing order of acidity of substituted benzoic acids:
  p-chlorobenzoic acid, 2, 4-dichlorobenzoic acid, 2,4,6-trichlorobenzoic acid.
Decreasing order of acidity of carboxylic acids:
  α-chlorophenylacetic acid, p-chlorophenylacetic acid, phenylacetic acid, α-phenylpropionic acid
Decreasing order of acidity of carboxylic acids:
  p-nitrobenzoic acid, p-nitrophenylacetic acid, β-(p-nitrophenyl) propionic acid
Increasing order of basicity:
  ammonia, aniline, cyclohexylamine
Decreasing order of basicity:
  ethylamine, 2-aminoethanol, 3-amino-1-propanol
Decreasing order of basicity:
  aniline, p-methoxyaniline, p-nitroaniline
Increasing order of acidity:
  benzene sulphonic acid, benzoic acid, benzyl alcohol, phenol
Decreasing order of acidity:
  m-bromophenol, m-cresol, m-nitrophenol, phenol
Decreasing order of acidity of substituted phenol:
  p-chlorophenol, 2, 4-dichlorophenol, 2, 4, 6-trichlorophenol
Decreasing order of reactivity towards KCN:
  benzyl chloride, chlorobenzene, ethyl chloride
Increasing order of nitration:
  benzene, chlorobenzene, nitrobenzene, toluene
Increasing order of reactivity towards alcoholic silver nitrate:
  1-bromo-1-butene, 3-bromo-1-butene, 4-bromo-1-butene

Explanation by solving in right order:

Increasing reactivity towards S_N1 solvolysis:
  benzyl chloride, p-chlorobenzyl chloride, p-methoxybenzyl chloride, p-methylbenzyl chloride,
  p-nitrobenzyl chloride
Increasing order of reactivity towards elimination by alcoholic KOH:
  1-phenyl-2-bromopropane, 1-phenyl-3-bromopropane
Decreasing order of reactivity towards aqueous HBr: Isomeric pentyl alcohols
Increasing order of reactivity towards aqueous HBr:
  1-phenyl-1-propanol, 3-phenyl-1-propanol, 1-phenyl-2-propanol
Decreasing order of reactivity towards aqueous HBr:
  benzyl alcohol, p-cyanobenzyl alcohol, p-hydroxybenzyl alcohol
Increasing order of reactivity towards aqueous HBr:
  benzyl alcohol, diphenylmethanol, methanol and triphenylmethanol
Decreasing order of reactivity towards bromination:
  anisole, benzene, toluene, chlorobenzene, nitrobenzene, phenol
Increasing order of reactivity towards bromination:
  hydroquinone, p-methoxyphenol, p-methylphenol, p-chlorophenol, p-nitrophenol, sym-trihydroxybenzene
Decreasing order of acidity of carboxylic acids:
  butanoic acid, 2-bromobutanoic acid, 3-bromobutanoic acid, 4-bromobutanoic acid

Decreasing reactivity towards S_N1 substitution:
  1-chloropropane 3-chloropropane, n-propylchloride
Decreasing acidity: H_2O, HO==CH, NH_3, RH, ROH
Decreasing basicity: R', HO==C', NH_2, OH', OR'
Decreasing order of reactivity towards the addition of HCl:
  styrene, p-chlorostyrene, p-methylstyrene, p-nitrostyrene
Decreasing order of reactivity towards dehydration:
  α-phenyl ethyl alcohol, α-(p-nitrophenyl) ethyl alcohol, α-(p-aminophenyl) ethyl alcohol
Various Increasing Properties

In the reaction of free radicals:
- CH₃, 1°, 2°, 3°, allyl, vinyl

In the reaction of carbocation:
- CH⁺, 1°, 2°, 3°

Increasing enthalpy of reaction:
\[
\begin{align*}
\text{C}_3\text{H}_7\text{Br} & \rightarrow \text{C}_3\text{H}_7^+ + \text{Br}^- & \Delta H_1 \\
\text{C}_3\text{H}_8\text{Br} & \rightarrow \text{C}_3\text{H}_7\text{H}^+_2 + \text{Br}^- & \Delta H_2 \\
\text{C}_3\text{H}_8\text{H} & \rightarrow \text{C}_3\text{H}_7\text{H} + \text{Br}^- & \Delta H_3 \\
\text{C}_3\text{H}_8\text{C} & \rightarrow \text{C}_3\text{H}_7\text{H} + \text{Br}^- & \Delta H_4
\end{align*}
\]

Increasing order of reactivity towards Sₐ2 displacement:
1-bromopentane, 2-bromopentane, 2-bromo-2-methylbutane

The compound that undergoes decarboxylation most readily under mild condition is

\[\text{(a)} \quad \text{COOH} \quad \text{CH}_2\text{COOH} \quad \text{COOH} \quad \text{CH}_2\text{COOH} \quad \text{COOH} \quad \text{CH}_2\text{COOH}\]

**Solution:**

β-Ketocarboxylic acid is unstable acid. It readily undergoes decarboxylation through a cyclic transition state.

Therefore, the choice (b) is correct.

A compound contains 88.89% C and 11.1% H. It gives white precipitate with ammoniacal silver nitrate. The compound is

(a) C₅H₅C ≡ CH \quad (b) C₅H₆C ≡ CH \quad (c) C₆H₇C ≡ CH \quad (d) C₆H₈C ≡ CH

**Ans:**
More the number of electron withdrawing group stronger is the acid

\[ \text{Cl}_2\text{CCOOH} > \text{Cl}_2\text{CHCOOH} > \text{ClICH}_2\text{COOH} > \text{CH}_3\text{COOH} \]

The more the number of electron-withdrawing group, the stronger the acid.

Try to figure out what sequence is depicted

\[ \text{FCH}_2\text{COOH} > \text{ClCH}_2\text{COOH} > \text{BrCH}_2\text{COOH} > \text{ICH}_2\text{COOH} \]

The more the electronegativity of halogen, the stronger the acid.

\[ \text{CH}_3\text{COOH} < \text{ClCH}_2\text{CH}_2\text{COOH} < \text{ClICH}_2\text{COOH} \]

\[ \text{HO} - \text{COOH} < \text{H}_2\text{C} - \text{COOH} < \text{C}_6\text{H}_5\text{COOH} < \text{Br} - \text{COOH} \]

\[ \text{H}_2\text{O} < \text{C}_2\text{H}_5\text{OH} < \text{C}_6\text{H}_5\text{COOH} < \text{HCOOH} < \text{HCl} \]

phenol < p-nitrophenol < HCOOH < C6H5COOH

C6H5NH2 < C6H5NCH3 < NH3 < CH3NH2 < (CH3)2NH

(CH3)3N is the weakest because of steric factors.

1-butanol < 2-butanol < 2-methyl-2-propanol. A tertiary alcohol reacts immediately, a secondary alcohol reacts within five minutes and a primary alcohol does not.

2-methyl-2-propanol < 2-butanol < 1-butanol. Reactivity of ROH is CH3OH > 1° > 2° > 3°.

An organic compound A contains 49.32% C, 9.59% H and 19.18% N and rest oxygen. The compound A on boiling with NaOH gives nitrogen free carboxylic acid whose silver salt contains 59.67% Ag. The compound A is

(a) CH3CONH2  (b) C2H5CONH2  (c) C3H7CONH2  (d) C4H9CONH2

Ans:

(b) C2H5CONH2

An organic compound A contains 49.32% C, 9.59% H and 19.18% N and rest oxygen. The compound A on boiling with NaOH gives nitrogen free carboxylic acid whose silver salt contains 59.67% Ag. The compound A is

(a) CH3CONH2  (b) C2H5CONH2  (c) C3H7CONH2  (d) C4H9CONH2

Hence \( \frac{108}{M_R + 44 + 108} = 0.5967 \). This gives \( M = 29 \) i.e. \( R = \text{C}_2\text{H}_5 \).

Hence, the compound is C2H5CONH2.
An aromatic compound contains 69.4% C and 5.8% H.

An aromatic compound contains 69.4% C and 5.8% H. A sample of 0.3 g of this compound gave ammonia which neutralizes 25 mL of 0.05 M H₂SO₄. The empirical formula of the compound is

(a) C₃H₅N₂O
(b) C₅H₇NO
(c) C₆H₇NO₂
(d) C₆H₆NO

Ans:

Amount of H₂SO₄ neutralized = \((25 \times 10^{-3} \text{ L})(0.05 \text{ M})\) = \(1.25 \times 10^{-3} \text{ mol}\)
The neutralization reaction is \(2\text{NH}_3 + \text{H}_2\text{SO}_4 \rightarrow (\text{NH}_4)_2\text{SO}_4\)

Amount of NH₃ evolved = \(2 \times 1.25 \times 10^{-3} \text{ mol} = 2.5 \times 10^{-3} \text{ mol}\)

Mass of N in the compound = \((2.5 \times 10^{-3} \text{ mol})(14 \text{ g mol}^{-1}) = 0.035 \text{ g}\)

Per cent of N in the compound = \(\frac{0.035}{0.3} \times 100 = 11.67\%

Per cent of O in the compound = \(100 - (69.4 + 5.8 + 11.67) = 13.13\%

Ratio of atoms in the compound is \(\text{C : H : N : O} = 69.4 : 5.8 : 11.67 : 13.13\)

Hence, empirical formula: \(\text{C}_6\text{H}_7\text{NO}\)
Alternatively, calculate per cent of C in the given choices which comes out to be (a) 0.55, (b) 0.69, (c) 0.58 and (d) 0.65 only for choice (b), the answer tallies.
Zwitter Ions - The Hydrogen atom (rather ion) moves to different positions depending on the pH (So can have a positive charge at some position, or a negative charge at some other position, depending on the pH)

Electrophoresis - is a method to confirm if a positive charge is present or a negative charge is present in the ion.

Iso-Electric point - the pH at which the ion (molecule) does not migrate to any electrodes.

What happens if oxalic acid is heated with conc sulphuric acid?

We get Carbonmonoxide

\[
\text{COOH} + (\text{H}_2\text{SO}_4)\text{(I)} \rightarrow \text{H}_2\text{O} + \text{CO}_2\text{(g)} + \text{CO}_2\text{(g)} + (\text{H}_2\text{SO}_4)\text{(I)}
\]

IMPORTANT ORDER AND FACTS OF ORGANIC CHEMISTRY

1. RCOCl > RCOCOR > RCOOR > RCONH₂ Nucleophilic substitution reaction.
2. HI > HBr > HCl > RCOOH > C₆H₅OH > H₂O > CH≡CH > NH₃ (Acidic nature).
3. CCl₃CHO > HCHO > CH₂CHO > CH₃COH₂ Nucleophilic addition reaction.

4. CH₂ = CH₂ > CH ≡ CH > C₂H₆ Electrophilic addition reaction.

5. Substitution Reaction

6. Nucleophilic Substitution Reaction

7. (D)₃C - Cl > (C₂H₅)₃C - Cl > C₂H₅Cl > CH₂Cl > CH₂ = CH₂ > C₂H₅Cl > CH₂ = CHCl

8. (D)₃C > (C₂H₅)₃C > C₂H₅CH₂ > CH₂ = CH₂ > CH₂ = CH₂ > C₃H₅Cl > CH₂Cl > C₂H₅Cl

9. (D)₃C > CH₂ = CH₂ > CH₂ = CH₂ > CH₂ = CH₃ > CH₃ > CH₂Cl > C₂H₅Cl

10. (CH₃)₂C = C(CH₃)₂ > CH₃ > CH = C > (CH₃)₂ > CH₃ > CH = C > CH₂CH₃ > CH₃ > CH₂ = CH₂ > C₂H₅Cl > CH₂Cl > C₂H₅Cl

11. CH₂ = CH₂ > CH₃ > CH = CH₂ > CH₂ = CH₃ > CH₂ = CH₂ > CH₂ = CH₂ > CH₂ = CH₂ > CH₂ = CH₂ > CH₂ = CH₂ > CH₂ = CH₂ > CH₂ = CH₂ > CH₂ = CH₂

12. NH₃ > OH > NH₃ > H₂O (Basic strength)

13. NI₃ > NBr₃ > NCl₃ > NF₃ (Basic strength)

14. Br₂ > Cl₂ > I₂ (Selectivity for halogenation)

15. Halogenation of alkenes by cyclic halonium state, so anti attack takes place.

16. Hydroboration followed by oxidation is always anti markownikoff’s addition due to steric effect.

17. Oximercuration - demercuration is markownikoff’s addition of water because some carbocation character in cyclic mercurium state.

18. CHCl₃ in the presence of strong bases forms biradical : CCl₂ which undergo addition with double or triple bonds.

19. When conjugated diene reacts with alkene or alkyne it is known as diel’s elder synthesis.

20. Ozonolysis of cyclo alkene forms one mole dialdehyde while ozonolysis of cyclo alkadiene forms two moles of dialdehyde.

21. Ozonolysis with (CH₃)₂S is known as reductive ozonolysis.

22. Hydration of alkyne occur’s in HgSO₄ and dil H₂SO₄.
23. 1-alkynes forms ppt with Ag(NH₃)₂⁺ and Cu(NH₃)₄²⁺.

24. Cis-2-butene reacts with Br₂ to form dl(±) pair of enantiomers of 2,3-dibromobutane while in case of trans-2-butene forms meso-2,3-dibromo butane due to anti addition always.


26. Chloral reacts with chloroform group to form insecticide DDT.

27. NBS is used for free radical allylation.

28. Rate for S¹ reaction is 3° > 2° > 1° in protic polar solvent.

29. Rate for S² reaction is 1° > 2° > 3° in polar aprotic solvent like DMSO, DMF, HMPT.

30. Chemical reactions like Hoffmann carbamidine and Reimer Tiemann’s reaction active species is biradical CCl₂.

31. If cyclo 1,3-penta diene reacts with CHCl₃ and potassium tert. butoxide to form chlorobenzene.

32. Alkyl halides reacts with AgCN to form isocyanides due to ambident nature of nucleophile, other ambident nucleophiles are $\text{NO}_2^-$ and $\text{SO}_3^{2-}$.

33. In dehydration of alcohols active species is carbocation so rearrangement occurs like hydride shift or alkyl shift.

34. Dehydration of cyclobutyl methyl alcohol ring expansion takes place, formation of cyclo pentene occurs.

35. In esterification where acid reacts with alcohol to form ester, -OH given by acid while -H by alcohol

$$\text{CH}_3\text{COOH} + \text{C}_2\text{H}_5\text{OH} \xrightarrow{\text{H}^+} \text{CH}_3\text{COOC}_2\text{H}_5 + \text{H}_2\text{O}$$

36. Ether’s reacts with HI to form alcohol and halide where fission of lower ether by S² mechanism while higher ethers like tert. butyl methyl ether or alkyl methyl ether by S¹ mechanism.

37. Quantitative estimation of ethers is done by ziesal’s method.

38. If unsym. cyclic ether undergo fission it depends upon medium weather it is acid or basic like in acidic medium some character of carbocation so nucleophile goes to carbon where more alkyl groups are there while reverse in basic medium due to steric factor.

39. Aldehydes are reducing agent while ketones are not.

40. Aldehydes and ketones are separated by tollen’s reagent.

41. Carbonyl and noncarbonyl are separated by sodium bi sulphite and bradye’s reagent.

42. As the size of alkyl group increases steric hindrance comes into play, reactivity towards nucleophilic addition decreases.

43. Aldehydes with hydrogen atom in the presence of dil base undergo enolization and form $\text{C}=\text{O}$.
44. If there is two-CHO group with $\equiv$hydrogen atom to form cyclic intra aldol product.

45. Aldehydes without -hydrogen atom in the presence of con. alkali to form each molecule of acid and alcohol by hydride active species.

46. Glyoxal reacts with con. KOH to form glycolate ion by Intra Cannizaro’s.

47. Pinacol pinacolone type reactions involve protonation, deprotonation and alkyl shift.

48. Aldol, Perkin, reformatsky and knovengel’s reactions are classified as carbanion active reaction.

49. In Beckmann’s rearrangement migration of group which is anti to-OH group takes place.

50. Beckmann’s rearrangement is a reaction of oximes in the presence of H$_2$SO$_4$ or PCl$_5$ to produce N-alkyl amide derivatives.

51. Migratory attitude of alkyl group in Pinacol-pinacolone, beckmann’s and bayer villegar oxidation is $\rightarrow$ C$_6$H$_5$ $\rightarrow$ (CH$_3$)$_3$C $\rightarrow$ (CH$_3$)$_2$CH $\rightarrow$ C$_2$H$_5$ $\rightarrow$ CH$_3$.

52. Cyclo hexanone oxime on beckmann’s reaction gives caprolectum which on reaction with $\overset{180}{\overset{\circ}{\overset{180}{\circ}}}$ to give polymer nylon-6.

53. 2-methyl propanal even contains $\equiv$hydrogen atom but does not give aldol reaction.

54. Diphenyl glyoxal reacts with con. KOH to form salt of benzillic acid.

55. Propanone in the presence of dry HCl gas by enolic intermediate to form diacetone alcohol undergo heating form mesityl oxide. If this again reacts with propanone to form phoron.

56. Benzaldehyde reacts with alc. KCN to form Benzoin which on oxidation form benzil.

57. By wittig reaction carbonyl compounds are converted into E-Z form of alkene.

58. Benzaldehyde do not respond to benedict’s and fehling’s solution due to less reducing power of aromatic aldehyde.

59. HCOOH respond to oxidising agent due to presence of $\equiv$CHO group.

60. HI $\rightarrow$ H$_2$SO$_4$ $\rightarrow$ HNO$_3$ $\rightarrow$ RCOOH $\rightarrow$ H$_2$CO$_3$ $\rightarrow$ C$_4$H$_5$OH $\rightarrow$ H$_2$O $\rightarrow$ CH $\equiv$CH $\rightarrow$ NH$_3$ $\rightarrow$ CH$_3$ $\rightarrow$ C $\equiv$CH $\rightarrow$ CH$_2$ $\rightarrow$ CH$_3$ $\rightarrow$ this is decreasing acidic nature.

61. Acids with $\equiv$hydrogen atom when reacts with halogen in the presence of P to form $\equiv$haloacid (HVZ).

62. Carboxylic acid on reaction with PCl$_5$, CH$_3$N$_2$ and H$_2$O to form higher acid or next homologue.

63. For reactivity of acid derivatives use funda weaker the base better the leaving group.

64. Acid amide on reaction with PCl$_5$ to form alkane nitrile.

65. Anhydride on reaction with carbonyl compound in the presence of base (carbanion) forms $\equiv$, $\beta$-unsaturated carboxylic acid (perkin reaction).
66. Amide on reaction with Br₂ and alkali to form primary amine of lower homologue. Intermediate species is nitrene which undergo intra rearrangement to form RNH₂.

67. Ester’s with -hydrogen atom in the presence of strong base to form carbanian undergo nucleophilic substitution reaction forms β-keto ester for example ethyl acetate in the presence of pot. Butoxide form aceto acetic ester (AAE). reaction is known as clesen’s ester condensation of four types

(a) Simple clesen’s ester condensation.
(b) CROSS clesen’s ester condensation.
(c) Intra CEC (Dieckmann’s condensation).
(d) mixed clesen ester condensation.

68. For a compound to be aromatic it must be cyclic, planar and obey (4n + 2) π electron rule.

69. Cyclooctatetraene is non aromatic compound while pyrrole, pyredene, furan, cyclopentadienylanion all are aromatic.

70. In aromatic electrophilic substitution reaction there is no hydrogen isotopic effect except sulphonation and iodination.

71. m- directing groups like nitrobenzene and benzaldehyde cannot undergo fridal craft reaction.

72. Phenol is less acidic than general carboxylic acid, cannot react with NaHCO₃.

73. Anilene is more reactive than phenol towards electrophilic substitution reaction because less energy difference between nitrogen and carbon.

74. Benzene diazonium chloride reacts with phenol or anilene to form azo compounds.

75. C₆H₅NH₂ > C₆H₅OH > C₆H₅OR > C₆H₅Br > C₆H₅NO₂ (Reactivity towards electrophilic substitution reaction

### General Series - Chemistry Fact Sheet - 3

<table>
<thead>
<tr>
<th>General series</th>
<th>Order</th>
<th>Why?</th>
</tr>
</thead>
<tbody>
<tr>
<td>B.P. of CH₃CH₂OH, CH₃COCH₃, CH₃CH₂</td>
<td>I &gt; II &gt; III</td>
<td>There is intermolecular H-bonding I. III has weak force of attraction and is most volatile. Intramolecular H-bonding in o-isomer makes it more volatile.</td>
</tr>
<tr>
<td>B.P. of o, m, p-nitro phenol</td>
<td>o &lt; m &lt; p</td>
<td></td>
</tr>
</tbody>
</table>

CBSE Standard 12 Chemistry Survival Guide - Stoichiometry Titration by Prof. Subhashish Chattopadhyay
SKMClasses Bangalore Useful for IIT-JEE, I.Sc. PU-II, Boards, IGCSE IB AP-Chemistry and other exams
3. Reactivity of ... with Tollen's reagent

\[ \text{CHO, CH}_2\text{CHO, CH}_2\text{COCH}_3\text{, C}_2\text{H}_5\text{CHO} \]

\[ \text{I, II, III, IV} \]

---CHO group is easily oxidised compared to keto group due to reducing hydrogen.

4. Reactivity of ... with Fehling's solution

\[ \text{I > II > IV > III} \]

---do---

5. Extent of hydration of

\[ \text{I < II < III < IV} \]

Aldehydes are more hydrated than ketones. Halide makes C of carbonyl group more electropositive.

6. Electrophilic nature of ........ for nucleophilic attack

\[ \text{I > II > III} \]

CH\(_3\) group decreases +ve charge on C hence nucleophilic attack.

7. Reactivity of isomeric 1º, 2º, 3º butyl halide towards elimination (E1 or E2)

\[ 3º < 2º < 1º \]

due to stability of intermediate carbocation

8. Dehydration of

\[ \text{OH} \]

\[ \text{IV < I < II < III} \]

Alcohol leading to increase in conjugation due to dehydration is more easily dehydrated. IV is vinylic, hence least.

9. Stability of
Substituted alkenes are more stable. More the alkyl groups are attached to the doubly bonded carbon atom more is the stability.

10. Stability of
   \[ \text{I} < \text{III} < \text{II} \]
   II is more substituted than III (More hyperconjugation more stability)

11. Stability of
   \( \text{I} > \text{II} > \text{IV} \)
   IV is vinylic while in conjugative, II allylic.

12. Stability of
   \[ \text{I} < \text{IV} < \text{II} < \text{III} \]
   III is 3° allylic and II is 1° allylic

13. Dehydration of
   \( 1^\circ, 2^\circ, 3^\circ \) isomeric butyl alcohol
   \[ 3^\circ < 2^\circ < 1^\circ \]
   More the stability of intermediate, greater the reactivity of chemical reaction.

14. Boiling points of
   \( n \)-butyl amine, \( n \)-butyl alcohol, \( n \)-pentane
   \[ \text{I} > \text{II} > \text{III} \]
   II have H-bonding but electronegativity of O > N hence H-bonding in II > I greater the stability, easier the formation of particular species.

15. Formation of
   \[ \text{I} > \text{II} > \text{III} > \text{IV} \]
   (easiest I)

16. Reactivity of C—H bond (abstraction of H)
17. Leaving nature (tendency) of ... in \( S_n \) reaction.
   
   I < II < III < IV < V < VI < VII < VIII

   If acid is strong, its conjugate base is weak and greater the leaving tendency.

18. Rate of esterification of the following acids with MeOH
   
   MeCH_2COOH, Me_2CHCOOH, Me_3COOH, Et_2COOH (I - Pr)_2CHCOOH

   I > II > III > IV > V

   As the size of the substituents on the \(-\text{C} \) increases, the tetrahedrally bonded interme.- diate becomes more crowded and these slower the rate.

19. Relative reactivity of ... with electrophile in \( S_E \) reaction

   \[
   \begin{align*}
   & \text{(I)} \quad \text{(II)} \quad \text{(III)} \quad \text{(IV)} \\
   & \quad \text{CH}_3 \quad \text{CH}_3 \quad \text{OH} \quad \text{CH}_3
   \end{align*}
   \]

   I > II > IV > III > V

   \(-\text{CH}_3 \) is o-, p-directing and responsible for activation.

20. Relative reactivity of these compounds with electrophile in \( S_E \) reaction

   \[
   \begin{align*}
   & \text{(I)} \quad \text{(II)} \quad \text{(III)} \quad \text{(IV)} \\
   & \quad \text{CH}_3 \quad \text{CH}_3 \quad \text{NO}_2 \quad \text{OH}
   \end{align*}
   \]

   II > I > III > IV

   \(-\text{CH}_3 \) is o-, p-directing due to activation while \(-\text{COOH} \) is m-directing and deactivating group.

21. Relative reactivity of ... with electrophile in \( S_E \) reaction

   \[
   \begin{align*}
   & \text{(I)} \quad \text{(II)} \quad \text{(III)} \quad \text{(IV)} \\
   & \quad \text{CH}_3 \quad \text{CH}_3 \quad \text{COOH} \quad \text{COOH}
   \end{align*}
   \]

   II > I > III > IV

   As the number of \( \text{sp}^3 \) hybridised C
Activating effects of the following o, p-directors.

22. 

\[ \begin{align*}
\text{II} & > \text{I} > \text{III} \\
-\text{OH}, -\text{O}^\cdot & -\text{CH}_3\text{C}=\text{O} \\
\end{align*} \]

Relative reactivity of & towards SN1 reaction
benzyl chloride, & p-methoxy benzyl chloride
and p-nitro benzyl chloride

23. 

Relative reactivity of & towards SN1 and SN2 reaction
PhCH\text{Cl} (I) & PhCH\text{ClMe} (II) & PhCClMe\text{e} (III)
(Ph stands for phenyl, C\text{6}H\text{5})

24. 

Relative reactivity of & with E\text{+} (electrophile) in SE reaction.
S\text{N1} & S\text{N2} & SE
\begin{align*}
\text{SN1} & : 1^\circ < 2^\circ < 3^\circ \\
\text{SN2} & : 3^\circ < 2^\circ < 1^\circ \\
\text{SE} & -\text{NO}_2 \text{ deactivates benzene ring for SE}
\end{align*}
26. Order of $S_N2$ reactivity of alkoxide nucleophiles

$$\text{Me}_3\text{CO}^-, \text{MeO}^-, \text{MeCH}_2\text{O}^-, \text{Me}_2\text{CHO}^-$$

$I < IV < V < III < II$

$S_N2$ reactivity is susceptible to steric hindrance by the nucleophile as well as by the size of alkyl group.

Basic power - Chemistry Fact Sheet - 2

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<tr>
<td>I &gt; II &gt; III &gt; IV</td>
<td>OCH$_3$</td>
<td>$-\text{OCH}_3$ is strong electron donating group. This is due to ortho effect, all the aniline are less basic than $p$-substituted aniline due to steric hindrance. I (hyper conjugation and induction) II (induction) IV (ortho effect), ortho effect normally decreases basic nature.</td>
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<tr>
<td>I &gt; II &gt; III &gt; IV</td>
<td>NH$_2$</td>
<td>$-\text{NH}_2$ is strong electron donating group. This is due to ortho effect, all the aniline are less basic than $p$-substituted aniline due to steric hindrance. I (hyper conjugation and induction) II (induction) IV (ortho effect), ortho effect normally decreases basic nature.</td>
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<td>I &gt; III &gt; II</td>
<td>$\text{NH}_2$</td>
<td>lone pair on N is not used in resonance of $\pi$-electrons in I. In II lone pair of the ring is itself used in delocalisation while that of outside ring in III.</td>
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<td>lone pair on N is not used in resonance of $\pi$-electrons in I. In II lone pair of the ring is itself used in delocalisation while that of outside ring in III.</td>
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</table>
4. In II there is $sp^3$ hybridised C, In I, $sp^2$. NO$_2$ is electron withdrawing.

5. lone pair on N is used in delocalisation of $\pi$-electrons in aromatic amines while cyclohexyl is electron repelling (III); in II, lone pair on N is used by two benzene ring. NO$_2$ is electron-withdrawing, thus nitro-anilines are less basic than aniline. IV is less basic than III because $-NO_2$ is closer and exerts a stronger inductive effect.

6. phenyl and $-COCH_3$ are electronwithdrawing and $-C_6H_5 < COCH_3$

7. Electron donating nature of C$_2$H$_5 > CH_3$ So more basic strength.
Acidic Powers & Their Orders - Chemistry Fact Sheet

<table>
<thead>
<tr>
<th>Acidic power</th>
<th>Order</th>
<th>Why?</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. 3- chlorobutanoic acid</td>
<td>III &gt; II &gt; I</td>
<td>Farther the (-I) group (Cl), lesser the acidic strength.</td>
</tr>
<tr>
<td>2. 3- methyl pentanoic acid</td>
<td>I &gt; II &gt; III</td>
<td>Farther the (+I) group, greater the acidic power.</td>
</tr>
<tr>
<td>3.</td>
<td>II &gt; I &gt; III</td>
<td>-CH₃ is electron donating and -NO₂ is electron attracting.</td>
</tr>
<tr>
<td>4.</td>
<td>II &gt; I &gt; III</td>
<td>-CH₃ is electron repelling; decreases acidic strength of phenol.</td>
</tr>
<tr>
<td>5.</td>
<td>III &gt; I &gt; II</td>
<td>-OCH₃ group contains +M effect and decreases acidic power.</td>
</tr>
</tbody>
</table>
6. I < II < III

- NO₂ is electron attracting; III is more resonance stabilised than I and also than II. In I, only inductive effect is operative. sp² hybridised carbon of I, II are more electronegative hence acid strength is increased.

- Benzylic (C₆H₅CH₂) is more stabilised than allylic (CH₂==CHCH₂).

7. I > II > III

- Oxalic acid, succinic acid, malonic acid, adipic acid
- I > II > III (all dibasic)

8. I > II > III > IV

- Adipic acid

9. II < III < I

- Oxygenic, mellitic, p-nitrobenzoic acid

10. III < II < I

- Hydroxy benzoic acid

-- OH shows electron withdrawing nature at o- and m- and electron repelling at p-. o- isomer due to intramolecular bonding in salicylate ion is stronger than m- isomer
11. o- m- p- methoxy benzoic acid III < II < I —do—

12. o- m- p- amino benzoic acid I < III < II —NH$_2$ is electron donating.

**Topic wise grouping of information kind of Dictionary of Inorganic Chemistry**

Given two Equations

\[
(1) \quad \text{B}_2\text{O}_3 + 6\text{NaOH} \rightarrow (\text{X}) + 3\text{H}_2\text{O}
\]

\[
(2) \quad \text{B}_2\text{O}_3 + 2\text{NaOH} \rightarrow (\text{Y}) + \text{H}_2\text{O}
\]

What are (X) and (Y)?

Ans: X is Sodium Borate and Y is Sodium meta borate

\[
\text{B}_2\text{O}_3 + 6\text{NaOH} \rightarrow 2\text{Na}_3\text{BO}_3 \quad (\text{Sodium Borate}) + 3\text{H}_2\text{O}
\]

\[
\text{B}_2\text{O}_3 + 2\text{NaOH} \rightarrow 2\text{NaBO}_2 \quad (\text{Sodium metaborate}) + \text{H}_2\text{O}
\]
Silica is soluble in which of the following acids?

1. HF  
2. HCl  
3. HBr  
4. HI

Solution:

\[
\text{SiO}_2 + 4\text{HF} \rightarrow \text{SiF}_4 + 2\text{H}_2\text{O}
\]

and again

\[
\text{SiF}_4 + 2\text{HF} \rightarrow \text{H}_2\text{SiF}_6 \quad (\text{Hydroflurosilicic acid})
\]

K$_3$Co(NO$_3$)$_6$ is Fischer's Salt

The brown ring test for NO$_3^-$ is due to formation of the complex \([\text{Fe(H}_2\text{O})_5\text{NO}]^{2+}\)

Merck's perhydrol is 30.4% H$_2$O$_2$

H$_2$SO$_5$, H$_2$S$_2$O$_8$ have peroxide linkages
<table>
<thead>
<tr>
<th>Chemical</th>
<th>Name</th>
<th>Chemical</th>
<th>Name</th>
</tr>
</thead>
<tbody>
<tr>
<td>KNO₂</td>
<td>Potassium Nitrite</td>
<td>BaSO₃</td>
<td>Barium Sulfite</td>
</tr>
<tr>
<td>Mg(NO₃)₂</td>
<td>Magnesium Nitrate</td>
<td>Na₂SO₄</td>
<td>Sodium Sulfate</td>
</tr>
<tr>
<td>LiClO₄</td>
<td>Lithium Perchlorate</td>
<td>Ca(BrO)₂</td>
<td>Calcium Hypobromite</td>
</tr>
<tr>
<td>NaClO₃</td>
<td>Sodium Chlorate</td>
<td>Al(IO₂)₃</td>
<td>Aluminum Iodite</td>
</tr>
<tr>
<td>RbClO₂</td>
<td>Rubidium Chlorite</td>
<td>KBrO₃</td>
<td>Potassium Bromate</td>
</tr>
<tr>
<td>CsClO</td>
<td>Cesium Hypochlorite</td>
<td>LiIO₄</td>
<td>Lithium Periodate</td>
</tr>
<tr>
<td>Calcium Nitrate</td>
<td>Ca(NO₃)₂</td>
<td>(NH₄)₂SO₃</td>
<td>Ammonium Sulfite</td>
</tr>
<tr>
<td>Strontium Sulfate</td>
<td>SrSO₄</td>
<td>LiNO₂</td>
<td>lithium nitrite</td>
</tr>
<tr>
<td>Potassium Hypochlorite</td>
<td>KClO</td>
<td>LiBrO₄</td>
<td>lithium perbromate</td>
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<td>Rubidium Chlorate</td>
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<td>Ca(IO₂)₂</td>
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<tr>
<td>Ammonium Chlorite</td>
<td>NH₄ClO₂</td>
<td>B(BrO₃)₃</td>
<td>Boron Bromate</td>
</tr>
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<td>Sodium Perchlorate</td>
<td>NaClO₄</td>
<td>Mg(IO)₂</td>
<td>Magnesium Hypoiodite</td>
</tr>
</tbody>
</table>
Bromine reacting with NaOH in cold and hot give different mix of products. Specify

\[ \text{Br}_2 + 2\text{NaOH (cold)} \rightarrow \text{NaBr} + \text{NaOBr} + \text{H}_2\text{O} \]

\[ 3\text{Br}_2 + 6\text{NaOH (Hot)} \rightarrow 5\text{NaBr} + \text{NaBrO}_3 + 3\text{H}_2\text{O} \]

Malachite decomposes to give A + CO\(_2\) + H\(_2\)O and compound A on reduction with Carbon gives CO + B

Identify A and B

Solution: CuCO\(_3\).Cu(OH)\(_2\) \(\rightarrow\) 2CuO (A) + CO\(_2\) + H\(_2\)O

CuO + C \(\rightarrow\) Cu (B) + CO

Some examples of Complex Anions

\[ \text{K}_2\text{PtCl}_6 \quad \leftrightarrow \quad 2\text{K}^+ + [\text{PtCl}_6]^{2-} \]

\[ \text{Na}_3\text{AlF}_6 \quad \leftrightarrow \quad 3\text{Na}^+ + \text{AlF}_6^{3-} \]

\[ \text{K}_2\text{SiF}_6 \quad \leftrightarrow \quad 2\text{K}^+ + \text{SiF}_6^{2-} \]
The ionization potential of Pb is higher than Sn due to poor shielding by 14f electrons present in Pb.

Selenous acid $\text{H}_2\text{SeO}_3$ (Oxidation Number of Se is +4)

Telluric acid $\text{H}_6\text{TeO}_6$ (Oxidation Number of Te is +6)
Electronegativity Values (in Pauling Scale)

Microcosmic Salt is Na(NH\textsubscript{4})HPO\textsubscript{4} Sodium ammonium hydrogen phosphate

Which of the following electrolyte will be most effective in coagulation of gold sol?

(1) NaNO\textsubscript{3}  (2) K\textsubscript{4}Fe(CN)\textsubscript{6}  (3) Na\textsubscript{3}PO\textsubscript{4}  (4) MgCl\textsubscript{2}

Gold sol is a negatively charged coagulation which is carried out by positively charged ion furnished by electrolyte.

Coagulating power depends on the valency of cation of electrolyte. (Schulze Hardy Law) Mg\textsuperscript{2+} ion has highest valency so MgCl\textsubscript{2} is most effective.

Organometallic compounds are Metal atoms directly linked with Carbon. So Ti(OC\textsubscript{2}H\textsubscript{5})\textsubscript{4} is not an organometallic compound.
IUPAC name of Diamminetetraaquacobalt III chloride

As per IUPAC nomenclature, the name of the complex $[\text{Co(H}_2\text{O})_4(\text{NH}_3)_2]\text{Cl}_3$ is
(a) Tetraaquadiaminecobalt(III) chloride  
(b) Tetraaquadiaminecobalt(III) chloride  
(c) Diaminetetraaquacobalt(III) chloride  
(d) Diaminetetraaquacobalt(III) chloride

**Solution:**

While naming a complex, the ligands are quoted in alphabetical order, regardless of their charges (followed by the metal). The oxidation state of the central metal is shown by a Roman numeral in brackets immediately following the metal name without any intervening blank. $\text{NH}_3$ is named as ammine and not as amine. Hence, the IUPAC name of the given compound is Diaminetetraaquacobalt(III) chloride.

Therefore, the choice (d) is correct.
Color of light absorbed by aqueous solution of CuSO₄

The colour of light absorbed by an aqueous solution of CuSO₄ is
(a) orange-red  (b) blue-green  (c) yellow  (d) violet

Solution:
An aqueous solution of CuSO₄ is blue. The colour of light absorbed by this solution is that of complimentary colour which is orange-red.
Therefore, the choice (a) is correct.
**BaCl₂ • 5 H₂O**  
Ba²⁺ is the cation Barium, Cl⁻ is the Chloride anion. There are five water molecules therefore the name is: **Barium Chloride Pentahydrate**

**Magnesium Perchlorate**  
Magnesium is the Mg²⁺ cation, and perchlorate is the ClO₄⁻ anion, therefore we need two perchlorate anions for each Mg cation therefore the formula is: **Mg(ClO₄)₂**

**(NH₄)₂SO₃**  
NH₄⁺ is the ammonium ion, and SO₃⁻² is the sulfite anion, therefore the name is: **Ammonium Sulfite**

**Calcium Nitrate**  
Calcium is the Ca²⁺ cation, and nitrate is the NO₃⁻ anion, therefore the formula is:  
**Ca(NO₃)₂**
**Group 1 Elements—Alkali Metals**

Group 1 contains lithium (Li), sodium (Na), potassium (K), rubidium (Rb), caesium (Cs) and francium (Fr).

All alkali metals contain one electron in their outermost orbitals. Due to contribution of one electron per atom towards the metallic bonding, all the elements are soft and low melting. Because of low ionization energies, the metals are strong reducing agents. The alkali metals dissolve in liquid ammonia and form a deep blue solution when dilute. The blue colour is due to the solvated electrons. The conducting ability of their ions in aqueous solution follows the order: Li$^+$ < Na$^+$ < K$^+$ < Rb$^+$ < Cs$^+$.

The alkali metals are very reactive and tarnish in air to form oxide. They burn in air, Li forms Li$_2$O, Na forms Na$_2$O$_2$ and higher alkali metals form superoxide (MO$_2$). The oxides of alkali metals are basic in nature. The peroxides are diamagnetic and are oxidizing agents. The superoxides are paramagnetic and are stronger oxidising agents than peroxides.

The melting points of alkali metal halides decrease in the order MF > MCI > MBr > MI. Lithium halides have melting points less than those of sodium halides due to the more covalent character possessed by lithium salts.

The solubilities of carbonates and bicarbonates increase on descending the group.

**Lother Meyer Plot of Atomic Volume vs Atomic mass**

In Lother Meyer plot of atomic volume versus atomic mass, the peaks are occupied by

(a) alkali metals  (b) alkaline earth metals  (c) halogens  (d) noble gases

**Ans:** (a)
KOBr Potassium hypobromite Structure

\[ \text{K}^+ \quad \text{O}^{-} \quad \text{Br} \]

**Group 2 Elements—Alkaline Earth Metals**

Group 2 contains beryllium (Be), magnesium (Mg), calcium (Ca), strontium (Sr), barium (Ba) and radium (Ra).

All alkaline earth metals contain two electrons in their outermost orbitals. These are soft. But compared to alkali metals, they are harder and have higher cohesive energy. Their melting points are higher than those of Group 1 elements and vary irregularly because of different structures. The atomic and ionic radii are smaller than those of the elements of Group 1. The alkaline earth metals dissolve in liquid ammonia giving bright blue colour due to the solvated electrons.

Alkaline earth metals are typically divalent and form colourless ionic compounds. When burnt in air, Be, Mg and Ca form monoxide while Sr, Ba and Ra form peroxides. When burnt in oxygen, they all form oxides which are basic in nature. The base strength of hydroxides increases on descending the group. Their solubility also increases on descending the group.

The solubilities of carbonates and sulphates decrease on descending the group. In deciding the solubilities of hydroxides, lattice energy predominates over hydration energy whereas in carbonates and sulphates, hydration energy predominates over lattice energy.
Temperature dependent Magnetic behaviour

NiCl₂(P(C₂H₅)₃)(C₅H₅)₂ exhibits temperature dependent magnetic behaviour (paramagnetic/diamagnetic). The coordination geometries of Ni²⁺ in the paramagnetic and diamagnetic states are respectively

(a) tetrahedral and tetrahedral  
(b) square planar and square planar  
(c) tetrahedral and square planar  
(d) square planar and tetrahedral.

Solution:

In the paramagnetic state, the complex will have unpaired 3d electrons in Ni²⁺ ion, whereas no such electrons (i.e. all the 3d electrons will be paired) are present in the diamagnetic state.

The electronic configuration of \( {\text{Ni}}^{2+} \) is

\[
\begin{align*}
\text{3d} & \quad 4s & \quad 4p \\
\uparrow & \downarrow & \uparrow
\end{align*}
\]

If the two unpaired electrons remain present in the ion, it will undergo \( \text{sp}^3 \) hybridization to accommodate ligands and the geometry of the complex ion will be tetrahedral.

If the two unpaired electrons are coupled, the ion will be diamagnetic and it can undergo \( \text{dsp}^3 \) hybridization to accommodate ligands leading to the square planar geometry to the complex ion.

Therefore, the choice (c) is correct.
Chlorine dioxide can be made by carefully adding sulfuric acid to potassium chlorate, KClO₃. It would be expected that this mixture would react to produce chloric acid, HClO₃, and then, because of the dehydrating power of sulfuric acid, to produce the anhydride of chloric acid, Cl₂O₅:

\[
\begin{align*}
KClO₃ + H₂SO₄ & \rightarrow KHSO₄ + HClO₃ \\
2HClO₃ & \rightarrow H₂O + Cl₂O₅
\end{align*}
\]

Dichlorine pentoxide, Cl₂O₅, however, is very unstable—its existence has never been verified. If it is formed at all, it decomposes at once to give chlorine dioxide and oxygen:

\[
2Cl₂O₅ \rightarrow 4ClO₂ + O₂
\]

The molecule has a triangular structure, with O—Cl—O angle 118° and bond lengths 1.49 Å. We assign it the structure \( \text{Cl}^{+} \cdot \cdot \cdot \text{O}^{-} \), with interchange of the two kinds of bonds (resonance).

Atomic radii of fluorine and neon in Angstrom units are respectively given by
(a) 0.72, 1.60  
(b) 1.60, 1.60  
(c) 0.72, 0.72  
(d) none of these values

Ans: (a)

Atomic radius of Neon will be larger than Fluorine
Reaction of white phosphorus with aqueous NaOH

The reaction of white phosphorus with aqueous NaOH gives phosphine along with another phosphorus containing compound. The reaction type, the oxidation states of phosphorus in phosphine and the other product, respectively, are:

(a) redox reaction, $-3$ and $-5$
(b) redox reaction, $3$ and $+5$
(c) disproportionation reaction, $-3$ and $+5$
(d) disproportionation reaction, $-3$ and $+3$

**Solution:**

The reaction is:

$$P_4 + 3\text{NaOH} + 3\text{H}_2\text{O} \rightarrow \text{PH}_3 + 3\text{NaH}_2\text{PO}_2$$

The salt NaH$_2$PO$_2$ undergoes the following changes on heating:

$$4\text{NaH}_2\text{PO}_2 \rightarrow \text{Na}_4\text{P}_2\text{O}_7 + 2\text{PH}_3 + \text{H}_2\text{O}$$

The reaction is disproportionation reaction and the oxidation states of phosphorus are $-3$ (in phosphine) and $+5$ in the second product.

Therefore, the choice (c) is correct.

Generally the stabilities of complexes decreases with increasing atomic number for the electropositive metals e.g. group IIA or hard Lewis acids, and increase with increasing atomic number for the noble metals (soft acids), following the general trend of ionization energy.

(a) For electropositive metals (hard acids) the order of stabilities of the Halide complexes $F > Cl > Br > I$ but for highly polarizing soft acid metal ions such as Hg$^{2+}$, we see the reverse order.

(b) The most electropositive metals (hardest acids) show a greater tendency of forming complexes, with hard ligands such as $F$ or Oxygen containing ligands.
Metals are classified according to their acceptor properties.

Class A - Hard Acids. Show affinities to ligands whose basicity is proportional to protons.

Class B - Soft acids form stable olefin complexes.

Class C - Borderline metals
Bromine forms only two stable oxygen acids—hypobromous acid and bromic acid—and their salts:

HBrO, hypobromous acid
HBrO₃, bromic acid

KBrO, potassium hypobromite
KBrO₃, potassium bromate

Their preparation and properties are similar to those of the corresponding compounds of chlorine. They are somewhat weaker oxidizing agents than their chlorine analogues.

Group 13 Elements—Boron Family

Group 13 contains boron (B), aluminium (Al), gallium (Ga), indium (In), and thallium (Tl). These elements have outer electronic configuration \((ns)^2 (np)^1\), where \(n\) varies from 2 to 6. Boron is nonmetal while others are metals. The atomic litter of Ga, In and Tl are smaller than the expected values due to d-block contraction. The atomic radius of Tl is a little larger than in due to lanthanide contraction. On descending the group, +1 oxidation state becomes more stable than +3 state due to the inert pair effect.

The very high melting point of boron is due to its icosahedral structure. In boron family, gallium has the lowest melting point. The ionization energies do not follow the expected trend of decreasing values on descending the group.

All elements burn in oxygen at high temperatures forming \(M_2O_3\). The reaction of aluminium with oxygen (known as thermite reaction) is strongly exothermic. Aluminium is amphoteric. It dissolves in dilute mineral acids and in aqueous sodium hydroxide.

The acidic character of hydroxides decreases on descending the group.

Boric acid is a very weak monobasic acid. It does not liberate hydrogen ion but accepts a hydroxyl ion. In the presence of \(\alpha\)-d-glucol (glycerol, mannitol or sugars), boric acid behaves as a strong acid and can be titrated with NaOH in the presence of phenolphthalein indicator.
**Results of the Flame Test for Various Cations**

<table>
<thead>
<tr>
<th>1</th>
<th>2</th>
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<td>Li</td>
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<td>Ne</td>
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<td>Al</td>
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<td>Cl</td>
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<td>Nd</td>
<td>Pm</td>
<td>Sm</td>
<td>Eu</td>
<td>Gd</td>
<td>Tb</td>
<td>Dy</td>
<td>Ho</td>
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<td>97</td>
<td>98</td>
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</tbody>
</table>

The fluorides of Al, Ga, In, and TI are ionic while the other halides are generally covalent and exist as dimer.

The trihalides of boron are electron-deficient compounds. Due to back bonding, the electron density on boron is increased. The tendency to form π-π bond is maximum in BF$_3$ and falls rapidly on passing to BCl$_3$ to BBr$_3$. The increasing order of acid strength follows the order BF$_3$ < BCl$_3$ < BBr$_3$. 
The halogens other than fluorine form stable compounds corresponding to nearly all values of the oxidation number from $-1$ to $+7$, as shown in the accompanying chart.

<table>
<thead>
<tr>
<th>Oxidation Number</th>
<th>Compound 1</th>
<th>Compound 2</th>
<th>Compound 3</th>
</tr>
</thead>
<tbody>
<tr>
<td>+7</td>
<td>$\text{HClO}_4$, $\text{Cl}_2\text{O}_7$</td>
<td>$\text{H}_5\text{IO}_6$</td>
<td></td>
</tr>
<tr>
<td>+6</td>
<td>$\text{Cl}_2\text{O}_6$</td>
<td></td>
<td></td>
</tr>
<tr>
<td>+5</td>
<td>$\text{HClO}_3$</td>
<td>$\text{HBrO}_3$</td>
<td>$\text{HIO}_3$, $\text{I}_2\text{O}_6$</td>
</tr>
<tr>
<td>+4</td>
<td>$\text{ClO}_2$</td>
<td>$\text{BrO}_2$</td>
<td>$\text{IO}_2$</td>
</tr>
<tr>
<td>+3</td>
<td>$\text{HClO}_2$</td>
<td></td>
<td></td>
</tr>
<tr>
<td>+2</td>
<td>$\text{HClO}$</td>
<td>$\text{HBrO}_2$, $\text{Br}_2\text{O}$</td>
<td>$\text{HIO}$</td>
</tr>
<tr>
<td>+1</td>
<td>$\text{Cl}_2$</td>
<td>$\text{Br}_2$, $\text{I}_2$</td>
<td></td>
</tr>
<tr>
<td>0</td>
<td>$\text{F}_2$</td>
<td>$\text{HCl}$, $\text{Cl}^-$</td>
<td>$\text{HBr}$, $\text{Br}^-$</td>
</tr>
<tr>
<td>-1</td>
<td>$\text{HF}$, $\text{F}^-$</td>
<td>$\text{HCl}$, $\text{Cl}^-$</td>
<td>$\text{HI}$, $\text{I}^-$</td>
</tr>
</tbody>
</table>
cyanide extraction process of silver from argentite ore

In the cyanide extraction process of silver from argentite ore, the oxidizing and reducing agents used are
(a) \(O_2\) and \(CO\), respectively
(b) \(O_2\) and \(Zn\) dust, respectively
(c) \(HNO_3\) and \(Zn\), respectively
(d) \(HNO_3\) and \(CO\), respectively

**Solution:**
The argentite ore contains \(Ag_2S\). After crushing and concentration by froth flotation process, the ore is leached with 0.4 – 7% solution of sodium cyanide kept agitated with a current of air.
Silver passes into the solution as argentocyanide. \(Ag_2S + 4NaCN \rightleftharpoons 2Na[Ag(CN)_2] + Na_2S\)
The air blown removes \(Na_2S\) as \(Na_2S_2O_3\) and \(Na_2SO_4\) causing the above reaction to proceed to completion. Silver is recovered by adding zinc into argentocyanide. \(2Ag(CN)_2^- + Zn \rightarrow [Zn(CN)_4]^{2-} + 2Ag\)
Thus, oxidizing agent is \(O_2\) (from air) and the reducing agent is \(Zn\).
Therefore, the choice (b) is correct.
Magnitude of electron affinity (kJ/mol), s-, p-, and d-block elements

<table>
<thead>
<tr>
<th>Common Name</th>
<th>Chemical Name</th>
</tr>
</thead>
<tbody>
<tr>
<td>acetone</td>
<td>dimethyl ketone; 2-propanone (usually known as acetone)</td>
</tr>
<tr>
<td>acid potassium sulfate</td>
<td>potassium bisulfate</td>
</tr>
<tr>
<td>acid of sugar</td>
<td>oxalic acid</td>
</tr>
<tr>
<td>ackey</td>
<td>nitric acid</td>
</tr>
<tr>
<td>alcali volatil</td>
<td>ammonium hydroxide</td>
</tr>
<tr>
<td>alcohol, grain</td>
<td>ethyl alcohol</td>
</tr>
<tr>
<td>alcohol sulfuris</td>
<td>carbon disulfide</td>
</tr>
<tr>
<td>Alcohol, wood</td>
<td>Methyl alcohol</td>
</tr>
<tr>
<td>-------------</td>
<td>----------------</td>
</tr>
<tr>
<td>Alum</td>
<td>Aluminum potassium sulfate</td>
</tr>
<tr>
<td>Alumina</td>
<td>Aluminum oxide</td>
</tr>
<tr>
<td>Antichlor</td>
<td>Sodium thiosulfate</td>
</tr>
<tr>
<td>Antimony black</td>
<td>Antimony trisulfide</td>
</tr>
<tr>
<td>Antimony bloom</td>
<td>Antimony trioxide</td>
</tr>
<tr>
<td>Antimony glance</td>
<td>Antimony trisulfide</td>
</tr>
<tr>
<td>Antimony red (vermillion)</td>
<td>Antimony oxysulfide</td>
</tr>
<tr>
<td>Aqua ammonia</td>
<td>Aqueous solution of ammonium hydroxide</td>
</tr>
<tr>
<td>Aqua fortis</td>
<td>Nitric acid</td>
</tr>
<tr>
<td>Aqua regia</td>
<td>Nitrohydrochloric acid</td>
</tr>
<tr>
<td>Aromatic spirit of ammonia</td>
<td>Ammonia in alcohol</td>
</tr>
<tr>
<td>Arsenic glass</td>
<td>Arsenic trioxide</td>
</tr>
<tr>
<td>Azurite</td>
<td>Mineral form of basic copper carbonate</td>
</tr>
<tr>
<td>Asbestos</td>
<td>Magnesium silicate</td>
</tr>
<tr>
<td>Aspirin</td>
<td>Acetylsalicylic acid</td>
</tr>
<tr>
<td>Baking soda</td>
<td>Sodium bicarbonate</td>
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<tr>
<td>Banana oil (artificial)</td>
<td>Isoamyl acetate</td>
</tr>
<tr>
<td>Barium white</td>
<td>Barium sulfate</td>
</tr>
<tr>
<td>Benzol</td>
<td>Benzene</td>
</tr>
<tr>
<td>Bicarbonate of soda</td>
<td>Sodium hydrogen carbonate or sodium bicarbonate</td>
</tr>
<tr>
<td>Bichloride of mercury</td>
<td>Mercuric chloride</td>
</tr>
<tr>
<td>Bichrome</td>
<td>Potassium dichromate</td>
</tr>
<tr>
<td>Bitter salt</td>
<td>Magnesium sulfate</td>
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<tr>
<td>Black ash</td>
<td>Crude form of sodium carbonate</td>
</tr>
<tr>
<td>Black copper oxide</td>
<td>Cupric oxide</td>
</tr>
<tr>
<td>Black lead</td>
<td>Graphite (carbon)</td>
</tr>
<tr>
<td>Blanc-fize</td>
<td>Barium sulfate</td>
</tr>
<tr>
<td>Bleaching powder</td>
<td>Chlorinated lime; calcium hypochlorite</td>
</tr>
<tr>
<td>Blue copperas</td>
<td>Copper sulfate (crystals)</td>
</tr>
<tr>
<td>Blue lead</td>
<td>Lead sulfate</td>
</tr>
<tr>
<td>Blue salts</td>
<td>Nickel sulfate</td>
</tr>
<tr>
<td>Blue stone</td>
<td>Copper sulfate (crystals)</td>
</tr>
</tbody>
</table>
blue vitriol  copper sulfate
bluestone   copper sulfate
bone ash    crude calcium phosphate
bone black  crude animal charcoal
boracic acid boric acid
borax       sodium borate; sodium tetraborate
bremen blue  basic copper carbonate
brimstone   sulfur
burnt alum   anhydrous potassium aluminum sulfate
burnt lime   calcium oxide
burnt ochre  ferric oxide
burnt ore    ferric oxide
brine        aqueous sodium chloride solution
butter of antimony antimony trichloride
butter of tin anhydrous stannic chloride
butter of zinc zinc chloride
calomel      mercury chloride; mercurous chloride
carbolic acid phenol
carbonic acid gas carbon dioxide
caucistic lime calcium hydroxide
caustic potash potassium hydroxide
caucistic soda sodium hydroxide
chalk        calcium carbonate
Chile saltpeter sodium nitrate
Chile nitre  sodium nitrate
Chinese red  basic lead chromate
Chinese white zinc oxide
chloride of soda sodium hypochlorite
chloride of lime calcium hypochlorite
chrome alum  chromic potassium sulfate
chrome green chromium oxide
chrome yellow lead (VI) chromate
chromic acid chromium trioxide
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<th>Equivalent Term</th>
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<td>corrosive sublimate</td>
<td>mercury (II) chloride</td>
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<tr>
<td>corundum (ruby, sapphire)</td>
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</tr>
<tr>
<td>cream of tartar</td>
<td>potassium bitartrate</td>
</tr>
<tr>
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<td>ferric oxide</td>
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<td>dechlor</td>
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<td>carbon crystal</td>
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<td>iron</td>
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<td>anhydride iron (III) chloride</td>
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<td>fluorspar</td>
<td>natural calcium fluoride</td>
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<td>fixed white</td>
<td>barium sulfate</td>
</tr>
<tr>
<td>flowers of sulfur</td>
<td>sulfur</td>
</tr>
<tr>
<td>'flowers of' any metal</td>
<td>oxide of the metal</td>
</tr>
<tr>
<td>formalin</td>
<td>aqueous formaldehyde solution</td>
</tr>
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<td>French chalk</td>
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<td>French vergidris</td>
<td>basic copper acetate</td>
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<td>Term</td>
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<td>jeweler's rouge</td>
<td>ferric oxide</td>
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<td>killed spirits</td>
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<td>magnesium oxide</td>
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<td>manganese dioxide</td>
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<td>chloride of the metal</td>
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<td>potassium nitrate</td>
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<td>fuming sulfuric acid</td>
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<td>phosphoric acid</td>
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<td>ferric ferrocyanide</td>
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<td>Formula</td>
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<td>--------------------------</td>
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<td>copper acetoarsenite</td>
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<td>barium sulfate</td>
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<td>graphite</td>
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<td>mercury</td>
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<td>lead tetraoxide</td>
</tr>
<tr>
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<td>aluminum acetate solution</td>
</tr>
<tr>
<td>red prussiate of potash</td>
<td>potassium ferrocyanide</td>
</tr>
<tr>
<td>red prussiate of soda</td>
<td>sodium ferrocyanide</td>
</tr>
<tr>
<td>Rochelle salt</td>
<td>potassium sodium tartrate</td>
</tr>
<tr>
<td>rock salt</td>
<td>sodium chloride</td>
</tr>
<tr>
<td>rouge, jewelers</td>
<td>ferric oxide</td>
</tr>
<tr>
<td>rubbing alcohol</td>
<td>isopropyl alcohol</td>
</tr>
<tr>
<td>sal ammoniac</td>
<td>ammonium chloride</td>
</tr>
<tr>
<td>sal soda</td>
<td>sodium carbonate</td>
</tr>
<tr>
<td>salt, table</td>
<td>sodium chloride</td>
</tr>
<tr>
<td>salt of lemon</td>
<td>potassium binoxalate</td>
</tr>
<tr>
<td>salt of tartar</td>
<td>potassium carbonate</td>
</tr>
<tr>
<td>saltpeter</td>
<td>potassium nitrate</td>
</tr>
<tr>
<td>silica</td>
<td>silicon dioxide</td>
</tr>
<tr>
<td>slaked lime</td>
<td>calcium hydroxide</td>
</tr>
<tr>
<td>soda ash</td>
<td>sodium carbonate</td>
</tr>
<tr>
<td>soda nitre</td>
<td>sodium nitrate</td>
</tr>
<tr>
<td>soda lye</td>
<td>sodium hydroxide</td>
</tr>
</tbody>
</table>
soluble glass  sodium silicate
sour water     dilute sulfuric acid
spirit of hartshorn  ammonium hydroxide solution
spirit of salt  hydrochloric acid
spirit of wine  ethyl alcohol
spirits of nitrous ether  ethyl nitrate
sugar, table  sucrose
sugar of lead  lead acetate
sulfuric ether  ethyl ether
talc or talcum  magnesium silicate
tin crystals  stannous chloride
trona  natural sodium carbonate
unslaked lime  calcium oxide
Venetian red  ferric oxide
verdigris  basic copper acetate
Vienna lime  calcium carbonate
vinegar  impure dilute acetic acid
vitamin C  ascorbic acid
vitriol  sulfuric acid
washing soda  sodium carbonate
water glass  sodium silicate
white caustic  sodium hydroxide
white lead  basic lead carbonate
white vitriol  zinc sulfate crystals
yellow prussiate of potash  potassium ferrocyanide
yellow prussiate of soda  sodium ferrocyanide
zinc vitriol  zinc sulfate
zinc white  zinc oxide

Westron - A refrigerant. \( \text{C}_2\text{H}_2\text{Cl}_4 \)

Colours of Compounds

\( \text{Cr}_2(\text{SO}_4)_3 \) - green
PbS - Black

[Fe(H2O)5NO]SO4 - brown

(CH3COO)3Fe - blood red
NH₄Cl - white fumes

AgCl - curdy white precipitate

AgCl + NH₄OH → Ag(NH₃)₂Cl + H₂O

CrO₃ + HCl → CrO₂Cl₂ (deep red fumes) + H₂O
NaOH + CrO2Cl2 -> Na2CrO4 (yellow) + NaCl + H2O

Na2CrO4 (yellow)

Na2CrO4 + Pb(CH3COO)2 -> PbCrO4 (yellow) + CH3COONa

HgO4 (yellow)

NaBr + AgNO3 -> AgBr (pale yellow) + NaNO3

AgI, AgBr, AgCl
Ammonia test for silver halides. Test tubes and the results of using concentrated ammonia to test for the presence of three silver halides. Each one of the silver halides is next to a fuller test tube that shows the result of adding concentrated ammonia. Halides are compounds containing halogens such as here, from left to right silver chloride, silver bromide and silver iodide. Adding dilute ammonia dissolves the chloride, while concentrated ammonia is needed to dissolve the bromide, but it fails to dissolve the iodide. The silver halides are themselves the results of standard tests using silver nitrate to test for the halide ions.

BrCl - pale yellow

NaI + AgNO3 -> Agl (yellow ppt) + NaNO3
Cu + HNO₃ → Cu(NO₃)₂ + NO₂ + H₂O

Cu(NO₃)₂

NO₂

H₃BO₃ + C₂H₅OH → (C₂H₅)₃BO₂ + H₂O (greenish flame)

H₃PO₄ + 12(NH₄)₂MoO₄ + HNO₃ → (NH₄)₃PO₄.12MoO₃ (canary yellow) + 21 NH₄NO₃ + 12 H₂O

In the presence of Arsenic, we get yellow precipitate of (NH₄)₃AsO₄.12MoO₃ (ammonium arsenomolybdate)
Mo$_3$O$_8$.xH$_2$O - molybdenum blue

SiF$_4$ + H$_2$O $\rightarrow$ H$_4$SiO$_4$ (silicic acid - white) + H$_2$SiO$_4$

Na$_2$S + Na$_2$[Fe(CN)$_5$NO] $\rightarrow$ Na$_4$[Fe(CN)$_5$NOS] - violet colour
Na₂S + CdCO₃ → CdS (yellow) + Na₂CO₃

Na₃[Co(NO₂)₆] + KCl → K₃[Co(NO₂)₆] (potassium cobalt nitrate – yellow ppt) + NaCl

AgCl, PbCl₂, Hg₂Cl₂ - white
\textbf{PbCl}_2 \text{ is white}

\textbf{Hg}_2\text{Cl}_2 \text{ is white}

\textbf{H}_2\text{S, PbS, Bi}_2\text{S}_3, \text{ CuS - black}

\textbf{H}_2\text{S is Black}
PbS is Black

Bi2S3 is Black

CuS - black

CdS, As2S3, SnS2 - Yellow

CdS is Yellow
As$_2$S$_3$ is Yellow

SnS$_2$ is Yellow

Sb$_2$S$_3$ is orange
SnS is Brown

Cr(OH)₃ is green

Chromium (III) Oxide is Green

CoS, NiS - black

CoS is black
NiS is Black Mineral Millerite

Cobalt Oxide is Black

Cobalt Sulphate is deep red brown
Cobalt Nitrate is deep red brown

Cobalt Chloride is deep Brown red

Cobalt Acetate is deep Brown

Azulene - Deep Bue

Iodoform - CHI$_3$ Yellow

Iron Phenol complex - Violet - [ Fe ( OC$_6$H$_5$)$_6$ ]$^3^-$

6 C$_6$H$_5$OH + FeCl$_3$ $\rightarrow$ [ Fe ( OC$_6$H$_5$)$_6$ ]$^3^-$ + 3 H$^+$ + Cl$^-$

Ammonium Phosphomolybdate - Yellow - (NH$_4$)$_3$PO$_4$.12MoO$_3$ ( Yellow )

Prusian Blue - Fe$_4$ [Fe(CN)$_6$]$_3$.xH$_2$O

[Fe(CN)$_3$NOS ]$^4^-$ - Violet

[Fe(SCN)]$^{2^+}$ Blood Red

[Co(NH$_3$)$_5$ Br]$^{2^+}$ SO$_4^{2-}$ - Violet - Pentaaminebromocobalt(III)sulphate
Test for Halogen:

Halogens present in an organic compound form sodium halide on fusion with sodium metal. Sodium halide extracted with water can be easily identified by adding silver nitrate solution after acidifying with dil. HNO₃.

If chlorine is present, a white curdy precipitate soluble in ammonium hydroxide solution is formed.

\[
\text{Na} + \text{Cl} \rightarrow \text{NaCl}
\]

\[
\text{NaCl} + \text{AgNO}_3 \rightarrow \text{AgCl} + \text{NaNO}_3
\]

If bromine is present, a yellowish white precipitate sparingly soluble in ammonium hydroxide solution is formed.

\[
\text{Na} + \text{Br} \rightarrow \text{NaBr}
\]

\[
\text{NaBr} + \text{AgNO}_3 \rightarrow \text{AgBr} + \text{NaNO}_3
\]

If iodine is present, a yellow precipitate insoluble in ammonium hydroxide solution is formed.

\[
\text{Na} + \text{I} \rightarrow \text{NaI}
\]

\[
\text{NaI} + \text{AgNO}_3 \rightarrow \text{AgI} + \text{NaNO}_3
\]

Test for Nitrogen:

The carbon and nitrogen present in the organic compound on fusion with sodium metal gives sodium cyanide (NaCN) soluble in water. This is converted into sodium ferrocyanide by the addition of sufficient quantity of ferrous sulphate. Ferric ions generated during the process react with ferrocyanide to form prussian blue precipitate of ferric ferrocyanide.

\[
\text{Na} + \text{C} + \text{N} \rightarrow \text{NaCN}
\]

\[
6\text{NaCN} + \text{FeSO}_4 \rightarrow \text{Na}_4[\text{Fe(CN)}_6] + \text{Na}_2\text{SO}_4
\]

Sodium ferrocyanide

\[
\text{Na}_4[\text{Fe(CN)}_6] + \text{Fe}^{3+} \rightarrow \text{Fe}_4[\text{Fe(CN)}_6]_3
\]

Ferric ferrocyanide
**Test for Sulphur:**

If sulphur is present in the organic compound, sodium fusion will convert it into sodium sulphide. Sulphide ions are readily identified using sodium nitroprusside.

\[
Na + S \rightarrow Na_2S
\]

\[
Na_2S + Na_2[Fe(CN)5NO] \rightarrow Na_4[Fe(CN)5NOS]
\]

Sodium nitroprusside violet colour

**Test for both Nitrogen and Sulphur:**

If both nitrogen and sulphur are present in an organic compound, sodium fusion will convert it into sodium thiocyanate which then react with Fe\(^{3+}\) to form blood colour complex [Fe(SCN)]\(^{2+}\)

\[
Na + C + N + S \rightarrow NaSCN
\]

\[
Fe^{3+} + 2Na \rightarrow [Fe(SCN)]^{2+}
\]

*Note:* If the Lassaigne's extract containing excess of sodium metal, sodium cyanide and sulphides are formed instead of sodium thiocyanate.

\[
NaSCN + 2Na \rightarrow NaCN + Na_2S
\]

Here in this type of cases, both sulphur and nitrogen are to be identified in separate tests.
BaCO₃, SrCO₃, CaCO₃, Mg(NH₄)PO₄ - white

BaCO₃ is white

SrCO₃ is white

CaCO₃ powder is white
Mg(NH₄)PO₄ is white

Pentaamminechlorocobalt(III)Bromide

\[
\text{[ Co(NH}_3\text{)}\text{5Cl}_2 \text{Cl}_2 \text{ ] }\text{ Pentaamminechlorocobalt(III)chloride}
\]

\[
\text{Pentaamminesulphatocobalt(III)Bromide}
\]

- Dark Yellow
- Pale Yellow

CBSE Standard 12 Chemistry Survival Guide - Stoichiometry Titration by Prof. Subhashish Chattopadhyay
SKMClasses Bangalore Useful for IIT-JEE, I.Sc. PU-II, Boards, IGCSE IB AP-Chemistry and other exams
Eta bonds - Also known as Hapticity - Hapticity is the coordination of a ligand to a metal center via an uninterrupted and contiguous series of atoms.

The hapticity of a ligand is described with the Greek letter η ('eta'). For example, η2 describes a ligand that coordinates through 2 contiguous atoms. In general the η-notation only applies when multiple atoms are coordinated (otherwise the κ-notation is used). In addition, if the ligand coordinates through multiple atoms that are not contiguous then this is considered denticity (not hapticity), and the κ-notation is used once again. Lastly, bridging ligands are described with the μ ('mu') notation.

Ferrocene - bis(η5-cyclopentadienyl)iron

Uranocene - bis(η8-1,3,5,7-cyclooctatetraene)uranium
W(CO)₃(PPri₃)₂(η₂-H₂) - the first compound to be synthesized with a dihydrogen ligand (also known as Dihydrogen Complexes)

IrCl(CO)[P(C₆H₅)₃]₂(η₂-O₂) - the dioxygen derivative which forms reversibly upon oxygenation of Vaska’s complex.

- Chromium (III) chloride is purple

HgCl₂ + NH₄OH → (Hg + Hg(NH₂)Cl) + NH₄Cl + H₂O

black ppt

BiCl₃ + H₂O → BiOCl (white) + HCl

Bi(OH)₃ Bismuth Hydroxide is white
Oxidation of Phenol

Test of Phenol
With Neutral FeCl₃, Phenol on reaction with neutral ferric chloride gives violet colour of ferric phenoxide

\[ C_6H_5OH + FeCl_3 \rightarrow (C_6H_5O)_2Fe \quad \text{Violet colour} \]

\[ Cu(NO_3)_2 + K_4[Fe(CN)_6] \rightarrow Cu_2[Fe(CN)_6] \quad \text{(Chocolate brown)} + KNO_3 \]

\[ Cd(NH_3)_4(NO_3)_2 + H_2S \rightarrow CdS \quad \text{(yellow)} + NH_4NO_3 + NH_3 \]
HgCl₂ + SnCl₂ → Hg₂Cl₂ (white) + SnCl₄

SbCl₃ + H₂O → SbOCl (white) + HCl

SbOCl Antimony oxychloride
Antimony Trioxide is white

Antimony Pentoxide is gray

Antimony Trichloride is white
Potassium antimonate white

FeCl₃ + KSCN → Fe(CNS)₃ (blood red) + KCN

**Can Test**

Treat a few drops of ceric ammonium nitrate with 3-4 ml of hot aqueous solution of the organic compound. A green or brown precipitate is formed.

\[
\text{(NH}_4\text{)}_2\text{Ce(NO}_3\text{)}_6 + 2\text{C}_6\text{H}_5\text{OH} \rightarrow [\text{Ce(NO}_3\text{)}_4 (\text{C}_6\text{H}_5\text{OH})_2] + 2\text{NH}_4\text{NO}_3
\]

Phenol gives blue colour with ammonia and sodium hypo chloride.

ZnCl₂ + 2NaOH → Zn(OH)₂ (white ppt) + 2NaCl
\[
\text{Zn(OH)}_2 + \text{NaOH} \rightarrow \text{Na}_2\text{ZnO}_2 \text{ (soluble)} + \text{H}_2\text{O}
\]

\[
\text{Na}_2\text{ZnO}_2 + \text{H}_2\text{S} \rightarrow \text{ZnS (white)} + \text{NaOH}
\]

\[
\text{Mn(NO}_3\text{)}_2 + \text{Pb}_3\text{O}_4 + \text{HNO}_3 \rightarrow \text{HMnO}_4 \text{ (pink)} + \text{Pb(NO}_3\text{)}_2 + \text{H}_2\text{O}
\]
Co(NO₂)₃ + KNO₂ → K₃[Co(NO₂)₆] (yellow)

Na₄Co(CO₃)₂ + H₂O + [O] → Na₃Co(CO₃)₂ (green coloration) + NaOH
NiCO₃ + [O] → Ni₂O₃ (black) + CO₂

MgCl₂ + Na₂HPO₄ + NH₄OH → Mg(NH₄)PO₄ (white) + NaCl + H₂O

magnesium ammonium phosphate
K₂HgI₄ + NH₄Cl + KOH → NH₂HgO/HgI (Iodide of millon’s base, brown ppt) + KCl + H₂O

Chromium oxide powder is green

KI Potassium iodide is white
Aluminum Oxide is white

Mercuric iodide $\text{HgI}_2$: It is a yellow solid below 400K but changes to red solid above 400K.

$\text{HgI}_2 \xrightarrow{400K} \text{HgI}_2$

It dissolves in excess of KI forming $\text{K}_2\text{HgI}_4$;

$\text{HgI}_2 + 2\text{KI} \rightarrow \text{K}_2\text{HgI}_4$

Alkaline solution of $\text{K}_2\text{HgI}_4$ is called Nessler’s reagent.
Add NH₄OH excess and dimethylglyoxime to NiCl₂ then a rosy red ppt of nickel appears

CuO + B₂O₃ (Glassy bead) → Cu(BO₂)₂ (Copper metabolite - blue)

Compounds of zinc

Zinc oxide ZnO: Zincite (ZnO) is also called Philosopher's wool. It is a white powder, becomes yellow on heating and again white on cooling. It is amphoteric in nature. It is used as a white pigment under the name Zinc white or Chinese white.

Zinc Sulphate (white vitriol), ZnSO₄ ⋅ 7H₂O: It is a colourless transparent crystal highly soluble in water. It is used as an eye-lotion and for preparing double salts. On heating it loses its molecules of water as,

\[ \text{ZnSO}_4\cdot 7\text{H}_2\text{O} \xrightarrow{375K} \text{ZnSO}_4\cdot \text{H}_2\text{O} \xrightarrow{725K} \text{ZnSO}_4 \xrightarrow{1075K} \text{ZnO} + \text{SO}_2 + \text{O}_2 \]
Verdigris - basic copper acetate

This pigment was used to the Middle Ages in paints. Until the nineteenth century this was the most vibrant shade of green available and it was used by artists widely. The color comes from copper acetate and you should know that it is highly toxic to humans.

Compounds of Mercury

(1) Mercuric oxide, HgO : It is obtained as a red solid by heating mercury in air or oxygen for a long time at 673 K

\[ 2Hg + O_2 \rightarrow 2HgO \text{ (red)} \text{ or by heating mercuric nitrate alone or in the presence of Hg} \]

Heat

\[ 2Hg \text{ (NO}_3\text{)}_2 \rightarrow 2HgO + 4NO_2 + O_2 \text{ red} \]
When NaOH is added to a solution of HgCl$_2$, yellow precipitate of HgO are obtained.

$$\text{Hg}_2\text{Cl}_2 + 2\text{NaOH} \rightarrow \text{HgO} \downarrow + \text{H}_2\text{O} + 2\text{NaCl}$$

(yellow)

Red and yellow forms of HgO differ only in their particle size. On heating to 673 K, yellow form changes to red form.

$$\text{HgO} \xrightarrow{673K} \text{HgO}$$

Yellow → red

It is used in oil paints or as a mild antiseptic in ointments.

(2) Mercuric chloride, HgCl$_2$: It is obtained by treating Hg with Cl$_2$ or by heating a mixture of NaCl and HgSO$_4$ in presence of small amount of MnO$_2$ (which oxidizes any Hg(I) salts formed during the reaction).

Heat

$$\text{HgSO}_4 + 2\text{NaCl} \rightarrow \text{HgCl}_2 + \text{Na}_2\text{SO}_4$$
HgCl₂ is a white crystalline solid and is commonly known as corrosive sublimate. It is a covalent compound since it dissolves in organic solvents like ethanol and ether.

It is extremely poisonous and causes death. Its best antidote is white of an egg.

When treated with stannous chloride, it is first reduced to white ppt. of mercurous chloride and then to mercury (black).

$$2\text{HgCl}_2 + \text{SnCl}_2 \rightarrow \text{Hg}_2\text{Cl}_2 + \text{SnCl}_4$$

white ppt.

$$\text{Hg}_2\text{Cl}_2 + \text{SnCl}_2 \rightarrow 2\text{Hg} + \text{SnCl}_4$$

grey

With ammonia it gives a white ppt. known as infusible white ppt.

$$\text{HgCl}_2 + 2\text{NH}_3 \rightarrow \text{Hg} (\text{NH}_2) \text{Cl} + \text{NH}_4 \text{Cl}$$

A dilute solution of HgCl₂ is used as an antiseptic.

(3) Mercuric iodide, HgI₂: It is obtained when a required amount of KI solution is added to a solution of . HgCl₂. HgCl₂ + 2KI → HgI₂ + 2KCl

Below 400 K, HgI₂ is red but above 400 K, it turns yellow.

$$\text{HgI}_2 \text{ readily dissolves in excess of KI solution to form the } \text{(HgI}_4 \text{)}^2-$$

complex ion. HgI₂ + 2KI → K₂HgI₄ Red ppt. soluble colourless solution

An alkaline solution of K₂ [HgI₄] is called Nessler’s reagent and is used to test NH₄⁺ ions.

It gives a brown ppt. of NH₂ - Hg - O Hg - I (iodide of Millon’s base) with NH₄⁺ ions.

$$2\text{K}_2 [\text{HgI}_4 ] + \text{NH}_3 + 3\text{KOH} \rightarrow \text{NH}_2. \text{HgO}. \text{Hgl} + 7\text{KI} + 2\text{H}_2 \text{O}$$

It is used in ointments for treating skin infections.

(4) Mercurous chloride, Hg₂Cl₂: It is obtained as under:

(a) Hg₂ (NO₃) + 2NaCl → Hg₂Cl₂ + 2NaNO₃

white ppt.

Heat in an iron retort
(b) $\text{HgCl}_2 + \text{Hg} \rightarrow \text{Hg}_2 \text{Cl}_2$ (condenses on cooling)

It is purified by sublimation.

Mercurous chloride is also called calomel. It is a white powder insoluble in $\text{H}_2\text{O}$. On heating, it decomposes to give $\text{HgCl}_2$ and $\text{Hg}$.

\[
\text{Heat} \quad \text{Hg}_2 \text{Cl}_2 \rightarrow \text{HgCl}_2 + \text{Hg}
\]

It dissolves in chlorine water forming mercuric chloride.

$\text{Hg}_2\text{Cl}_2 + \text{Cl}_2 \rightarrow 2\text{HgCl}_2$

With ammonia, it turns black due to the formation of a mixture of finely divided black $\text{Hg}$ and mercuric amino chloride.

$\text{Hg}_2\text{Cl}_2 + 2\text{NH}_3 \rightarrow \text{Hg} + \text{NH}_4\text{HgCl} + \text{NH}_4\text{Cl}$

(5) Mercuric sulphide, $\text{HgS}$: The solubility product of $\text{HgS}$ is lower than that of $\text{ZnS}$ and hence it gets precipitated as black solid when $\text{H}_2\text{S}$ is passed through an acidic solution of any mercury (II) salt.

$\text{HgCl}_2 + \text{H}_2\text{S} \rightarrow \text{HgS} + 2\text{HCl}$

It is insoluble in water and $\text{HCl}$ but dissolves in aqua regia (1 part conc.$\text{HNO}_3$ + 3 parts conc. $\text{HCl}$)

\[
3\text{HCl} + \text{HNO}_3 \rightarrow \text{NOCI} + 2\text{H}_2\text{O} + 2 \text{[ Cl ]}
\]

\begin{tabular}{|c|c|}
\hline
\text{Aqua regia} & \text{Nitrosyl chloride} \\
\hline
\text{HgS} + 2\text{[ Cl ]} & \rightarrow \text{HgCl}_2 + \text{S} \\
\hline
\end{tabular}

(Soluble)

On sublimation, its colour changes to red and hence it is used as a red pigment.

(6) Mercuric sulphate, $\text{HgSO}_4$: It is obtained when $\text{HgS}$ is treated with conc.$\text{H}_2\text{SO}_4$.

$\text{Hg} + 2\text{H}_2\text{SO}_4 \rightarrow \text{HgSO}_4 + \text{SO}_2 + 2\text{H}_2\text{O}$

It is a white solid which decomposes on heating to give mercurous sulphate.

\[
3\text{HgSO}_4 \xrightarrow{\Delta} \text{Hg}_2\text{SO}_4 + \text{Hg} + 2\text{SO}_2 + 2\text{O}_2
\]
It is used as a catalyst in the hydration of alkynes to give aldehydes or ketones. It is also used as a cosmetic under the name Vermillion and in ayurvedic medicine as makardhwaj.

(7) Amalgams: Mercury forms alloys commonly known as amalgams, with all metals except iron and platinum. Hence it is transported in iron containers.

(8) Alloy of transition metal: See in table discuss earlier in metallurgy.

The common Anions are divided into 3 groups for the purpose of identification:

Those which evolve gases with dilute hydrochloric acid:
- Carbonate ($CO_3^{2-}$)
- bicarbonate(HCO$_3^-$)
- Nitrite(NO$_2^-$)
- Sulphide ($S^{2-}$)
- Sulphite(SO$_3^{2-}$)
- Thiosulphate(S$_2$O$_3^{2-}$)

Those which do not react with dilute HCl, but which do evolve gases or volatile liquids with concentrated sulphuric acid:
- Chloride (Cl$^-$)
- Bromide (Br$^-$)
- Iodide (I$^-$)
- Nitrate (NO$_3^-$)

Those which do not react with either dilute hydrochloric acid or concentrated sulphuric acid:
- Phosphate (PO$_4^{3-}$)
- Borate (B$_4$O$_7^{2-}$)
- Sulphate (SO$_4^{2-}$)

A Radical is an atom, or group of atoms that behaves like a single atom. Examples: H, K, Na, Cu, Pb, simple radicals; NH$_4$, SO$_2$, NO$_2$, compound radicals.
A Basic Radical is a metal, or any radical behaving like a metal. Examples: Ag, Cu, Ca, Na, and NH 4 the last so classified because it behaves like such simple radicals as Na, K, and Li.

An Acid Radical is a nonmetal, or any radical behaving like a nonmetal. Examples: Cl, Br, I, SO 2 , NO2

An Acid is a substance containing H that may be replaced by a basic radical. When a nonmetal is combined with H only, as in HC1, HBr, H 2 S, the compound is said to be a binary (i.e. composed of two elements) acid, or an hydracid. When the nonmetal is combined with H and O, the compound is called a ternary (i.e. composed of three or more elements) acid, or an oxyacid; and the characteristic name of the acid is formed by adding the termination -ic to the name of the nonmetal to denote more oxygen, or the termination -ous to denote less. Examples: H2SO4 sulphuric, and H2SO3 sulphurous acid.

These are classified in six different groups in accordance with certain properties which permit a group precipitation by a single reagent. A reagent used for precipitation of a whole group is called a general reagent.

A reagent used as a test for a single substance is known as a special reagent.

Experiments showing Method of Classification

Use the following solutions: AgNO3, Cu(NO3)2, AsCl3, Fe2Cl6, Co(NO3)2, Ba(NO3)2, NaCl.

Take about 2 c.c. of each solution, and add HC1.*

Points to be Observed Whether a precipitate is formed.

Whether the precipitate is soluble in an excess of the reagent.

* Always use dilute acids unless otherwise directed.

GROUPING THE BASES

The color of each precipitate.

Whether there is change in color of the precipitate.

Change in color of solutions.

Chemical equations.

Tabulate all results carefully in your notebook, according to the following scheme:

HC1 was added to the seven solutions with the following results:

Typical of Gr. I AgNO 3 + HC1 = AgCl w. pp. + I IN ( ) 8 .
Typical of Gr. II A Cu(NO3)2 + II Cl = No change.
Typical of Gr. II B AsCl 3 + HC1 = No change.
Typical of Gr. III Fe 2 Cl 6 + HC1 = No change.
Typical of Gr. IV Co(NO 3 ) 2 + IIICl = No change.
Typical of Gr. V $\text{Ba(NO}_3\text{)}_2 + \text{HCl} = \text{No change.}$
Typical of Gr. VI $\text{NaCl} + \text{HCl} = \text{No change.}$

### 1) Anions which react with dilute hydrochloric acid

<table>
<thead>
<tr>
<th>Carbonates ($\text{CO}_3^{2-}$)</th>
<th>Bicarbonates ($\text{HCO}_3^-$)</th>
<th>Nitrites ($\text{NO}_2^-$)</th>
</tr>
</thead>
<tbody>
<tr>
<td>All carbonates except those of alkali metals, and ammonium are very slightly or difficulty soluble in water. Accordingly reactions in solution are only carried out in case of the soluble salts.</td>
<td>All bicarbonates are water soluble.</td>
<td>Most nitrites are water soluble except that of silver.</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Solid salt + dil. HCl</th>
<th>Effervescence and a colourless gas is evolved. Carbon dioxide ($\text{CO}_2$).</th>
<th>Effervescence and a colourless gas is evolved. Carbon dioxide ($\text{CO}_2$).</th>
<th>Pungent brown fumes are evolved consisting mainly of nitrogen dioxide ($\text{NO}_2$), which is produced by the combination of nitric oxide ($\text{NO}$) with the oxygen of the air:</th>
</tr>
</thead>
<tbody>
<tr>
<td>$\text{Na}_2\text{CO}_3 + 2\text{HCl} \rightarrow 2\text{NaCl} + \text{H}_2\text{O} + \text{CO}_2$</td>
<td>$\text{NaHCO}_3 + \text{HCl} \rightarrow \text{NaCl} + \text{H}_2\text{O} + \text{CO}_2$</td>
<td>$\text{NaNO}_2 + \text{HCl} \rightarrow \text{NaCl} + \text{H}_2\text{O} + \text{NO}_2$</td>
<td></td>
</tr>
<tr>
<td>$\text{Na}_2\text{HCO}_3 + \text{HCl} \rightarrow \text{NaCl} + \text{H}_2\text{O} + \text{CO}_2$</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

### To differentiate between $\text{CO}_3^{2-}$ and $\text{HCO}_3^-$:

1) Salt solution + magnesium sulphate solution:
- White precipitate of magnesium carbonate:
  - $\text{Na}_2\text{CO}_3 + \text{MgSO}_4 \rightarrow \text{MgCO}_3 \downarrow + \text{Na}_2\text{SO}_4$

2) Salt solution + mercuric chloride solution:
- red-brown precipitate of mercuric carbonate:
  - $\text{Na}_2\text{CO}_3 + \text{HgCl}_2 \rightarrow \text{HgCO}_3 \downarrow + 2\text{NaCl}$

### Confirmatory tests of nitrites:

1) The brown ring test:
- To a dilute solution of the nitrite, add two drops of freshly prepared ferrous sulphate solution ($\text{FeSO}_4$), and then add dil. $\text{H}_2\text{SO}_4$ very carefully drop by drop down the inside of the test tube: a brown ring forms due to the formation of (Fe(NO)SO). Note: If the addition of dil. $\text{H}_2\text{SO}_4$ is not down the inside of the test tube, the solution becomes brown color (we can not note the ring).

2) Solution of nitrite + potassium iodide solution (KI) + dilute sulphuric acid: a pale brown color appears due to the liberation of iodine.
   - (The nitrite solution is an oxidizing agent.)
   - $2\text{NO}_2^- \rightarrow \text{2NO} + \text{2I}^- \rightarrow 2\text{I}_2 + \text{2H}_2\text{O}$

3) Solution of nitrite + acidified solution of potassium permanganate $\text{KMnO}_4$ (i.e., $\text{KMnO}_4 + \text{H}_2\text{SO}_4$): the purple color of the permanganate disappears (the nitrite solution is a reducing agent).
   - $5\text{KNO}_2 + 3\text{KMnO}_4 + 3\text{H}_2\text{SO}_4 \rightarrow 5\text{KNO}_3 + 3\text{K}_2\text{SO}_4 + 3\text{H}_2\text{O}$

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CBSE Standard 12 Chemistry Survival Guide - Stoichiometry Titration by Prof. Subhashish Chattopadhyay
SKMClasses Bangalore Useful for IIT-JEE, I.Sc. PU-II, Boards, IGCSE IB AP-Chemistry and other exams
### Is HgCO₃ reddish Brown?

<table>
<thead>
<tr>
<th>Sulphites (SO₃²⁻)</th>
<th>Thiosulphates (S₈O₂⁴²⁻)</th>
<th>Sulphides (S²⁻)</th>
</tr>
</thead>
<tbody>
<tr>
<td>All sulphites are slightly soluble in water except the alkali metal sulphites and the ammonium salt.</td>
<td>Sodium thiosulphate is readily soluble in water, other thiosulphates are slightly soluble.</td>
<td>All sulphides, except those of alkali, alkali earth metals and the ammonium ion are slightly soluble in water.</td>
</tr>
</tbody>
</table>

**Solid salt + dilute HCl**

Sulphur dioxide gas (SO₂), is evolved, which is recognizable by its pungent smell.

N₂SO₃ + 2HCl → 2NaCl + H₂O + SO₂

Test for sulphur dioxide:

Dip a small piece of paper into an acid solution of potassium dichromate, and hold it over the mouth of the test tube. The paper will turn green, because the sulphur dioxide reduces the dichromate to a chromic sulphate.

3SO₂ + K₂Cr₂O₇ + H₂SO₄ →
K₂SO₄ + Cr₂(SO₄)₃ + H₂O

Orange → Green

\[ \text{color} \rightarrow \text{color} \]

Colourless gas, hydrogen sulphide (H₂S) is evolved, which is recognizable by its bad smell.

Na₂S + 2HCl → 2NaCl + H₂S

Test for hydrogen sulphide:

Dip a piece of paper in a solution of lead nitrate, and hold it over the mouth of the test tube in which H₂S is being generated.

The paper will turn black due to the formation of lead sulphide.

H₂S + 2Pb(CH₃COO)₂ → PbS + 2CH₃COOH

Black

### K₂SO₄ is white

### Cr₂(SO₄)₃ is green
### Sulphites

<table>
<thead>
<tr>
<th>Confirmation test:</th>
</tr>
</thead>
<tbody>
<tr>
<td>1) Salt solution + silver nitrate solution (AgNO₃)</td>
</tr>
<tr>
<td>A white precipitate forms (silver sulphate Ag₂SO₄), which dissolves in excess of sulphite due to the formation of soluble complex.</td>
</tr>
<tr>
<td>Na₂SO₃ + 2AgNO₃ → Ag₂SO₄ + 2NaNO₃ (Soluble complex)</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Confirmatory test:</th>
</tr>
</thead>
<tbody>
<tr>
<td>2) Salt solution + lead acetate solution Pb(Cl₃C₂COO)₂</td>
</tr>
<tr>
<td>A white ppt of lead sulphide ( Pb₃S₄ ) is formed.</td>
</tr>
<tr>
<td>Na₂SO₃ + Pb(Cl₃C₂COO)₂ → Pb₃S₄ + 2NaCl₃C₂COO (Soluble complex)</td>
</tr>
</tbody>
</table>

* *Salt solution + drops of acidified solution of K₂Cr₂O₇*  
The purple colour of the permanganate will disappear. The sulphite reduces the permanganate to Mn²⁺ which is almost colourless.  
5Na₂SO₃ + 2K₂Cr₂O₇ + 3H₂SO₄ → 5Na₂SO₄ + K₂SO₄ + 2K₂MnO₄ + 3H₂O

### Thiosulphates

| A white precipitate forms (silver thiosulphate Ag₂S₂O₇₂⁻), which is soluble in excess of the thiosulphate due to the formation of complex, which is unstable; it changes to yellow, brown and finally to black Ag₂S₄. |
| Na₂S₂O₇ + 2AgNO₃ → Ag₂S₂O₇₂⁻ + 2NaNO₃ (Soluble complex) |

| A black ppt of lead sulphide. ( Pb₃S₄ ) is formed. |
| Na₂S₂O₇ + Pb(Cl₃C₂COO)₂ → Pb₃S₄ + 2NaCl₃C₂COO (Black ppt) |

### Sulphides

- **Salt solution + 2 drops of dil H₂SO₄ + Iodine solution(I₂)**

| The brown colour of the iodine solution disappears. |
| Na₂S₂O₃ + I₂ + H₂O → Na₂S₄O₆ + 2HI |

| The brown colour of the iodine solution disappears because the iodine is reduced to the iodide ion, which is colourless. At the same time, the thiosulphate is oxidized to tetrasulphate. |
| 2Na₂S₂O₃ + I₂ → Na₂S₄O₆ + 2NaI |
2) Anions which react with concentrated sulphuric acid

<table>
<thead>
<tr>
<th>Chlorides (Cl(^-))</th>
<th>Bromides (Br(^-))</th>
<th>Iodides (I(^-))</th>
<th>Nitrates (NO(_3))</th>
</tr>
</thead>
<tbody>
<tr>
<td>All chlorides are water soluble except the chlorides of silver, mercuric, and cuprous.</td>
<td>Bromides resemble chlorides in their solubility.</td>
<td>Iodides resemble chlorides and bromides in their solubility. However, bismuth iodide is insoluble.</td>
<td>All nitrates are soluble in water except some basic nitrates.</td>
</tr>
</tbody>
</table>


2NaCl + H\(_2\)SO\(_4\) → Na\(_2\)SO\(_4\) + 2HCl

Test for HCl gas: Dip a clean glass rod into a bottle of conc. ammonium solution, and hold it close to the mouth of the test tube in which the HCl is being produced. Dense white fumes of ammonium chloride will be formed. HCl + NH\(_3\) → NH\(_4\)Cl

Reddish fumes evolve and the solution turns orange due to liberation of bromine (Br\(_2\)).

2NaBr + H\(_2\)SO\(_4\) → 2HBr + Na\(_2\)SO\(_4\)

2HBr + H\(_2\)SO\(_4\) → 2H\(_2\)O + SO\(_2\) + Br\(_2\)

If a small piece of copper metal and drops of water are added, dense brown fumes of nitrogen dioxide (NO\(_2\)) will be given off.

KIO\(_3\) + H\(_2\)SO\(_4\) → K\(_2\)SO\(_4\) + 2H\(_2\)O + SO\(_2\) + I\(_2\)

N\(_2\)

2KNO\(_3\) + H\(_2\)SO\(_4\) → K\(_2\)SO\(_4\) + 2H\(_2\)O + 2NO\(_2\)

4H\(_2\)O + Cu → Cu(NO\(_3\))\(_2\) + 2H\(_2\)O + 2NO\(_2\)

 Confirmation tests

<table>
<thead>
<tr>
<th>Chlorides</th>
<th>Bromides</th>
<th>Iodides</th>
<th>Nitrates</th>
</tr>
</thead>
<tbody>
<tr>
<td>1) Salt solution + silver nitrate (AgNO(_3))</td>
<td>A yellow precipitate of silver iodide (AgI) is formed.</td>
<td>A yellow precipitate of silver iodide (AgI) is formed.</td>
<td>A yellow precipitate of silver iodide (AgI) is formed.</td>
</tr>
<tr>
<td>NaBr + AgNO(_3) → AgBr↓ + NaNO(_3)</td>
<td>KI + AgNO(_3) → AgI↓ + KNO(_3)</td>
<td>KIO(_3) + AgNO(_3) → AgI↓ + KNO(_3)</td>
<td>KNO(_3) + AgNO(_3) → AgI↓ + KNO(_3)</td>
</tr>
</tbody>
</table>

2) Salt solution + Lead acetate (Pb(CH\(_3\)CO\(_2\))\(_2\)):

A white precipitate of lead bromide (PbBr\(_2\)) appears which is soluble in boiling water and recrystallizes on cooling.

2NaBr + Pb(CH\(_3\)CO\(_2\))\(_2\) → PbBr\(_2\)↓ + 2CH\(_3\)COONa

A white precipitate of lead bromide (PbBr\(_2\)) appears which is soluble in boiling water and recrystallizes on cooling.

2KI + Pb(CH\(_3\)CO\(_2\))\(_2\) → PbI\(_2\)↓ + 2CH\(_3\)COOK

A reddish precipitate of lead bromide (PbBr\(_2\)) appears which is soluble in boiling water and recrystallizes on cooling.

Hgl\(_2\) + 2KI → KI↓ + HgI\(_2\)↓ + 2KCl

Hgl\(_2\) + 2KI → KI↓ + HgI\(_2\)↓

Brown ring test:
The nitrate solution is mixed with freshly prepared FeSO\(_4\) solution, then conc. H\(_2\)SO\(_4\) is added and allowed to flow cautiously on the side of the test tube. A brown ring (Fe\(_2\)(NO\(_3\))\(_3\)SO\(_4\) is formed at the interface of the two layers. The brown ring disappears on shaking the solution.

Salt solution + copper sulphate (CuSO\(_4\)):

A brown ppt. of copper iodide CuI and iodine I\(_2\):

2KI + CuSO\(_4\) → CuI↓ + I\(_2\)↓ + 2K\(_2\)SO\(_4\) + 1/2I\(_2\)↓
AgCl is white

Cul Cuprous Iodide

Brown Ring Test
<table>
<thead>
<tr>
<th>Phosphates (H₃PO₄, H₂PO₄⁻, HPO₄²⁻, PO₄³⁻)</th>
<th>Borates (BO₃⁻, BO₂⁻, B₂O₇⁻)</th>
<th>Sulfates (SO₄²⁻)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Most phosphates are insoluble in water except those of ammonium and alkali metals.</td>
<td>Ammonium and alkali metal borates are water soluble while other borates are slightly soluble in water.</td>
<td>All sulfates are soluble in water except those of some divalent metals e.g. calcium, strontium, barium and lead.</td>
</tr>
<tr>
<td>Salt solution + Barium chloride solution (BaCl₂)</td>
<td>A white precipitate of barium phosphate (Ba₃(PO₄)₂) is produced, soluble in dilute acids e.g. HNO₃ or HCl and insoluble in excess of barium chloride.</td>
<td>A white precipitate of barium sulphate (BaSO₄) is formed, which is insoluble in dilute acids and in excess of barium chloride.</td>
</tr>
<tr>
<td>Na₂HPO₄ + BaCl₂ → Ba₃(PO₄)₂↓ + 2NaCl</td>
<td>A white precipitate of barium borate from concentrated solutions is produced, soluble in dilute acids and in excess of barium chloride.</td>
<td>Na₂SO₄ + BaCl₂ → BaSO₄↓ + 2NaCl</td>
</tr>
<tr>
<td>Confirmatory test: 1) Salt solution + silver nitrate (AgNO₃)</td>
<td>A yellow precipitate of silver phosphate (Ag₃PO₄) is formed, which is readily soluble in dil. HNO₃ and ammonia.</td>
<td>A white precipitate of silver sulphate (Ag₂SO₄) is formed from concentrated solution.</td>
</tr>
<tr>
<td></td>
<td>A white precipitate of silver borate (Ag₂BO₃) is formed, from concentrated solution, which gives brown ppt. after boiling (also a brown ppt. is formed with dilute solution).</td>
<td>Na₂SO₄ + 2AgNO₃ → Ag₂SO₄↓ + 2NaNO₃</td>
</tr>
<tr>
<td></td>
<td></td>
<td>2AgBO₃ + 3H₂O → Ag₂O + 2H₂BO₃</td>
</tr>
</tbody>
</table>

BaSO₄ is white
For phosphates:

0.5 ml of salt solution + 4 ml of ammonium molybdate solution + 0.5 ml of conc. HNO₃

Form a canary yellow precipitate of ammonium phosphomolybdate (NH₄)₂H₂Mo₇O₂₄·2H₂O in the cold or by gentle warming.

(States the volume needed in this test).

For sulphates:

Salt solution + lead acetate:

A white precipitate of lead sulphate is formed, which is readily soluble in hot concentrated ammonium acetate or conc. H₂SO₄

K₂SO₄ + Pb(CH₂COO)₂ → PbSO₄ + 2CH₂COOK

PbSO₄ + H₂SO₄ → Pb(HSO₄)₂
**Scheme for the identification of acid radicals**

<table>
<thead>
<tr>
<th>Observation</th>
<th>Effervescence and a colourless odorous gas is evolved.</th>
<th>Pungent brown fumes are evolved.</th>
<th>Colourless gas with pungent odour, which turns an acidified dichromate paper green, is evolved.</th>
<th>Colourless gas is evolved which is not bad and turns a lead acetate paper black.</th>
<th>The salt does not react with dil. HCl.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Results</td>
<td>The gas is CO₂. The anion is carbonate or bicarbonate.</td>
<td>The gas is NO₂. The anion is Nitrite.</td>
<td>The gas is SO₂ and the green colour is Cr₂(SO₄)₃. The anion is sulphite.</td>
<td>The gas is SO₂, the ppt. is S and the green colour is Cr₂(SO₄)₃. The anion is thiosulphate.</td>
<td>The salt is not from group 1. (Go to step 2).</td>
</tr>
</tbody>
</table>

To differentiate between carbonate and bicarbonate:

1. **Salt solution - magnesium sulphate solution:**
   - **For carbonate:** A white precipitate of magnesium carbonate (MgCO₃).
   - **For bicarbonate:** No ppt. in the cold, but on heating, a white ppt. of magnesium carbonate is obtained.

2. **Solid salt + conc. H₂SO₄:**
   - **Observation** | Effervescence with the evolution of a colourless gas which forms white fumes with NH₃·OH. | Reddish fumes are evolved and the solution turns orange. | Violet fumes are evolved, and a brown or black precipitate is formed in the test tube. | The salt does not react with conc. H₂SO₄. |
   - **Results** | The gas is HCl and the white fumes are NH₃·Cl. The anion is chloride. | The gas is Br₂. The anion is Bromide. | The gas is I₂. The anion is iodide. | The anion is not from group 2. (Go to step 3). |

3. **Salt solution - BaCl₂:**
   - **Observation** | A white precipitate is produced, soluble in dilute acids e.g. HNO₃ or HCl and insoluble in excess of barium chloride. | A white ppt. from concentrated solutions is produced, soluble in dilute acids and in excess of barium chloride | A white precipitate is formed which is insoluble in dilute acids and in excess of barium chloride. |
   - **Results** | The white precipitate is barium phosphate (Ba₃(PO₄)₃). The anion is phosphate. | The white ppt. is barium borate Ba₃(BO₃)₃. The anion is Borate. | A white precipitate is barium sulphate (BaSO₄). The anion is sulphate. |
To differentiate between phosphate, borate and sulphate:

Salt solution + AgNO₃ (silver nitrate):

<table>
<thead>
<tr>
<th></th>
<th>phosphate</th>
<th>borate</th>
<th>Sulphate</th>
</tr>
</thead>
<tbody>
<tr>
<td>Observation</td>
<td>A yellow precipitate is formed, which is readily soluble in dil. HNO₃ and ammonia.</td>
<td>A white ppt. is formed from concentrated solution, which gives brown ppt. after boiling. (Also, a brown ppt. is formed with dilute solution.)</td>
<td>A white ppt. is formed, with concentrated solution.</td>
</tr>
<tr>
<td>Results</td>
<td>The yellow ppt. is silver phosphate (Ag₂PO₄). The anion is phosphate.</td>
<td>The white ppt. is silver borate (Ag₂BO₄). The anion is borate.</td>
<td>The white ppt. is silver sulphate (Ag₂SO₄). The anion is sulphate.</td>
</tr>
</tbody>
</table>

Which one of the following is the smallest in size?
(a) N³⁻ (b) O²⁻ (c) F⁻ (d) Na⁺

Ans : ( d )

Na⁺ will have the smallest size

Group 14 Elements—Carbon Family

Group 14 contains carbon (C), silicon (Si), germanium (Ge), tin (Sn) and lead (Pb). Their outer electronic configuration is (ns)² (np)², where n varies from 2 to 6. The metallic character of elements increases on descending the group; C and Si are nonmetals, Ge is a nonmetal but also has some metallic characteristics, and Sn and Pb are metals. The melting points decrease on descending the group, with the exception of Pb whose melting point is slightly higher than that of Sn. Carbon has extremely high melting point. This is due to the stronger C—C bonds in the network of carbon atoms.

The ionization energies decrease from C to Si, but then change in an irregular way because of the effects of filling d and f sub-shells.

Carbon forms single, double and triple bonds with carbon itself and with other elements. The tendency to form multiple bond by other elements is rare. However, silicon can form double bond due to back bonding in which the lone pair in p orbitals of an atom is extended to an empty orbital of Si. One of the examples of back bonding is trisilylethane, N(SiH₂)₃.

The chemical reactivity of elements decreases down the group. The inert pair effect becomes increasingly effective on descending the group.

The stability of +4 oxidation state decreases while that of the +2 oxidation state increases on descending the group. C, Si and Ge are unaffected by dilute acids. Sn and Pb react with dilute nitric acid.

\[ 4Sn + 10HNO₃ \rightarrow 4Sn(NO₃)₂ + NH₄NO₃ + 3H₂O \]
\[ 3Pb + 8HNO₃ \rightarrow 3Pb(NO₃)₂ + 2NO + 4H₂O \]

C is not affected by alkalis, Si reacts forming silicates while Sn and Pb form stannate, \([Sn(OH)₄]²⁻\), and plumbate, \([Pb(OH)₄]²⁻\), respectively.
The elements of Group 14 form tetrahalides with the exception of PbCl₄, which is not known. The stability of halides decreases down the group. CCl₄ is stable while other halides are hydrolysed. The hydrolysis of SiCl₄ produces SiO₂ while SiF₄ produces SiOF₂ as well as [SiF₄]²⁻.

The acidic nature of the oxides of carbon family decreases down the group; CO₂ and SiO₂ are acidic, GeO₂ is weakly acidic and SnO₂ and PbO₂ are amphoteric.

Silicones are organosilicon polymers with general formula (R₂SiO)n, where R may be methyl, ethyl or phenyl group.

**Group 15 Elements—Nitrogen Family**

Group 15 contains nitrogen (N), phosphorus (P), arsenic (As), antimony (Sb) and bismuth (Bi). Their outer electronic configuration is (ns)² (np)³, where n varies from 2 to 6. The metallic character of these elements increases on descending the group; N and P are nonmetals, As and Sb are metalloids and Bi is a metal. The melting and boiling points follow the order:

- Melting point: N < P < As < Sb < Bi
- Boiling point: N < P < As < Sb < Bi

Phosphorus has two common allotropes; white and red. White phosphorus is more reactive than red form due to highly strained structure (P—P—P angle is 60°).

Black phosphorus is a highly polymerized form and is most stable. Nitrogen forms triple bond in dinitrogen because bond enthalpy ε(N—N) is greater than three times bond enthalpy ε(N—N). In phosphorus, the reverse is true, hence, it involves single bonds.

The melting points of hydrides follow the order NH₃ > PH₃ > AsH₃ > SbH₃. All the five elements of Group 15 form trihalides. Of these, nitrogen halides are least stable. All the trihalides with the exception of NF₃ hydrolyse in aqueous solution.

\[
\begin{align*}
\text{NCl₃} + \text{3H₂O} & \rightarrow \text{NH₃} + \text{3HCl} \\
\text{PCl₃} + \text{3H₂O} & \rightarrow \text{H₃PO₃} + \text{3HCl} \\
\text{AsCl₃} + \text{3H₂O} & \rightarrow \text{H₃AsO₃} + \text{3HCl} \\
\text{SbCl₃} + \text{H₂O} & \rightarrow \text{SbO}^{+} + \text{2H}^{+} + \text{3Cl}^{-} \\
\text{BiCl₃} + \text{H₂O} & \rightarrow \text{BiO}^{+} + \text{2H}^{+} + \text{3Cl}^{-}
\end{align*}
\]

Nitrogen forms many oxides: N₂O, NO, N₂O₃, NO₂ and N₂O₅. The most common oxides of phosphorus are P₂O₅ and P₄O₁₀. Phosphorus trioxide is an acidic oxide and anhydride of orthophosphorous acid (H₃PO₃). Phosphorus pentoxide is anhydride of phosphoric acid (H₃PO₄).

Nitrogen forms two acids; nitrous acid (HNO₂) and nitric acid (HNO₃). Phosphorus forms two series of oxoacids. These are phosphorus and phosphoric series. In phosphorus series, the acids include pyrophosphoric acid (H₂P₂O₇), orthophosphorous acid (H₃PO₄), metaphosphoric acid (H₄P₂O₇) and hypophosphorous acid (H₃PO₃). The acids H₃P₂O₇ and H₄P₂O₇ are reducing agents as they involve P—H bond.

In phosphoric series, the acids include orthophosphoric acid (H₃PO₄), pyrophosphoric acid (H₄P₂O₇) and polyphosphoric acid, (H₃PO₄)ₙ.

Which one of the following has the minimum atomic size?

(a) Li  (b) Be  (c) Na  (d) Mg

**Ans : (b)**

In the periodic table, size decreases in a period and it increases in a group. In the given elements, Be belongs to lower period and higher group and thus will have least size.

**Group 1B**

The size of Zr is very similar to Hf due to Lanthanide contraction

**Group 4A**
The size of Nb is very similar to Tantalum (Ta) due to Lanthanide contraction

Oxygen Family

Group 16 Elements—Oxygen Family

Group 16 contains oxygen (O), sulphur (S), selenium (Se), tellurium (Te) and polonium (Po). Their electronic configuration is \((nsp)^2\)\(^{(n-1)}p\)^{n}, where \(n\) varies from 2 to 6.

The metallic character of these elements increases on descending the group; O and S are nonmetallic, Se and Te are weaker nonmetallic and Po is metallic.

Electron affinity of sulphur is larger than that of oxygen, this is due to the more repulsion experienced by the incoming electron from the smaller, more compact electronic cloud of oxygen atom.

Oxygen is diatomic with two unpaired electrons. Sulphur exists in two allotropic forms—rhombic and monoclinic sulphur. Rhombic sulphur is stable at room temperature while monoclinic sulphur is stable above 369 K. Selenium exists in six allotropic forms. Tellurium has only one crystalline form and polonium has two allotrophic forms (cubic and rhombohedral).

Ozone is another allotrope of oxygen. It is very reactive. It is formed in the upper layer of atmosphere (about 20 km from the earth) by the action of ultraviolet radiation on oxygen. Ozone is a strong oxidizing agent. In organic chemistry, ozone is used to locate the carbon-carbon double and triple bonds. The ozone molecule is angular with bond angle about 117° and bond length 1.278 pm.

The melting and boiling points of hydrogen compounds of elements of Group 16 follow the order \(H_2O > H_2S < H_2Se < H_2Te\). The exceptional high values of \(H_2O\) is due to hydrogen bondings.

\(H_2O\) is a strong oxidizing agent. With stronger oxidizing agents such as \(KMnO_4, KIO_3, \text{O}_2\), hydrogen peroxide acts as a reducing agent.

Amongst Co, Ni, Cu and Zn; Zn has maximum size
The monohydrogen and dihydrogen phosphates have very important uses in real-world analytical laboratories. Mostly, these use centers around the control of the acidity of solutions. Solutions of these partially neutralized acids resist changes in acidity level even when strong acids or bases are added. For this reason, they are called “buffer solutions.” For example, dilute solutions of potassium dihydrogen phosphate, \( \text{KH}_2\text{PO}_4 \), will maintain an acidity level near the neutral point between acids and bases. Solutions of sodium monohydrogen, \( \text{NaH}_2\text{PO}_4 \), will maintain an acidity level that is considerably below the neutral point, which means that base-like properties predominate.

The size of Mo is very similar to W due to
(a) the difference of atomic number by one
(b) the contraction in size in the first transition series elements
(c) lanthanide contraction
(d) actinide contraction

Mo belongs to 5\(^{th}\) period and 6\(^{th}\) group. W belongs to 6\(^{th}\) period and 6\(^{th}\) group. Their sizes are more or less identical due to intervening lanthanides.

The solution from which excess \( \text{Br}_2 \) is removed by heating must be acidic, otherwise a portion of \( \text{Br}_2 \) shall be fixed in the form of potassium hypobromite (KBrO).

With alkalis it forms a hypobromite in cold dilute solution and a bromate in a hot solution

\[
2\text{KOH} + \text{Br}_2 \rightarrow \text{KBr} + \text{KBrO} + \text{H}_2\text{O}
\]

\[
6\text{KOH} + 3\text{Br}_2 \rightarrow 5\text{KBr} + \text{KBrO}_3 + 3\text{H}_2\text{O}
\]

The first compound of xenon, \( \text{XePtF}_6 \), was discovered in 1962. Xenon reacts with fluorine to give three compounds they are \( \text{XeF}_2 \), \( \text{XeF}_4 \), and \( \text{XeF}_6 \). Other known compounds of xenon are \( \text{XeO}_2 \), \( \text{XeO}_4 \), \( \text{XeOF}_2 \), \( \text{XeO}_2\text{F}_2 \), etc. No compound of He, Ne and Ar has been prepared.
Many salts of ternary acids are used in medicine. Lithium carbonate, Li₂CO₃, has been used successfully to combat severe jet lag. Lithium carbonate is also useful in the treatment of mania, depression, alcoholism, and schizophrenia. Magnesium sulfate, MgSO₄, sometimes helps to prevent convulsions during pregnancy and to reduce the solubility of toxic barium sulfate in internally administered preparations consumed before gastrointestinal X-ray films are taken.

Other salts of ternary acids that you may find in your home include potassium chlorate, KClO₃, in matches as an oxidizing agent and oxygen source; sodium hypochlorite, NaClO, in bleaches and mildew removers; and ammonium carbonate, (NH₄)₂CO₃, which is the primary ingredient in smelling salts.

The tips of “strike anywhere” matches contain tetraphosphorus trisulfide, red phosphorus, and potassium chlorate. Friction converts kinetic energy into heat, which initiates a spontaneous reaction.

$$P₄S₃(s) + 8O₂ \rightarrow P₄O₁₀(s) + 6SO₂(g)$$

The thermal decomposition of KClO₃ provides additional oxygen for this reaction.

Group 17 Elements—Halogen Family

Group 17 contains fluorine (F), chlorine (Cl), bromine (Br), iodine (I) and astatine (At). Their outer electronic configuration is \((n+1)p^5\), where \(n\) varies from 2 to 6. The trends in this group are as follows.

- Covalent and ionic radii—增加 down the group.
- Electronegativity and ionization energy—decrease down the group.
- Electron affinity—increases from F to Cl followed by a decrease.
- Melting and boiling points—Increase down the group.
- Bond enthalpy \(\varepsilon(X-X)\)—increases from F to Cl followed by a decrease.
- Oxidizing ability—decreases down the group.
- Halogens are very reactive and do not occur in free state. Fluorine is most electronegative atom, there exists hydrogen bondings in gaseous HF.
- HF is a weak acid and HCl, HBr and HI behave as strong acids. In the glacial acetic acid medium, the acid strength follows the order HI > HBr > HCl > HF.
Halogenes with the exception of F form a number of oxoacids—hypohalous acids (H\(_2\)X), halous acids (H\(_2\)X\(_2\)), halic acid (H\(_2\)X\(_3\)) and perhalic acid (H\(_2\)X\(_4\)). The acid strength follows the order H\(_2\)X\(_4\) > H\(_2\)X\(_3\) > H\(_2\)X\(_2\) > H\(_2\)X. Fluorine forms only hypohalous acid (HOF).

Halogenes also form interhalogen compounds AX, AX\(_2\), AX\(_3\) and AX\(_4\). The compounds AX and AX\(_2\) are formed where the electronegativity is not very large. The compounds AX\(_2\) and AX\(_3\) are formed by large Br and I atoms surrounded by small atom F. The molecule AX\(_4\) is T-shaped with two lone pair of electrons at the equilateral positions. The orientations of five pair of electrons around the atom A is trigonal bipyramidal.

The six pairs of electrons around atom A in AX\(_4\) acquire octahedral orientation. The seven pairs of electrons around atom A in AX\(_3\) acquire pentagonal bipyramidal orientation.

Halogenes are nonmetallic and have high electron affinity. The nonmetallic character decreases down the group and iodine shows some metallic character. With metals, they form ionic compounds by accepting one electron and with nonmetals covalent compounds are formed by sharing an electron.

All halogenes exhibit -1 oxidation state. Except fluorine, rest of the halogenes also exhibit 1, +3, +5 and +7 oxidation states. Fluorine is the most electronegative and thus there exists hydrogen bonding in HF with the result that it has exceptionally high melting and boiling points as compared to those of HCl, HBr and HI.

Because of high reactivity, halogenes do not exist in the free state.

**OXIDES OF CHLORINE**

Chlorine reacts with oxygen to form a number of oxides like Cl\(_2\)O, ClO\(_2\), ClO\(_3\), and ClO\(_4\). All these oxides are highly reactive and very unstable.

**Dichloro Oxide (Cl\(_2\)O)**

**Preparation**

- It can be prepared by passing dry chlorine over a fresh precipitate of yellow mercuric oxide as follows:
  
  \[ \text{HgO} + 2\text{Cl}_2 \text{(dry)} \rightarrow \text{HgCl}_2 + \text{Cl}_2 \text{O} \]

**Physiochemical Properties**

1. It is a brownish yellow gas with a penetrating odour.
2. It can be condensed to an orange coloured liquid in a freezing mixture (boiling point = 275 K).
3. Decomposition: It decomposes on heating or in sunlight as follows:
   
   \[ 2\text{Cl}_2\text{O} \rightarrow 2\text{Cl}_2 + \text{O}_2 \]
Group 18 Elements—Noble Gases

The group 18 contains helium (He), neon (Ne), argon (Ar), krypton (Kr), xenon (Xe) and radon (Rn). Their outer electronic configurations is \((n{s})^2 (n\pi)^6\) with the exception of the electronic configuration of He which is \((1s)^2\). All elements exist as monatomic gas.

Xenon forms a number of fluorides—\(\text{XeF}_2\), \(\text{XeF}_4\) and \(\text{XeF}_6\). The other compounds are \(\text{XeO}_3\), \(\text{XeOF}_4\), \(\text{XeO}_2\text{F}_2\), \(\text{XeOF}_2\) and \([\text{XeOF}_4]^{2-}\)

4. With Water: It dissolves in water forming a golden yellow solution of hypochlorous acid (HClO) i.e., it is an anhydride of hypochlorous acid.

\[
\text{Cl}_2\text{O} + \text{H}_2\text{O} \rightleftharpoons 2\text{HClO}
\]

5. Oxidizing Action: Being a strong oxidizing agent, it oxidizes HCl to \(\text{Cl}_2\) as follows:

\[
\text{Cl}_2\text{O} + 2\text{HCl} \rightarrow 2\text{Cl}_2 + \text{H}_2\text{O}
\]

6. With Ammonia: Its reacts with ammonia with an explosion as follows:

\[
3\text{Cl}_2\text{O} + 10\text{NH}_3 \rightarrow 2\text{N}_2 + 6\text{NH}_4\text{Cl} + 3\text{H}_2\text{O}
\]

**Structure** It has angular or V-shaped structure. Here oxygen atom is \(sp^3\) hybridized. \(\text{Cl} - \text{O} - \text{Cl}\) bond angle is 110.9\(^\circ\) and \(\text{O} - \text{Cl}\) bond length is 1.71 Å
Chlorine Dioxide (ClO₂)

Preparation

1. Pure ClO₂ can be obtained by passing dry Cl₂ over AgClO₃ heated to 363 K as follows:
   \[ 2\text{AgClO}_3 + \text{Cl}_2 (\text{dry}) \rightarrow 2\text{AgCl} + 2\text{ClO}_2 + \text{O}_2 \]

2. It can also be prepared by the action of Cl₂ on sodium chlorite as follows:
   \[ 2\text{NaClO}_2 + \text{Cl}_2 \rightarrow 2\text{NaCl} + 2\text{ClO}_2 \]

Physiochemical Properties

1. It can be condensed by cooling to a coloured liquid (boiling point 284 K).
2. It is a powerful oxidizing and bleaching agent.
3. Decomposition: It explodes and decomposes to Cl₂ and O₂ by an electric spark.
4. With H₂O: It dissolves in water to give a mixture of chlorous acid and chloric acid.
From the X-ray diffraction of nitric oxide it is found to be present in a dimeric orientation. It has a resonating dimeric structure.

Structure of $\text{(NO)}_2$

Group 12 - Electrode potential of Mercury is $+\text{ve}$, 1st ionisation Energy higher than Cd

5. **With Alkalies:** It gives a mixture of chlorite and chlorate with alkalies as follows:

$$2\text{ClO}_2 + \text{H}_2\text{O} \rightarrow \text{HClO}_2 + \text{HClO}_3$$

**Structure:** It is an odd electron molecule and paramagnetic in nature. It is an angular molecule with $\text{sp}^3$ hybridization of chlorine. Here $\text{O} - \text{Cl} - \text{O}$ bond angle is 118° and $\text{Cl} - \text{O}$ bond length is 1.47 Å.
Dichlorine Hexoxide \( (\text{Cl}_2\text{O}_6) \)

**Preparation**

1. It can be prepared by mixing chlorine dioxide with ozonized air at 273 K.

\[
2\text{ClO}_2 + 2\text{O}_3 \rightarrow \text{Cl}_2\text{O}_6 + 2\text{O}_2
\]

**Physicochemical Properties**

1. It is a dark red unstable liquid.
2. Decomposition: It is quite unstable and decomposes into \( \text{ClO}_2 \) and \( \text{O}_2 \).
3. It is a strong oxidizing agent.
4. With Water: It reacts with water to give chloric acid and perchloric acids.

\[
\text{Cl}_2\text{O}_6 + \text{H}_2\text{O} \rightarrow \text{HClO}_3 + \text{HClO}_4
\]

5. With HF: It gives \( \text{HClO}_4 \) and chloryl fluoride with HF as follows:

\[
\text{Cl}_2\text{O}_6 + \text{HF} \rightarrow \text{HClO}_4 + \text{ClO}_2\text{F}
\]

6. With Alkalis: It reacts with alkalis to give chlorates and perchlorates.

5. With HF: It gives \( \text{HClO}_4 \) and chloryl fluoride with HF as follows:

\[
\text{Cl}_2\text{O}_6 + \text{HF} \rightarrow \text{HClO}_4 + \text{ClO}_2\text{F}
\]

6. With Alkalis: It reacts with alkalis to give chlorates and perchlorates.
Structure: In liquid state it is a diamagnetic molecule with uncertain structure which is not known so far.

$2\text{Cl}_2\text{O}_5 \rightleftharpoons \text{Cl}_2\text{O}_6$

In the vapour state it exists as $\text{ClO}_3$ molecule which has odd number of electrons and hence paramagnetic.

In solid state it has $\text{ClO}_4^-$, $\text{ClO}_2^+$ ions.

Although the exact structure is unknown for it yet the following structure having Cl−Cl linkage is proposed for it in which each chlorine atom is sp$^3$ hybridized.

![Structure Diagram](image)

**Chlorine Heptoxide ($\text{Cl}_2\text{O}_7$)**

**Preparation**

It can be formed by the dehydration of perchloric acid with $\text{P}_2\text{O}_5$ at 263 K.

$$2\text{HClO}_4 \xrightarrow{\text{P}_2\text{O}_5} \text{Cl}_2\text{O}_7 + \text{H}_2\text{O}$$

**Physicochemical Properties**

1. It is a colourless oily explosive liquid.
2. With Water: It slowly dissolves in water to form perchloric acid i.e., it is an anhydride of perchloric acid.

$$\text{Cl}_2\text{O}_7 + \text{H}_2\text{O} \rightarrow 2\text{HClO}_4$$

**Structure:** In it two $\text{ClO}_4$ units are joint through oxygen at an angle of 118.5° or 118°36' giving tetrahedral look. Here Cl−O bond length is 1.72 Å while Cl−O$_3$ is 1.42 Å.
### (1) Oxoacids of Nitrogen

(i) Hyponitrous acid \( \text{H}_2\text{N}_2\text{O}_2 \)  
(ii) Hydronitrous acid \( \text{H}_2\text{NO}_2 \)  
(iii) Nitrous acid \( \text{HNO}_2 \)  
(iv) Hyponitric acid \( \text{H}_2\text{N}_2\text{O}_3 \)  
(v) Nitric acid \( \text{HNO}_3 \)  
(vi) Pernitric acid \( \text{HNO}_4 \)  

### (2) Oxoacids of Phosphorous

(i) Hypophosphorous acid \( \text{H}_3\text{PO}_2 \)  
(ii) Phosphorous acid \( \text{H}_3\text{PO}_3 \)  
(iii) Hypophosphoric acid \( \text{H}_3\text{P}_2\text{O}_6 \)  
(iv) Orthophosphoric acid \( \text{H}_3\text{PO}_4 \)  
(v) Pyrophosphoric acid \( \text{H}_4\text{P}_2\text{O}_7 \)  
(vi) Metaphosphoric acid \( \text{HPO}_3 \)  
(vii) Peroxophosphoric acid \( \text{H}_3\text{PO}_5 \text{ or } \text{H}_4\text{P}_2\text{O}_8 \)  
(viii) Triplyphosphoric acid \( \text{H}_3\text{P}_3\text{O}_{10} \)  
(ix) Tetra polyphosphoric acid \( \text{H}_6\text{P}_4\text{O}_{13} \)
Oxy-acids of Chlorine

Chlorine forms all the four oxy-acids, which are hypochlorous acid (HClO), chlorous acid (HClO₂), chloric acid (HClO₃) and perchloric acid (HClO₄). It may be noted that Cl-atom is in +1, +3, +5 and +7 oxidation state respectively in these acids.

HClO, HClO₂, HClO₃, HClO₄

The acidic strength of these acids increases as follows:

HClO < HClO₂ < HClO₃ < HClO₄

Hypochlorous Acid (HClO)

Preparation

1. From Chlorine

   (i) It can be prepared by passing Cl₂ into H₂O or into a suspension of CaCO₃ in water or into an aqueous solution of potassium hypochlorite (KClO) or bleaching powder (CaOCl₂).

   \[
   \text{Cl}_2 + \text{H}_2\text{O} \rightarrow \text{HCl} + \text{HClO}
   \]

   \[
   \text{Cl}_2 + \text{H}_2\text{O} \rightarrow \text{Cl}^- + \text{ClO}^- + 2\text{H}^+
   \]
Co-ordination Complexes

2. It can be prepared by the action of atmospheric CO₂ on sodium hypochlorite, NaOCl as shown.

\[
\text{NaOCl} + \text{CO}_2 + \text{H}_2\text{O} \rightarrow \text{NaHCO}_3 + \text{HClO}
\]

3. From bleaching powder Hypochlorous acid can be prepared by distilling the aqueous solution of bleaching powder (CaOCl₂) with a calculated quantity of 5 per cent HNO₃ or by passing CO₂ into aqueous solution of CaOCl₂ and then distilling it.

\[
2\text{CaOCl}_2 + 2\text{HNO}_3 \rightarrow \text{CaCl}_2 + \text{Ca(NO}_3)_2 + 2\text{HClO}
\]

\[
\text{CaOCl}_2 + \text{H}_2\text{O} + \text{CO}_2 \rightarrow \text{CaCO}_3 + 2\text{HClO}
\]

Physicochemical properties

1. The dilute solution of hypochlorous acid is colourless while its concentrated solution is yellow in colour.

2. It is a weak acid, even weaker than H₂CO₃ with a dissociation constant of 3 × 10⁻⁸ at 20°C.

3. Acidic nature: (Monobasic nature) HClO is a monobasic acid as its aqueous solution on ionization gives only one H⁺ ion.

\[
\text{HClO (aq)} \rightleftharpoons \text{H}^+ + \text{(aq)} + \text{ClO}^-\text{(aq)}
\]

The monobasic nature shows that this molecule has one –OH group attached directly with the central Cl atom.
Being an acid it reacts with alkalies to give salts which are called hypochlorites.

\[
\text{NaOH} + \text{HClO} \rightarrow \text{NaOCl} + \text{H}_2\text{O} \\
\text{OH}^- + \text{HClO} \rightarrow \text{OCl}^- + \text{H}_2\text{O}
\]

4. Disproportion

(i) The dilute solution of hypochlorous acid is quite stable in the dark but when the concentrated solution is exposed to light, it becomes unstable and undergoes disproportionation to form Cl₂ and HClO₄.

\[
5\text{HClO} \rightarrow 2\text{Cl}_2 + \text{HClO}_4 + 2\text{H}_2\text{O}
\]

(ii) When aqueous solution of HClO is heated, it undergoes disproportionation into HCl and HClO₄.

\[
\text{3HClO} \underset{\Delta}{\rightarrow} 2\text{HCl} + \text{HClO}_4 \\
\text{3ClO}^- \underset{\Delta}{\rightarrow} 2\text{Cl}^- + \text{ClO}_4^-
\]

(iii) On distillation it decomposes into H₂O and Cl₂O₃.

\[
2\text{HClO} \rightarrow \text{Cl}_2\text{O} + \text{H}_2\text{O}
\]

5. Reaction with metals: HClO reacts with Mg with the evolution of H₂.

\[
\text{Mg} + 2\text{HClO} \rightarrow \text{Mg(OCl)}_2 + \text{H}_2
\]

It reacts with Fe and Al with the evolution of H₂ and Cl₂ while with Co, Ni and Cu, Cl₂ and O₂ are evolved.
6. Reaction with mercury: When it is shaken with Hg, a light brown precipitate of basic mercuric chloride, HgCl(OH) is formed which is soluble in HCl.

\[ 2\text{Hg} + 2\text{HClO} \rightarrow 2\text{HgCl} + \text{H}_2\text{O} \]
Light brown ppt.

7. Oxidizing and bleaching properties: The aqueous solution of HClO and its salts (like, NaOCl) are oxidizing and bleaching agents. It is due to the fact that HClO or NaOCl decomposes to give nascent oxygen as follows.

\[
\begin{align*}
\text{HClO} & \xrightarrow{\text{Reduction}} \text{HCl} + [\text{O}] \\
\text{NaOCl} & \xrightarrow{\text{Reduction}} \text{NaCl} + [\text{O}]
\end{align*}
\]

\[ \text{ClO}^- + 2\text{H}^+ + 2e^- \xrightarrow{\text{Reduction}} \text{Cl}^- + \text{H}_2\text{O} \]
(lion electron equation)

8. Action of AgNO₃: It reacts with AgNO₃ to give silver hypochlorite (AgClO) which is unstable and undergoes disproportionation into AgCl and AgClO₃.

\[ \text{HOCI} + \text{AgNO}_3 \rightarrow \text{AgClO} + \text{HNO}_3 \]

\[ 3\text{AgClO} \rightarrow 2\text{AgCl} + \text{AgClO}_3 \]

Uses:

(i) Hypochlorous acid is used for bleaching paper pulp etc.
**Structure**  
It is a linear molecule in which chlorine atoms is sp² hybridized.

![H-O-Cl](image)

**Figure 16.38  Structure of Hypochlorous Acid**

**Chlorous acid (HClO₂)**

**Preparation**

1. **From Barium hypochlorite**  
   It is prepared in the aqueous solution by treating a suspension of barium hypochlorite, **Ba(ClO₂)₂**, with dil. **H₂SO₄** and filtering off the precipitate of **BaSO₄**.
   
   \[ \text{Ba(ClO}_2\text{)}_2 + \text{H}_2\text{SO}_4 \rightarrow 2\text{HClO}_2 + \text{BaSO}_4 \]

2. **From ClO₂**  
   It can also be obtained by the action of ClO₂ on H₂O₂.
   
   \[ 2\text{ClO}_2 + \text{H}_2\text{O} \rightarrow 2\text{HClO}_2 + \text{O}_2 \]

**Physicochemical properties**

1. The freshly prepared solution of chlorous acid is colourless but it soon decomposes to ClO₂ which colours the solution yellow.

2. It gives a violet colour with FeSO₄.

3. **Decomposition:** In acid solution, HClO₂ rapidly decomposes as shown.
4. Disproportionation: HClO₄ and its salts undergo disproportionation on heating in an alkaline solution as shown.

\[ 3\text{HClO}_4 \rightarrow 2\text{HClO}_3 + \text{HCl} \]

\( \text{Cl} = +5 \quad \text{Cl} = -1 \)

\[ 3\text{ClO}_2 \rightarrow 2\text{ClO}_3^- + \text{Cl}^- \]

It undergoes auto-oxidation.

\[ 2\text{HClO}_4 \rightarrow \text{HClO}_4 + \text{HClO}_3 \]

5. Oxidizing properties: Both HClO₄ and its salts show oxidizing properties due to nascent oxygen.

\[ \text{HClO}_4 \rightarrow \text{HCl} + 2[\text{O}] \]

\[ \text{ClO}_2^- + 4\text{H}^+ + 4\text{e}^- \rightarrow \text{Cl}^- + 2\text{H}_2\text{O} \]

HClO₄ and its salts liberate I₂ from KI solution.

\[ \text{HClO}_4 + 2\text{H}_2\text{O} + 4\text{KI} \rightarrow 4\text{KOH} + \text{HCl} + I_2 \]

\[ 4\text{I}^- + \text{ClO}_2^- + 4\text{H}^+ \rightarrow \text{Cl}^- + 2\text{H}_2\text{O} + 2I_2 \]

Structure: ClO₂⁻ is angular in shape as chlorine atom is sp³ hybridized.

Parke’s process-Pyrometallurgical extraction of Ag (Silver) from Pb (Lead) Chemistry

MaC Arthur Forrest cyanide process to extract Au, Ag by KCN, Na Cyanide Chemistry

Deacon’s process-Obtaining Cl₂ from HCl passed on hot saturated slag Chemistry

Cu₂Fe(CN)₆ Chocolate Brown Precipitate

Cd(CN)₂, K₂SO₄, KCN are all white - Cd(CN)₂ white precipitate

Ag₂S₂O₃ White Precipitate

2FeSO₄ + 2H₂SO₄ green vitriol, ZnSO₄ White Vitriol, CuSO₄ Blue Vitriol

Ti⁺³ Violet, V⁺³ Grey, Fe⁺³ yellow, Co⁺² Pink - ion colours

Silver nitrate (Lunar caustic) AgNO₃
# Oxidation States and Colors

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<th>Metal</th>
<th>+1</th>
<th>+2</th>
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<th>+4</th>
<th>+5</th>
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<td>blue</td>
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<tr>
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<td></td>
<td>Green</td>
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<td></td>
<td>Yellow (CrO$_4^{2-}$)</td>
<td>orange (Cr$_2$O$_7^{2-}$)</td>
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<tr>
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<td>Brown</td>
<td></td>
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<td>Intense purple (MnO$_4$)</td>
</tr>
<tr>
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<tr>
<td>Co</td>
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<td>Orange/yellow</td>
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