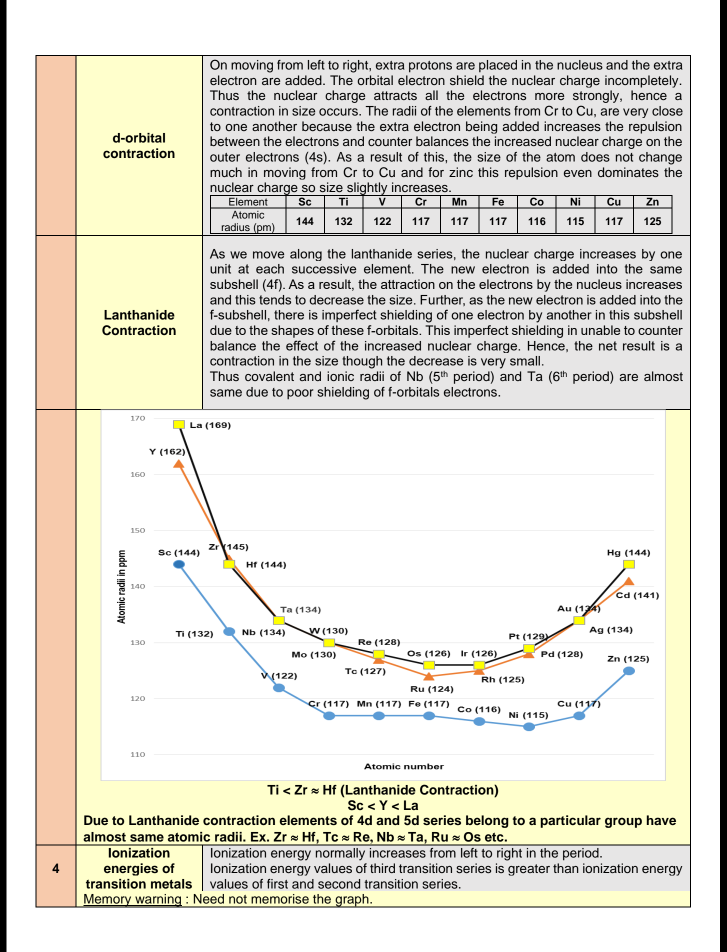
d & f-block elements & their important compounds

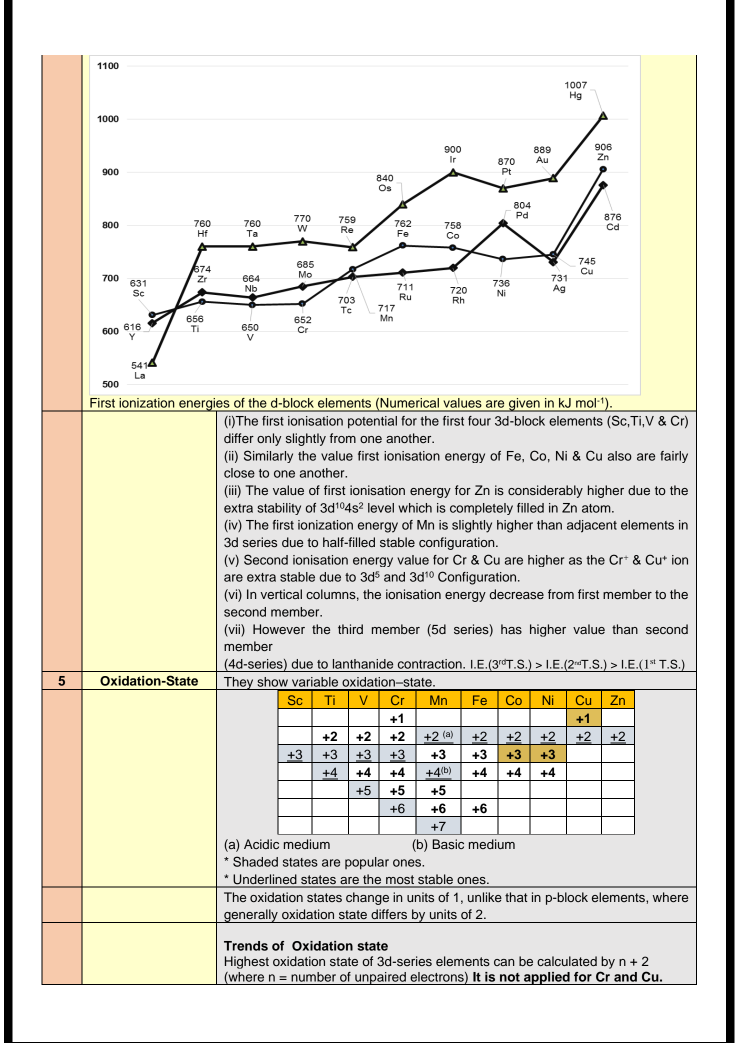
Section (A) : Electronic configuration, atomic size and ionic size, density, melting and boiling points, lonization enthalpy and oxidation state.

Element		Sc	Ti	V	Cr	Mn	Fe	Со	Ni	Cu	Zn
Atomic number		21	22	23	24	25	26	27	28	29	30
Electronic con	figuration										
	М	3d ¹ 4s ²	3d ² 4s ²	3d ³ 4s ²	3d ⁵ 4s ¹	3d ⁵ 4s ²	3d ⁶ 4s ²	3d ⁷ 4s ²	3d ⁸ 4s ²	3d ¹⁰ 4s ¹	3d ¹⁰ 4s ²
	M+	3d ² 4s ⁰	3d ³ 4s ⁰	3d ⁴ 4s ⁰	3d⁵	3d ⁶ 4s ⁰	3d ⁷ 4s ⁰	3d ⁸ 4s ⁰	3d ⁹ 4s ⁰	3d ¹⁰	3d ¹⁰ 4s ¹
	M ²⁺	3d ¹	3d ²	3d ³	3d ⁴	3d⁵	3d ⁶	3d ⁷	3d ⁸	3d ⁹	3d ¹⁰
	M ³⁺	[Ar]	3d ¹	3d ²	3d ³	3d ⁴	3d⁵	3d ⁶	3d ⁷	_	_
Enthalpy of ate	omisation,	Δ _a H⊕ / kJ	mol ⁻¹								
		326	473	515	397	281	416	425	430	339	126
Ionisation enth	nalpy / ∆ _i H⁻	⁻ / kJ mol⁻	-1								
∆iH⊕	I	631	656	650	653	717	762	758	736	745	906
∆iH⊕	II	1235	1309	1414	1592	1509	1561	1644	1752	1958	1734
∆iH⊕		2393	2657	2833	2990	3260	2962	3243	3402	3556	3829
Metallic/ionic	М	164	147	135	129	137	126	125	125	128	137
radii/pm	M ²⁺	_	_	79	82	82	77	74	70	73	75
	M ³⁺	73	67	64	62	65	65	61	60	_	-
Standard electrode		I	-1.63	-1.18	-0.90	-1.18	-0.44	-0.28	-0.25	+0.34	-0.76
Potential E [⊕] / V	M ³⁺ /M ²⁺	-	-0.37	-0.26	-0.41	+1.57	+0.77	+1.97	_	-	_

	d-block Elements									
Last	Last electron enters in (n – 1)d subshell. d-block elements lie in middle of periodic table.									
	General Introduction									
1.	Occurrence	Three of transition metals are very abundant in the earth's crust.								
		$Fe \rightarrow 4^{th}$ most abundant elements in earth's crust by weight. Ti $\rightarrow 5^{th}$ most abundant elements in earth's crust by weight. Mn $\rightarrow 12^{th}$ most abundant elements in earth's crust by weight.								
	First row Element with even atomic no. are in general more abundant than their neighbours with odd atomic no. except for Mn ($Z = 25$) but still 12 th most abundant element by weight.									
	2nd and 3rd row	Less abundant than 1 st d- series elements. Tc → does not occur in nature. Last six members of 2 nd and 3 rd row occur less than 0.16 ppm in earth's crust.								
	Coinage metals	Cu, Ag and Au are called coinage metals.								
	Noble metals	Cu, Ag, Au, Hg and Pt, etc.								
2.	No. of elements	Total d-block elements – (40) Total transition elements – (36). If 112 elements are included in periodic table (Except Zn, Cd, Hg and Uub)	-							
3.	Groups	IIIB to VIIB, VIII, IB, IIB or group 3 to 12 (IUPAC).								
4.	Period	$\begin{array}{c c} 4^{th} \text{ to } 7^{th} \\ \hline & 4^{th} \text{ period} & 3d\text{- series } (\text{Sc - Zn }) \\ \hline & 5^{th} \text{ period} & 4d\text{- series } (\text{Y - Cd }) \\ \hline & 6^{th} \text{ period} & 5d\text{- series } (\text{La - Hg }) \\ \hline & 7^{th} \text{ period} & 6d\text{- series } (\text{Ac - Uub }) \end{array}$								
	Periodic trends and chemical properties									

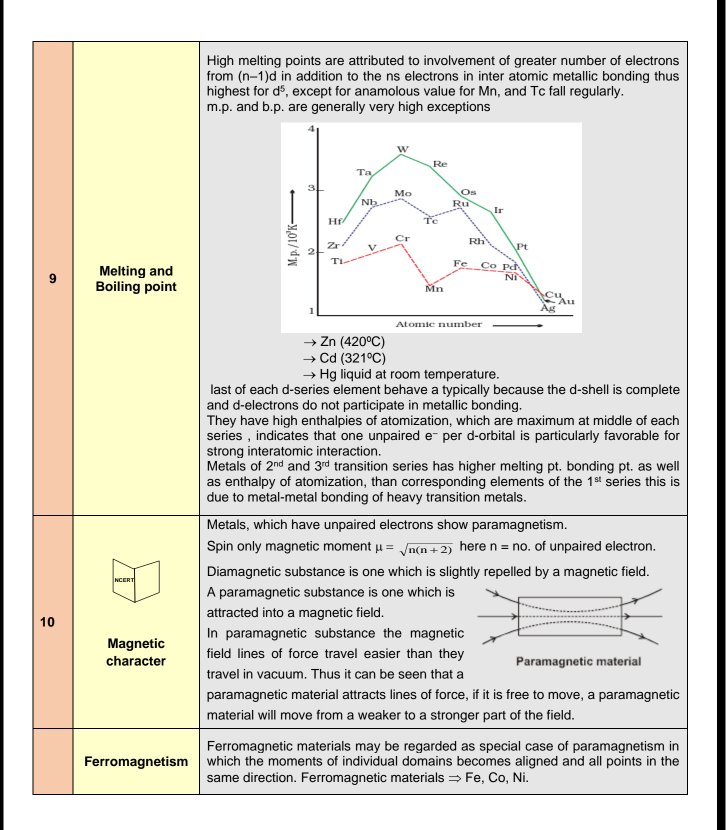
S.N.		Propert	ies		DISCRIPTION									
1		Metalli charact	ter	All the transition elements are metals, therefore they are showing all the metallicproperties. Good conductor of heat and electricity. Metallic lustre, strong and ductile. With the exception of Zn, Cd, Hg, Mn all the transition metals have one or more regular metallic structure. The transition metals are hard and have low volatility (except Zn, Cd and Hg).										
2	Electronic configuration			(n – 1)d ^{1 – 10} ns ^{1 or 2}										
			3 IIIB	4 IVB	5 VB	6 VIB	7 VIIB	8	9 VIII	10	11 IB	12 IIB		
			21	22	23	24	25	26	27	28	29	30		
		3d Series	Sc	Ti	v	Cr	Mn	Fe	Со	Ni	Cu	Zn		
			3d ¹ 4s ²	3d ² 4s ²	3d ³ 4s ²	3d ⁵ 4s ¹	3d ⁵ 4s ²	3d ⁶ 4s ²	3d ⁷ 4s ²	3d ⁸ 4s ²	3d ¹⁰ 4s ¹	3d ¹⁰ 4s ²		
			39	40	41	42	43	44	45	46	47	48		
		4d Series	Y	Zr	Nb	Мо	Тс	Ru	Rh	Pd	Ag	Cd		
			4d ¹ 5s ²	4d ² 5s ²	4d ⁴ 5s ¹	4d ⁵ 5s ¹	4d ⁵ 5s ²	4d ⁷ 5s ¹	4d ⁸ 5s ¹	4d ¹⁰	4d ¹⁰ 4s ¹	4d ¹⁰ 4s ²		
			57	72	73	74	75	76	77	78	79	80		
		5d Series		Hf	Та	w	Re	Os	Ir	Pt	Au	Hg		
			5d ¹ 6s ²	4f ¹⁴ 5d ² 6s ²	5d ³ 6s ²	5d ⁴ 6s ²	5d ⁵ 6s ²	5d ⁶ 6s ²	5d ⁷ 6s ²	5d ⁹ 6s ¹	5d ¹⁰ 6s ¹	5d ¹⁰ 6s ²		
			89	104	105	106	107	108	109	110	111	112		
		6d Series	Ac**	Rf	Db	Sg	Bh	Hs	Mt	Ds	Rg	Cn		
			6d ¹ 7s ¹											
									lls, thus ses, e.g.		ectronic d Cu.	configura	tion	
					r and Cu attain extra stability of half-filled and full-filled electronic configuration y changing the expected pattern of electronic configuration.									
					ctronic st , e.g. Gr					lways fo	llow the p	attern of	the	
							Ni Pd	3d ⁸ 4						
							Pu	5d ⁹ (
				d-orbita	ls are co	mplete a				(0.14)	2)			
						Ni (3d Pd(4d	,	Cu(3d ¹⁰ / Ag(4d ¹⁰ 5		n <u>(3d¹º4s²</u> d(4d¹º5s				
				Even th	ough gro		/		/	•	/	and coin	age	
				Even though ground state of the atom has a d ¹⁰ configuration Pd and coina metals behaves as typical transition elements, as in their most common oxidati states they have incomplete d-orbital.										
				A transition element is defined as the one which has incompletely filled d orbitals									itals	
				in its ground state or in any one of its oxidation states. Zinc, Cadmium and Mercury of group 12 have full <i>d</i> ¹⁰ configuration in their gro									und	
		Transiti		state as well as in their common oxidation states and hence, they are som										
		elemen	115	not rega	arded as	transitior	n metals.				·			
			All transition elements are d-block elements but all d-block elements are not transition elements. (as per NCERT)									not		
	c	ize of at	ome						sition se	ries), the	e covaler	t radii of	the	
3	3	and ior		element	s decrea	se from					he end w			
				increase	es slightl	у.								





					stable. Most c Highes by Ru The hig oxide a elemen +7 Ex. Mu Lower ligands bondin They fo	Ex. Sc ⁺ ommon (44) and ghest ox and oxyf nts and g $^{+6}$ nO ₄ , Cr ₂ oxidation s such as g Ni(CO	3^3 , Ti ⁺⁴ , V ⁺ constates Os (76). idation s luoride b lood oxid O_7^{-2} , Cr C n states s carbon i) ₄ and Fe c compou	$^{-5}$ Fe ⁺³ , M state an shown by tates (+4 because lants. $^{2^{-}}_{4}$, Mn O ₂ even neg monoxide e(CO) ₅ , o	In ⁺² , Zn ⁺² nong the r transitio 4, +5, +6 fluorine a $_{2}^{+6}$, MnO ₄ ²⁻ gative (-2 e and dip xidation	² etc. transition on element 5, +7, +8 and $0xyg$ $, Cr O_2F_2$ 2, -1, 0, - byridine c state of N	n elemer nts of '4d) are fou gen are r +1) is sta lue to ph Ni and Fe	ts is +2. and '5d' nd in co nost ele bilised b enomena is zero.	d ¹⁰ are mor series is + mpounds o ctronegativ y π-bondin a of synergi	-8 of /e ng ic
					Form	ulas of (Dxides o	f 3d Met	als (* mi	xed oxic	les)			
		XIDATIOI NUMBER		Sc	Ti	V	Cr	Mn	Fe	Co		Cu	Zn	
		+7						Mn ₂ O ₇						
		+6					CrO ₃							
		+5				V ₂ O ₅								
		+4	TiO2 V2O4 CrO2 MnO2 Image: Contract of the second s											
		+3	SC	2 O 3	Ti ₂ O ₃	V ₂ O ₃	Cr ₂ O ₃	Mn ₂ O ₃ Mn ₃ O ₄ *	Fe ₂ O ₃ Fe ₃ O ₄ ³) ₄ *			
		+2			TiO	VO	(CrO)	MnO	FeO	CoO	NiC	CuO Cu2O	ZnO	
6		Colou	ır	- 	They for an unpa lower ei excitatio MnO4 ⁻ is	m differe ired elec nergy d- n lies in coloure	tron that orbital is visible ra	red comp compou exited t inge, con of d ⁰ cor	oounds. T nd may l to a high nplimenta nfiguratic	The comp be colour her energe ary colour on of Mn.	oounds ir red. Whe gy d-orb ir is obse It is not	which m n an ele ital, if th rved.	ured. netal ion ha ctron from e energy o l-d transitio	a of
	Me	mory wa	rnina ·								u O.			
		Oxidation State			Ti	V	Cr	Mn	Fe	Со	Ni	Cu	Zn	
	(I) Colurless													
		(II) Blue Pink Green Pink Green Blue Colurless												
	(III) Colurless Purple Green Green Violet Yellow Blue													
	(IV) Colurless Blue													
Sect	ection (B) : Electrode potential and chemical reactivity, Magnetic properties,													
	formation of coloured ions, Catalytic properties, formation of interstitial compounds, alloy formation.													
		ompou ectrode	nus,	all	by for	nation								
7		tential												
		$E_{M^{2^+}/M}^{0}$ There is no regular trend in these values. This is attributed to the irregular variation of ionisation enthalpies (IE ₁ + IE ₂), hydration energies and the sublimation energies in the period.												

		Memory warning : Ne	ed not memorize	the table but remember the comments.						
		Element (M) $E_{M^{3+}/I}^{0}$	H ²⁺ E ⁰ _{M²⁺/M}	Comments						
		Ti -0.3	7 –1.63	+3 Oxidation states is more stable						
		V -0.2	.6 –1.18	+3 Oxidation states is more stable						
		Cr -0.4	.1 –0.9	+3 Oxidation states is more stable						
		Mn 1.57	7 –1.18	+2 Oxidation states is more stable						
		Fe 0.77	7 –0.44	+2 Oxidation states is more stable						
		Co 1.97	7 –0.28	+2 Oxidation states is more stable						
		Ni –	-0.25	+3 Oxidation states does not exist for Ni						
		Cu –	0.34	+3 Oxidation states does not exist for Cu. 0 Oxidation states is more stable						
		Zn –	0.76	+3 Oxidation states does not exist for Zn						
	Comparative graph of $E_{M^{2+}/M}^{0}$ and $E_{M^{3+}/M^{2+}}^{0}$	configuration in Zn ²⁺ a to the highest negativ It may be noted that th comparison to element have fairly large ionisa	If-filled <i>d</i> sub-she are related to their the electrode potents of group 2 (e.g	entials of transition metals are low in g., Ca = -2.87 V, the transition elements and very large enthalpies of atomisation).						
		their hydration enthal	Compared to group 2 elements. These reduce their electrode potentials though their hydration enthalpies are large.							
8	Density	-	Elements with highest densities are Osmium (Os) = 22.51 g/cm ³ , Iridium (Ir) = 22.61 g/cm ³							



			and their compounds are known for their catalytic activity. to adopt multiple oxidation states and to form complexes.					
		Catalyst	Used					
		Fe	Haber's process for manufacture of NH ₃ .					
		V ₂ O ₅	Contact process for H ₂ SO ₄ manufacture.					
	NCERT	Pt	Ostwald's process of nitric acid.					
11	NCENT	Ni	Hydrogenation of oils.					
	Catalytic Properties	FeSO4 & H2O2	Fenton's reagent for oxidising alcohol to aldehyde.					
		Cu	Dehydrogenation of alcohols					
		Al ₂ (C ₂ H ₅) ₆ + TiCl ₄	Ziegler-Natta catalyst in the production of Polyethene.					
		FeCl₃	For making CCl ₄ from CS ₂ and Cl ₂ .					
		Pt/PtO Adams catalyst, used for reduction.						
12	FORMATION OF INTERSTITIAL COMPOUNDS	N are trapped insid stoichiometric and are Fe ₃ H, VH _{0.56} and TiH to any normal oxida composition, these of principal physical are follows: (i) They have high me	•					
13	FORMATION OF ALLOYS	 (i) Transition elements have maximum tendency to form alloys. (ii) The reactivity of transition elements is very less and their sizes are almost similar. Due to this a transition metal atom in the lattice can be easily replaced be other transition metal atom and hence they have maximum tendency to form alloys. (iii) In the alloys, ratio of component metals is fixed. (iv) These are extremely hard and have high melting point. 						

Λ	Note- Men	norize the * marked al	loys
	*(a)	Bronze	Cu (75 - 90 %) +Sn (10 - 25 %)
	*(b)	Brass	Cu (60 - 80 %) +Zn (20 - 40 %)
	(c)	Gun metal	(Cu + Zn + Sn) (87 : 3 : 10)
	(d)	German Silver	Cu + Zn + Ni (2 : 1 : 1)
	(e)	Bell metal	Cu (80 %) + Sn(20 %)
	(f)	Nichrome	(Ni + Cr + Fe)
	(g)	Alnico	(Al, Ni,Co)
	(h)	Type Metal	Pb + Sn + Sb
	(i)	Alloys of steel	
		• Vanadium steel	V (0.2 - 1 %)
		Chromium steel	Cr (2 - 4 %)
		Nickel steel	Ni (3 -5 %)
		• Manganese steel	Mn (10 -18 %)
SOME IMPORTANT		Stainless steel	Cr (12 - 14 %) & Ni (2 - 4 %)
ALLOY		Tunguston steel	W (10 - 20 %)
	(j)	14 Carat Gold	54 % Au + Ag (14 to 30 %) + Cu (12 - 28 %)
	(k)	24 Carat Gold	100 %Au
	*(I)	Solder	Pb + Sn
	(m)	Magnellium	Mg (10%) + AI (90%)
	(n)	Duralumin	(Al + Mn + Cu)
	*(0)	Artificial Gold	Cu (90 %) + Al (10%)
	(p)	Constantan	Cu(60%) + Ni (40%)
		% of Carl	bon in different type of Iron
		Name	% of C
	(a)	Wrought Iron	0.1 to 0.25
	(b)	Steel	0.25 to 2.0
	(c)	Cast Iron/Pig Iron	2.6 to 4.3

Section (C) : Important d-block metal compounds.

Preparations and properties of some important d-Block metal compounds

Sulphates (SO₄^{2–})

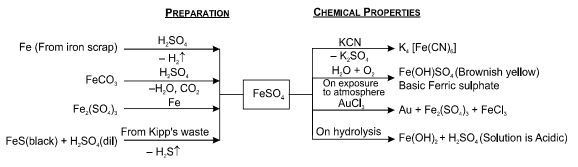
1. FeSO₄.7H₂O (Green Vitriol)

Ferrous sulphate.

Commonly known as harkasis.

Physical properties

- Hydrated ferrous sulphate is a green coloured compound.
- Effloresces on exposure to air.
- Anhydrous FeSO₄ is colourless.



Heating effect :

 $\mathsf{FeSO}_4.7\mathsf{H}_2\mathsf{O} \xrightarrow{140^{\circ}\mathsf{C}} \mathsf{FeSO}_4.\mathsf{H}_2\mathsf{O} \xrightarrow{300^{\circ}\mathsf{C}} 2\mathsf{FeSO}_4 \xrightarrow{\mathsf{High}} \mathsf{Fe}_2\mathsf{O}_3 + \mathsf{SO}_2 \uparrow + \mathsf{SO}_3 \uparrow$

Uses : It is used

- (i) for making Blue Black ink.
- (ii) as mordant in dyeing.
- (iii) as insecticide in agriculture.
- (iv) for making laboratory reagents like Mohr's salt etc.

$$FeSO_4 + (NH_4)_2 SO_4 + 6H_2O) \longrightarrow FeSO_4 (NH_4)_2SO_4.6H_2O (Mohr's salt)$$

(v) $FeSO_4 + H_2O_2$ known as Fenton's reagent is used as catalyst.

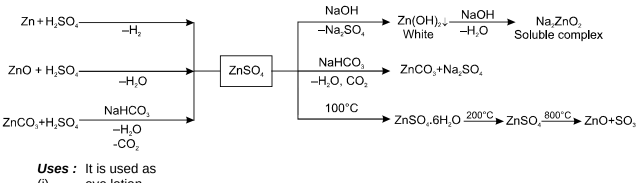
2. Zinc Sulphate, ZnSO₄. 7H₂O (white Vitriol)

Physical Properties

- Colourless, crystalline solid, soluble in water.
- It slowly effloresces when exposed to air.
- It is isomorphos with Epsom salt (MgSO₄.7H₂O).

PREPARATION

CHEMICAL PROPERTIES

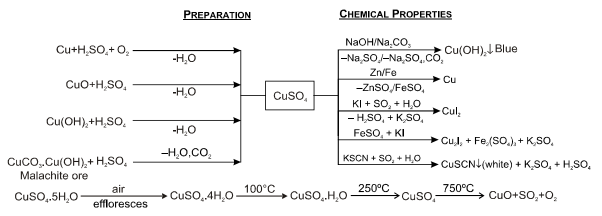


- (i) eye lotion.
- (ii) for making lithophone-mixture of BaS + ZnSO₄ (while paint).
- (iii) mordant in dyeing.

3. Copper sulphate, CuSO₄.5H₂O (blue vitriol)

Also called as 'Nilathotha'

Physical Properties : Blue crystalline compound soluble in water.



Uses : It is used

- (i) for making other copper compounds.
- (ii) for electroplating, electrotyping, as mordant in dyeing.
- (iii) in making Bordeaux mixture which is used in agriculture as fungicide and germicide.
- in making Fehlings solution. (iv)
- in medicine as antiseptic. (v)
- in electric batteries. (vi)

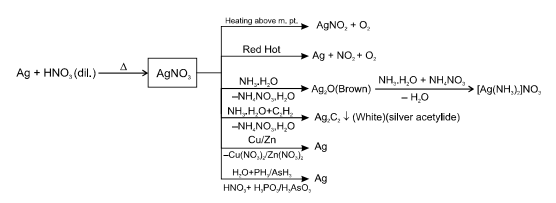
4. Silver nitrate, AgNO₃ (Lunar Caustic) **Physical Properties**

- (i) It is a colourless crystalline compound.
- (ii) Soluble in water and alcohol.
- (iii) It melts at 212°C.

Chemical Properties

- (i) It possesses powerful corrosive action on organic tissues, which at turns black especially in presence of light. The blackening is due to finely divided metalic silver, reduced by organic tissure It is therefore, stored in colored Bottles.
- (ii) Solutions of halides phosphates, sulfides chromates thiocyanates, sulphates and thiosulphates salt with silver nitrate solution.
- (iii) Ammonical silver nitrate is called as Tollen's reagent and used to identify reducing sugars and aldehvdes. PREPARATION

CHEMICAL PROPERTIES



It is known as silver mirror test of aldehydes and reducing sugars.

 $Ag_2O + HCHO \longrightarrow 2Ag + HCOOH.$

 $Ag_2O + C_6H_{12}O_6 \longrightarrow 2Ag + C_6H_{12}O_7.$

(iv) <u>Reactions with lodine :</u>

 $\begin{array}{l} 6 \ \mbox{AgNO}_3 \ (excess) + 3I_2 + 3H_2O \longrightarrow \ \mbox{AgIO}_3 + 5AgI + 6HNO_3 \\ 5 \ \mbox{AgNO}_3 + 3I_2(excess) + 3H_2O \longrightarrow \ \mbox{HIO}_3 \ + 5AgI + 5HNO_3 \end{array}$

Uses : It is used

- (i) as a laboratory reagent for the identification of various acidic especially for Cl, Br and I.
- (ii) Tollen's reagent is used in organic chemistry for testing aldehydes reducing sugars etc.
- (iii) for making AgBr, used in photography.
- (iv) in the preparation of inks and hair dyes.
- (v) in preparation of silver mirror.

5. Photography :

(i) A photographic film consists of a light sensitive emulsion of fine particles (grains) of silver salts in gelatine spread on a clear celluloid strip or a glass plate. AgBr is mainly used as the light sensitive material.

(ii) The film is placed in a camera. When the photograph is exposed, light from the subject enters the camera and is focussed by the lens to give a sharp image on the film. The light starts a photochemical reaction by exciting a halide ion, which loses an electron. The electron moves in a conduction band to the surface of the grain, where it reduces a Ag⁺ ion to metallic silver.

$2AgBr(s) \xrightarrow{light} 2Ag + Br_2$

(iii) In modern photography only a short exposure of perhaps 1/100th of a second is used. In this short time, only a few atoms of silver (perhaps 10–50) are produced in each grain exposed to light. Parts of the film which have been exposed to the bright parts of the subject contain a lot of grains with some silver.

(iv) Next the film is placed a developer solution. This is a mild reducing agent, usually containing quinol. Its purpose is to reduce more silver halide to Ag metal. Ag is deposited mainly where there are already some Ag atoms. Thus the developing process intensifies the latent image on the film so it becomes visible.

$$2AgBr(s) + 2OH^{-}(aq) + HO \longrightarrow OH (aq) \longrightarrow 2Ag(s) + 2H_2O(\Box) + O \implies O + 2Br^{-}(aq)$$

hydroquinol hydroquinone

(v) If the film was brought out into daylight at this stage, the unexposed parts of the emulsion would turn black and thus destroy the picture. To prevent this happening any unchanged silver halides are removed by placing the film in a fixer solution. A solution of sodium thiosulphate is used as fixer. It forms a soluble complex with silver halides.

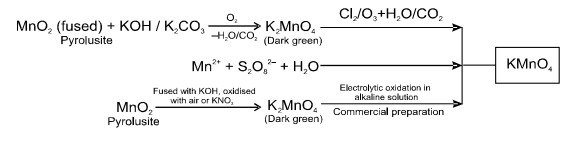
AgBr + $2Na_2S_2O_3 \rightarrow Na_3[Ag(S_2O_3)_2] + NaBr$

After fixing, the film can safely be brought out into daylight. This is called "negative".

Light is passed through the negative onto a piece of paper coated with AgBr emulsion. This is then developed and fixed in the same way as before.

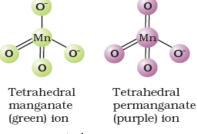
6. Potassium Permanganatic (KMnO₄)

PREPARATION



Physical Properties :

Purple coloured crystalline compound. Moderately soluble in water at room temperature.



Structure of manganate and permanganate ion.

Chemical Properties

(i) Heating effect

(ii) Reactions with MnO₄²⁻ in dil. alkaline, water and acidic solutions

$$O_{2} + H_{2}O + MnO_{4}^{-} \xrightarrow{\Lambda} MnO_{4}^{2-} \xrightarrow{H^{+}} 2MnO_{4}^{-} + MnO_{2} \downarrow + H_{2}O \xrightarrow{H_{2}O} 2MnO_{4}^{-} + MnO_{2} \downarrow + ClO4^{-} \xrightarrow{H_{2}O} 2MnO_{4}^{-} + MnO_{2} \downarrow + ClO4^{-} \xrightarrow{H_{2}O} 2MnO_{4}^{-} + MnO_{2} \downarrow + OlO4^{-} \xrightarrow{H_{2}O} X_{2} \downarrow + OlO4^{-} X_{2}$$

 $OH^- + MnO + 2H_2O$

(iii) with conc. H₂SO₄,

$$3H_2SO_4 \xrightarrow{2KMnO_4} K^+ + MnO_3^+ + 3HSO_4^- + H_3O^+$$
(green)
$$2MnO_2 + \frac{3}{2}O_2$$

$$Mn_2O_7 + H_2SO_4$$

$$Mn_2O_7 + H_2SO_4$$

(iv) KMnO₄ is a powerful oxidising agent

Potassium permanganate acts as an oxidising agent in alkaline, neutral or acidic solutions. A mixture of sulphur, charcoal and KMnO₄ forms an explosive powder. A mixture of oxalic acid and KMnO₄ catches fire spontaneous after a few seconds. The same thing happens when glycerine is poured over powdered KMnO₄

In alkaline & neutral medium :

In strongly alkaline medium KMnO4 is reduced to manganate.

 $2KMnO_4 + 2KOH (conc.) \longrightarrow 2K_2 MnO_4 + H_2O + [O]$

or $e^- + MnO_4^- \longrightarrow MnO_4^{2-}$ However if solution is dilute then K₂MnO₄ is converted in to MnO₂ which appears as a brownish precipitate. $2K_2MnO_4 + 2H_2O \longrightarrow 2MnO_2 + 4KOH + 2[O]$

or
$$2e^- + 2H_2O + MnO_4^{2-} \longrightarrow MnO_2 + 4OH^-$$

This type of behaviour is shown by KMnO₄ itself in neutral medium.

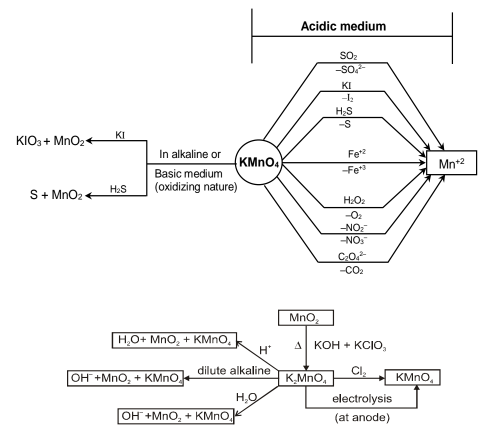
 $3e^{-} + 2H_2O + MnO_4^{-} \longrightarrow MnO_2 + 4OH^{-}$

$$3e^{-} + 2H_2O + MnO_4^{-} \xrightarrow[dilute]{} MnO_2 \\ brown \ ppt \\ \downarrow + H_2O \xleftarrow{-2[O]}{} 2MnO_4^{2-} + 2H_2O$$

Note : $MnO_4^- + 8H^+ + 5e^- \rightarrow Mn^{+2} + 4H_2O$

This medium is used in quantitative (Volumetric) Estimations. The eq. mass of KMnO₄ in acidic medium is = $\frac{\text{Molecular Mass}}{5}$

Reactions of KMnO₄:



Uses : It is used

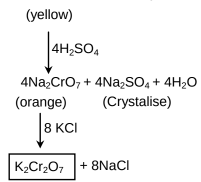
- (i) KMnO₄ is used as an oxidising agent in laboratory and industry.
- (ii) Alkaline potassium permanganate is called Bayer's reagent. This reagent is used in organic chemistry for the test of unsaturation. KMnO₄ is used in the manufacture of saccharin, benzoic acid, acetaldehyde etc.
- (iii) KMnO₄ is used in qualitative analysis for detecting halides, sulphites, oxalates, etc.

7. POTASSIUM DICHROMATE (K₂Cr₂O₇) :

Preparation :

The chromite ore is roasted with sodium carbonate in presence of air in a reverberatory furnace.

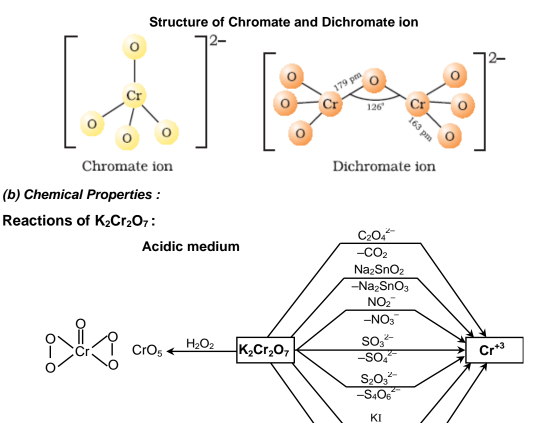
 $4FeO.Cr_2O_3 + 8Na_2CO_3 \xrightarrow{\text{Roasting in air}} 8Na_2CrO_4 + 2Fe_2O_3 + 8CO_2$



Properties

(a) Physical :

It is orange-red coloured crystalline compound. It is moderately soluble in cold water but freely soluble in hot water. It melts at 398°C.



Note- $K_2Cr_2O_7$ is preferred over Na₂Cr₂O₇ as a primary standard in volumetric estimation because Na₂Cr₂O₇ is hygroscopic in nature but $K_2Cr_2O_7$ is not.

-I₂ Fe⁺² -Fe⁺³

(i) Effect of heating : On heating strongly, it decomposes liberating oxygen.

$$\begin{array}{c} \begin{array}{c} Cr_2O_7^{-2} & \Delta \end{array} \\ \xrightarrow{(Orange)} & 2K_2CrO_4 + Cr_2O_3 + \frac{3}{2}O_2 \\ \end{array} \\ \xrightarrow{(Orange)} & 2KOH \\ & 2CrO_4^{2-} + H_2O \\ & (yellow) \end{array} \\ \begin{array}{c} 2CrO_4^{2-} + 2H^+ \overleftrightarrow{(V_2O_4)} & 2HCrO_4^{-} \end{array} \\ \end{array}$$

 $CrO_{4^{2-}}$ and $Cr_2O_{7^{2-}}$ exist in equilibrium and are interconvertable by altering the pH of solution. In alkaline solution, chromate ions are present while in acidic solution, dichromate ions are present.

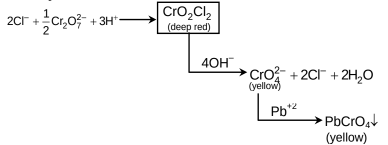
Chrome alum preparation \rightarrow

 $\begin{array}{l} K_2Cr_2O_7 \ + \ H_2SO_4 + \ 3SO_2 + \ 23H_2O & \xrightarrow{T < 70^\circ C} \\ \text{Oxidizes ethyl alcohol to acetaldehyde to acetic acid} \end{array} \\ \end{array} \\ \begin{array}{l} K_2SO_4.Cr_2(SO_4)_{3.24H_2O} \\ \text{Oxidizes of the set of$

$$\rightarrow$$
 C₂H₅OH—^[O] \rightarrow CH₂CHO—^[O] \rightarrow CH₂COOH

- \rightarrow It also oxidizes nitrites to nitrates, arsenates to arsenates, HBr to Br₂, HI to I₂ etc.
- \rightarrow K₂Cr₂O₇ + 2C(Charcoal) \longrightarrow Cr₂O₃ + K₂CO₃ + CO

Chromyl chloride test :



Reaction of potassium dichromate with Ag+ -

$$2BaCrO_{4}\downarrow + 2H^{+} \xleftarrow{Ba^{2+} + H_{2}O}_{Partial \ precipitation} \underbrace{Cr_{2}O_{7}^{2-}}_{(conc.)} \underbrace{2Ag^{+}}_{(reddish \ brown)} Ag_{2}Cr_{2}O_{7}\downarrow \\ (reddish \ brown) \underbrace{H_{2}O}_{boil} Ag_{2}CrO_{4}\downarrow + CrO_{4}^{2-} + 2H^{+}$$

 $Cr_2O_7^{2-} + Ba^{2+} + H_2O \implies 2BaCrO_4 \downarrow + 2H^+$

As strong acid is produced, the precipitation is only partial. But if NaOH or CH₃COONa is added, precipitate becomes quantitative.

Uses: It is used :

- (i) as a volumetric reagent in the estimation of reducing agents such as oxalic acid, ferrous ions, iodide ions, etc. It is used as a primary standard.
- (ii) for the preparation of several chromium compounds such as chrome alum, chrome yellow, chrome red, zinc yellow, etc.
- (iii) in dyeing, chrome tanning, calico printing, photography etc.
- (iv) as a cleansing agent for glass ware in the form of chromic acid.
- (iv) in leather industry and as an oxidant for preparation of azo compounds.

Section (D) : Lanthanoids and actinoids

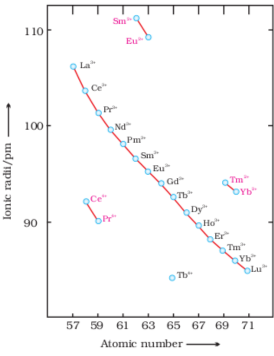
	f-block elements (JEE-Mains only)									
	Differentiating electrons enters in (n – 2)f subshell. f-block elements lie on the Bottom portion of periodic table.									
S.No.	Properties		DISCRIPTION							
1.	General character's	It shows high melting and	Il the f-block elements are heavy metals. shows high melting and boiling point. he most common oxidation state of these elements is +3.							
2.	No. of elements	Total number of f-block e	lements -	- (28)						
3.	Groups		Sc Y La Lanthanides (14) Ce ₅₈ – Lu ₇₁							
4.	E.C.	Lanthanide series 4f ¹⁻¹⁴ Actinide series 5f ¹⁻¹⁴ 6d ⁰		2						
5.	Period	Period III 6 th period 7 th period	B/ 3rd Sc Y La Ac	Lanthanides (14) Ce ₅₈ – Lu ₇₁ Actinides (14) Th ₉₀ – Lr ₁₀₃						
6.	Inner transition elements	and pre or antipenultimat elements.Ce ₅₈ = [Xe] 6s ²	e (n – 2) : , 5d¹, 4f¹	e shells that is ultimate (n) penultimate $(n-1)$ shell are incomplete are called inner transition ed into two series.						
	(i) Lanthanide series or Rare earth elements or Lanthenones	Ce ₅₈ – Lu ₇₁ 14 elements Lanthanides are found ra The first element of this s In these elements, last el They are present in IIIB g	Lanthanides are found rarely on earth so these are called rare earth metals. The first element of this series is Cerium and not Lanthanum. In these elements, last electron enters into 4f subshell. They are present in IIIB group and 6 th period of the periodic table. Promethium (₆₁ Pm) is the only lanthanide which is synthetic and radioactive in							
	(ii) Actinide series or Man made elements or Actinones	 Th₉₀ – Lr₁₀₃14 elements. All the actinides are radioactive elements. The first element of this series is Thorium and not Actinium. In these elements, last electron enters into 5f subshell. They are present in IIIB group and 7th period of the periodic table. All the actinides are radioactive in nature. First three elements (Th, Pa, U) are found in nature while others are synthetic in nature. Transuranic actinides are man-made elements (Np₉₃ – Lw₁₀₃) After U₉₂ i.e. from Np₉₃ onwards elements are called transuranic elements because (i) They are heavier than uranium. (ii) They are derived from uranium by nuclear reactions. 								

The Lanthanides : The names, symbols, electronic configurations of atomic and some ionic states and atomic and ionic radii of lanthanum and lanthanide (for which the general symbol Ln is used) are given in Table.

Electronic Configurations : It may be noted that atoms of these elements have electronic configuration with $6s^2$ common but with variable occupancy of 4f level (Table). However, the electronic configurations of all the tripositive ions (the most stable oxidation state of all the lanthanides) are of the form $4f^n$ (n = 1

to 14 with increasing atomic number). Atomic and Ionic Sizes : The overall decrease in atomic and ionic radii from lanthanum to lutetium (the lanthanide contraction). The shielding of one 4 f electron by another is less than one d electron by another with the increase in nuclear charge along the series. There is fairly regular decrease in the sizes with increasing atomic number. The cumulative effect of the contraction of the lanthanide series, known as lanthanide contraction, causes the radii of the members of the third transition series to be very similar to those of the corresponding members of the second series. The almost identical radii of Zr (160 pm) and Hf (159 pm), a consequence of the lanthanide contraction. **Oxidation States :** In the lanthanides, La(III) and Ln(III) compounds are predominant species. However, occasionally +2 and +4 ions in solution or in solid compounds are also obtained.

This irregularity (as in ionisation enthalpies) arises mainly from the extra stability of empty, half-filled or filled f subshell. Thus, the formation of Ce^{IV} is favoured by its noble gas configuration, but it is a strong oxidant reverting to the common +3 state. The E^{\odot} value for Ce⁴⁺/Ce³⁺ is + 1.74 V which suggests that it can oxidise water.



However, the reaction rate is very slow and hence Ce(IV) is a good analytical reagent.

Pr, Nd, Tb and Dy also exhibit +4 state but only in oxides, MO_2 . Eu^{2+} is formed by losing the two s electrons and its f⁷ configuration accounts for the formation of this ion.

However, Eu²⁺ is a strong reducing agent changing to the common +3 state. Similarly Yb²⁺ which has f 14 configuration is a reductant.

Tb^{IV} has half-filled f-orbitals and is an oxidant. The behaviour of samarium is very much like europium, exhibiting both +2 and +3 oxidation states.

	Table :- Electronic Configurations and Radii of Lanthanum and Lanthanides								
Atomic	Nome	aymhal	Ele	ectronic co	ns*	Radii/pm			
Number	Name	symbol	Ln	Ln ²⁺	Ln ³⁺	Ln ⁴⁺	Ln	Ln ³⁺	
57	Lanthanum	La	5d ¹ 6s ²	5d ¹	4f ⁰		187	106	
58	Cerium	Ce	4f ¹ 5d ¹ 6s ²	4f ²	4f ¹	4f ⁰	183	103	
59	Praseodymium	Pr	4f ³ 6s ²	4f ³	4f ²	4f ¹	182	101	
60	Neodymium	Nd	4f ⁴ 6s ²	4f ⁴	4f ³	4f ²	181	99	
61	Promethium	Pm	4f ⁵ 6s ²	4f ⁵	4f ⁴		181	98	
62	Samarium	Sm	4f ⁶ 6s ²	4f ⁶	4f ⁵		180	96	
63	Europium	Eu	4f ⁷ 6f ²	4f ⁷	4f ⁶		199	95	
64	Gadolinium	Gd	4f ⁷ 5d ¹ 6s ²	4f ⁷ 5d ¹	4f ⁷		180	94	
65	Terbium	Tb	4f ⁹ 6s ²	4f ⁹	4f ⁸	4f ⁷	178	92	
66	Dysprosium	Dy	4f ¹⁰ 6s ²	4f ¹⁰	4f ⁹	4f ⁸	177	91	
67	Holmium	Ho	4f ¹¹ 6s ²	4f ¹¹	4f ¹⁰		176	89	
68	Erbium	Er	4f ¹² 6s ²	4f ¹²	4f ¹¹		175	88	
69	Thulium	Tm	4f ¹³ 6s ²	4f ¹³	4f ¹²		174	87	
70	Ytterbium	Yb	4f ¹⁴ 6s ²	4f ¹⁴	4f ¹³		173	86	
71	Lutetium	Lu	4f ¹⁴ 5d ¹ 6s ²	4f ¹⁴ 5d ¹	4 f ¹⁴	-	-	-	

General Characteristics :

All the lanthanides are silvery white soft metals and tarnish rapidly in air.

The hardness increases with increasing atomic number, samarium being steel hard.

Their melting points range between 1000 to 1200 K but samarium melts at 1623 K.

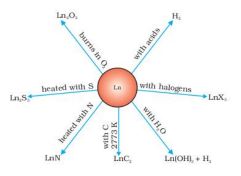
They have typical metallic structure and are good conductors of heat and electricity. Density and other properties change smoothly except for Eu and Yb and occasionally for Sm and Tm.

Many trivalent lanthanide ions are coloured both in the solid state and in aqueous solutions. Colour of these ions may be attributed to the presence of f electrons. Neither La³⁺ nor Lu³⁺ ion shows any colour but the rest do so. However, absorption bands are narrow, probably because of the excitation within f level.

The lanthanide ions other than the f° type (La³⁺ and Ce⁴⁺) and the f^{14} type (Yb²⁺ and Lu³⁺) are all paramagnetic. The paramagnetism rises to maximum in neodymium.

The first ionisation enthalpies of the lanthanides are around 600 kJ mol^{-1} , the second about 1200 kJ mol^{-1} comparable with those of calcium.

A detailed discussion of the variation of the third ionisation enthalpies indicates that the exchange enthalpy consideration (as in 3d orbitals of the first transition series), appear to impart a certain degree of stability to empty, half-filled and completely filled orbitals f level. This is indicated from the abnormally low value of the third ionization enthalpy of lanthanum, gadolinium and lutetium.



In their chemical behaviour, in general, the earlier members of the series are quite reactive similar to calcium but, with increasing atomic number, they behave more like aluminium.

Values for E[−] for the half-reaction:

 Ln^{3+} (aq) + $3e^- \rightarrow Ln(s)$ are in the range of -2.2 to -2.4 V except for Eu for which the value is - 2.0 V. This is, of course, a small variation.

The metals combine with hydrogen when gently heated in the gas.

The carbides, Ln_3C , Ln_2C_3 and LnC_2 are formed when the metals are heated with carbon.

They liberate hydrogen from dilute acids and burn in halogens to form halides.

They form oxides M₂O₃ and hydroxides M (OH)₃. The hydroxides are definite compounds, not just hydrated oxides.

They are basic like alkaline earth metal oxides and hydroxides.

The best single use of the lanthanides is for the production of alloy steels for plates and pipes. A well known alloy is mischmetall which consists of a lanthanide metal (~ 95%) and iron (~ 5%) and traces of S, C, Ca and AI. A good deal of mischmetall is used in Mg-based alloy to produce bullets, shell and lighter flint. Mixed oxides of lanthanides are employed as catalysts in petroleum cracking. Some individual Ln oxides are used as phosphors in television screens and similar fluorescing surfaces.

The Actinides :

The actinides include the fourteen elements from Th to Lr. The names, symbols and some properties of these elements are given in Table.

	Table : Electronic Configurations and Radii of Actinium and Actinoids									
Atomic	Name	overhol	Electron	ic configu	rations*	Radii/pm				
Number		symbol	М	M ³⁺	M ⁴⁺	M ³⁺	M ⁴⁺			
89	Actinium	Ac	6d ¹ 7s ²	5f ⁰		111				
90	Thorium	Th	6d ² 7s ²	5f ¹	5f ⁰		99			
91	Proctactiniium	Ра	5f ² 6d ¹ 7s ²	5f ²	5f ¹		96			
92	Uranium	U	5f ³ 6d ¹ 7s ²	5f ³	5f ²	103	93			
93	Neptunium	Np	5f ⁴ 6d ¹ 7s ²	5f ⁴	5f ³	101	92			
94	Plutonium	Pu	5f ⁶ 7s ²	5f⁵	5f ⁴	100	90			
95	Americium	Am	5f ⁷ 7s ²	5f ⁶	5f⁵	99	89			
96	Curium	Cm	5f ⁷ 6d ¹ 7s ²	5f ⁷	5f ⁷	99	88			
97	Berkelium	Bk	5f ⁹ 7s ²	5f ⁸	5f ⁷	98	87			
98	Californium	Cf	5f ¹⁰ 7s ²	5f ⁹	5f ⁸	98	86			
99	Einstenium	Es	5f ¹¹ 7s ²	5f ¹⁰	5f ⁹	-	-			
100	Fermium	Fm	5f ¹² 7s ²	5f ¹¹	5f ¹⁰	-	-			
101	Mendelevium	Md	5f ¹³ 7s ²	5f ¹²	5f ¹¹	-	-			
102	Nobelium	No	5f ¹⁴ 7s ²	5f ¹³	5f ¹²	-	-			
103	Lawrencium	Lr	5f ¹⁴ 6s ¹ 7s ²	5f ¹⁴	5f ¹³	-	-			

The actinides are radioactive elements and the earlier members have relatively long half-lives, the latter ones have half-life values ranging from a day to 3 minutes for lawrencium (Z = 103). The latter members could be prepared only in nanogram quantities. These facts render their study more difficult.

Electronic Configurations :

All the actinides are believed to have the electronic configuration of 7s² and variable occupancy of the 5f and 6d subshells.

The fourteen electrons are formally added to 5f, though not in thorium (Z = 90) but from Pa onwards the 5f orbitals are complete at element 103.

The irregularities in the electronic configurations of the actinides, like those in the lanthanides are related to the stabilities of the f° , f^{7} and f^{14} occupancies of the 5f orbitals. Thus, the configurations of Am and Cm are [Rn] $5f^{7}7s^{2}$ and [Rn] $5f^{7}$ 6d¹ 7s².

Ionic Sizes :

The general trend in lanthanides is observable in the actinides as well. There is a gradual decrease in the size of atoms or M^{3+} ions across the series. This may be referred to as the actinide contraction (like lanthanide contraction). The contraction is, however, greater from element to element in this series resulting from poor shielding by 5f electrons.

Oxidation States :

There is a greater range of oxidation states, which is in part attributed to the fact that the 5f, 6d and 7s levels are of comparable energies. The known oxidation states of actinides are listed in Table.

	Table : Oxidation States of Actinium and Actinides													
Ac	Ac Th Pa U Np Pu Am Cm Bk Cf Es Fm Md No Lr													
3		3	3	3	3	3	3	3	3	3	3	3	3	3
	4	4	4	4	4	4	4	4						
		5	5	5	5	5								
			6	6	6	6								
				7	7									

The actinides show in general +3 oxidation state.

The elements, in the first half of the series frequently exhibit higher oxidation states. For example, the maximum oxidation state increases from +4 in Th to +5, +6 and +7 respectively in Pa, U and Np but decreases in succeeding elements.

The actinides resemble the lanthanides in having more compounds in +3 state than in the +4 state. However, +3 and +4 ions tend to hydrolyse.

Because the distribution of oxidation states among the actinides is so uneven and so different for the earlier and latter elements, it is unsatisfactory to review their chemistry in terms of oxidation states.

General Characteristics and Comparison with Lanthanides :

The actinide metals are all silvery in appearance but display a variety of structures. The structural variability is obtained due to irregularities in metallic radii which are far greater than in lanthanides.

The actinides are highly reactive metals, especially when finely divided. The action of boiling water on them, for example, gives a mixture of oxide and hydride and combination with most non metals takes place at moderate temperatures. Hydrochloric acid attacks all metals but most are slightly affected by nitric acid owing to the formation of protective oxide layers; alkalies have no action.

The magnetic properties of the actinides are more complex than those of the lanthanides. Although the variation in the magnetic susceptibility of the actinides with the number of unpaired 5 f electrons is roughly parallel to the corresponding results for the lanthanides, the latter have higher values.

It is evident from the behaviour of the actinides that the ionisation enthalpies of the early actinides, though not accurately known, but are lower than for the early lanthanides. This is quite reasonable since it is to be expected that when 5f orbitals are beginning to be occupied, they will penetrate less into the inner core of electrons. The 5f electrons, will therefore, be more effectively shielded from the nuclear charge than the 4f electrons of the corresponding lanthanides. Because the outer electrons are less firmly held, they are available for bonding in the actinides.

A comparison of the actinides with the lanthanides, with respect to different characteristics as discussed above, reveals that behaviour similar to that of the lanthanides is not evident until the second half of the actinide series. However, even the early actinides resemble the lanthanides in showing close similarities with each other and in gradual variation in properties which do not entail change in oxidation state. The lanthanide and actinide contractions, have extended effects on the sizes, and therefore, the properties of the elements succeeding them in their respective periods. The lanthanide contraction is more important because the chemistry of elements succeeding the actinides are much less known at the present time.

Some Applications of d- and f-Block Elements :

Iron and steels are the most important construction materials. Their production is based on the reduction of iron oxides, the removal of impurities and the addition of carbon and alloying metals such as Cr, Mn and Ni.

TiO for the pigment industry and MnO₂ for use in dry battery cells. The battery industry also requires Zn and Ni/Cd.

The 'silver' UK coins are a Cu/Ni alloy.

MISCELLANEOUS SOLVED PROBLEMS (MSPs)

- 1. Among the following statements choose the true or false statement(s).
 - (a) $K_2Cr_2O_7$ on heating with charcoal gives metallic potassium and Cr_2O_3 .
 - (b) On heating in current of H₂ the crystalline KMnO₄ is converted into KOH and Mn₃O₄.

(c) Hydrated ferric chloride on treatment with 2, 2-dimethoxypropane gives anhydrous ferric chloride.

- Ans. (a
 - (a) False (b) False (c) True
- Sol. (a) $K_2Cr_2O_7 + 2C$ (charcoal) $\xrightarrow{\Delta} Cr_2O_3 + K_2CO_3 + CO^{\uparrow}$.

(b)
$$2KMnO_4 + 5H_2 \xrightarrow{\Delta} 2 KOH + 2MnO + 4H_2O$$

- (c) FeCl₃ . 6H₂O + 6CH₃— $\overset{OCH_3}{\overset{}{\underset{}}}$ $\xrightarrow{}$ FeCl₃ (anhydrous) + 12CH₃OH + 6CH₃COCH₃.
- 2. A compound (A) is used in paints instead of salts of lead. Compound (A) is obtained when a white compound (B) is strongly heated. Compound (B) is insoluble in water but dissolves in sodium hydroxide forming a solution of compound (C). The compound (A) on heating with coke gives a metal (D) and a gas (E) which burns with blue flame. (B) also dissolves in ammonium sulphate solution mixed with ammonium hydroxide. Solution of compound (A) in dilute HCl gives a bluish white / white precipitate (F) with excess of K4[Fe(CN)₆]. Identify (A) to (F) and explain the reactions.
- Ans. (A) ZnO, (B) Zn(OH)₂, (C) Na₂ ZnO₂, (D) Zn, (E) CO, (F) K₂Zn₃ [Fe(CN)₆]₂

Sol.
$$Zn(OH)_2$$
 (B) $\xrightarrow{\Delta}$ ZnO (A) + H₂O.

 $Zn(OH)_2$ (B) \downarrow + 2OH⁻ \longrightarrow [Zn(OH)₄]²⁻ (C) (soluble complex).

 $ZnO(A) + C \xrightarrow{\Delta} Zn(D) + CO(E).$

 $Zn(OH)_2$ (B) + 4NH₃ \longrightarrow [Zn(NH₃)₄]²⁺ (soluble complex) + 2OH⁻.

 $ZnO + 2HCI \longrightarrow ZnCl_2 + H_2O.$

 $3ZnCl_2 + 2K_4[Fe(CN)_6] \longrightarrow K_2Zn_3[Fe(CN)_6]_2 \downarrow$ (bluish white/white) (F) + 6KCl.

- **3.** An unknown inorganic compound (X) gave the following reactions:
 - (i) The compound (X) on heating gave a residue, oxygen and oxide of nitrogen.
 - (ii) An aqueous solution of compound (X) on addition to tap water gave a turbidity which did not dissolve in HNO₃.
 - (iii) The turbidity dissolves in NH₄OH.
 - Identify the compound (X) and give equations for the reactions (i), (ii) & (iii).
- **Ans.** $X = AgNO_3$
- **Sol.** $2AgNO_3(X) \xrightarrow{\Delta} 2Ag + 2NO_2 + O_2$.

AgNO₃ (aq.) + CI⁻ \longrightarrow AgCl \downarrow (white) + NO₃⁻.

AgCl + 2NH₃ \longrightarrow [Ag(NH₃)₂]⁺ (soluble complex).

4. Amongst [TiF₆]^{2–}, [CoF₆]^{3–}, Cu₂ Cl₂ and [NiCl₄]^{2–} [Atomic number; Ti = 22, Co = 27, Cu = 29, Ni = 28] the colourless species are :

(A) [TiF₆]²⁻ and [Cu₂Cl₂] (B) Cu₂Cl₂ and [NiCl₄]²⁻ (C) [TiF₆]²⁻ and [CoF₆]³⁻ (D) [CoF₆]³⁻ and [NiCl₄]²⁻

Sol. (A) In $[TiF_6]^{2-}$ the titanium is in +4 oxidation state having the electronic configuration $[Ar]^{18} 3d^0 4s^0$. Similarly in Cu₂Cl₂ the copper is in +1 oxidation state having the electronic configuration $[Ar]^{18} 3d^{10} 4s^0$. As they do not have any unpaired electrons for d-d transition, they are therefore colourless. In [NiCl₄]^{2–} the nickel is in +2 oxidation state and electronic configuration is [Ar]¹⁸ 3d⁸ 4s⁰. As it has two unpaired electrons, so the complex is coloured.

In $[CoF_6]^{3-}$, the cobalt is in +3 oxidation state having electron configuration [Ar] $3d^6 4s^0$. As it has four unpaired electrons, so the complex is coloured.

- 5. On the basis of trends in the properties of the 3d-series elements, suggests possible M²⁺ aqua ions for use as reducing agents, and write a balanced chemical equation for the reaction of one of these ions with O₂ in acidic solution.
- Sol. Because oxidation state +2 is most stable for the later elements of 3d-series elements, strong reducing agents include ions of the metals on the left of the series: such ions include V²⁺ (aq) and Cr²⁺ (aq) The Fe²⁺ (aq) ion is only weakly reducing. The Co²⁺ (aq), Ni²⁺ (aq), and Cu²⁺ (aq) ions are not oxidized in water.

 $Fe^{3+} \xrightarrow{+ 0.77} Fe^{2+} \xrightarrow{- 0.44} Fe$

The chemical equation for the oxidation is then

4 Fe²⁺ (aq) + O₂(g) + 4H⁺ (aq) \longrightarrow 4Fe³⁺ (aq) + 2H₂O (\Box).

6. Match the reactions given in column-I with the characteristic(s) of the reaction products given in column-II.

(A) TiCl₄ Zn→
(b) FeCl₃ 573 K→
(c) KMnO₄ 750 K→
(c) KMnO₄ 750 K→
(c) KMnO₄ 4 (cold & conc.)
(c) KMnO₄ 750 K→

Sol. (A) TiCl₄ \xrightarrow{Zn} TiCl₃, violet (one unpaired electron so d-d transition is possible).

(B) $2FeCl_3 \xrightarrow{573 \text{ K}} Cl \xrightarrow{Cl} Fe \xrightarrow{Cl} Cl gas dimer.$

(C) 2KMnO₄ $\xrightarrow{750 \text{ K}}$ K₂MnO₄ green (one unpaired electron so d-d transition is possible) + MnO₂ + O₂

(D) $K_2Cr_2O_7 + 2H_2SO_4 \longrightarrow 2CrO_3$ bright orange (diamagnetic) + 2KHSO₄ + H₂O.

- 7. Which of the following is true for the species having 3d⁴ configuration ?
 (A) Cr²⁺ is reducing in nature.
 (B) Mn³⁺ is oxidising in nature.
 (C) Both (A) and (B)
 (D) None of these
- **Sol.** Cr²⁺ is reducing as its configuration changes from d⁴ to d³, the latter having a half-filled t³_{2g} energy level of 3d orbitals in octahedral crystal field spliting. On the other hand, the change from Mn³⁺ to Mn²⁺ results in the half-filled (d⁵) configuration which has extra stability. Therefore, (C) option is correct.
- 8. Which of the following increasing order of oxidising power is correct for the following species ? $VO_{2^{+}}$, $MnO_{4^{-}}$, $Cr_2O_7^{2^{-}}$

(A) $VO_2^+ < Cr_2O_7^{2-} < MnO_4^-$	(B) $VO_2^+ < MnO_4^- < Cr_2O_7^{2-}$
(C) $Cr_2O_7^{2-} < VO_2^+ < MnO_4^-$	(D) $Cr_2O_7^{2-} < MnO_4^- < VO_2^+$

Sol. This is attributed to the increasing stability of the lower species to which they are reduced. MnO₄⁻ is reduced to Mn²⁺ which has stable half filled valence shell electron configuration [3d⁵]. Cr₂O₇²⁻ is reduced to Cr³⁺ which has half filled t³_{2g} energy level of 3d orbitals in octahedral crystal field spliting

 $VO_{2^{+}}$ is reduced to $V^{3_{+}}$ which has electronic configuration [Ar]¹⁸3d²4s⁰.

So the order of increasing stability of the reduced species is $Mn^{2+} > Cr^{3+} > V^{3+}$ and, therefore, the increasing order of oxidising power is $VO_2^+ < Cr_2O_7^{2-} < MnO_4^-$. Therefore, (A) option is correct.

- 9. Which of the following statement(s) is/are correct ?
 - (A) Transition metals and many of their compounds show paramagnetic behaviour.
 - (B) The enthalpies of atomisation of the transition metals are high
 - (C) The transition metals generally form coloured compounds
 - (D) Transition metals and their many compounds act as good catalyst.
- **Sol.** (A) As metal ions generally contain one or more unpaired electrons in them & hence their complexes are generally paramagnetic.

(B) Because of having larger number of unpaired electrons in their atoms, they have stronger inter atomic interaction and hence stronger bonding between the atoms.

(C) According to CFT, in presence of ligands the colour of the compound is due to the d-d transition of the electrons.

(D) This activity is ascribed to their ability to adopt multiple oxidation state and to form complexes. Therefore, (A,B,C,D) options are correct.

- **10.** When CO₂ is passed into aqueous :
 - (A) Na₂CrO₄ solution, its yellow colour changes to orange.
 - (B) K_2MnO_4 solution, it disproportionates to $KMnO_4$ and MnO_2 .
 - (C) Na₂Cr₂O₇ solution, its orange colour changes to green.
 - (D) KMnO₄ solution, its pink colour changes to green.
- **Sol.** (A) Na₂CrO₄ $\xrightarrow{H^+}$ Na₂Cr₂O₇ (orange colour)
 - (B) $MnO_{4^{2-}} \xrightarrow{H^+} MnO_{4^-} + MnO_2$, in neutral or acidic medium
 - (C) False In acidic medium no colour change takes place.

(D) $MnO_{4^-} + e^- \xrightarrow{OH^-} MnO_{4^{2-}}$; in strong alkaline medium pink colour of KMnO₄ changes to green. Therefore, (A,B) options are correct.

- 11. Which of the following statement(s) is (are) not correct with reference to ferrous and ferric ions
 - (A) Fe³⁺ gives brown colour with potassium ferricyanide
 - (B) Fe²⁺ gives blue precipitate with potassium ferricyanide
 - (C) Fe^{3+} gives red colour with potassium sulphocyanide
 - (D) Fe^{2+} gives brown colour with potassium sulphocyanide
- **Sol.** Fe³⁺ produces red colouration with KSCN but Fe²⁺ does not give brown colour with KSCN. Therefore, (D) option is correct.
- **12. Statement-1** : Ammonical silver nitrate converts glucose to gluconic acid and metallic silver is precipitated.

Statement-2 : Glucose acts as a weak reducing agent.

- (A) Statement-1 is true, statement-2 is true; statement-2 is a correct explanation for statement-1.
- (B) Statement-1 is true, statement-2 is true; statement-2 is NOT a correct explanation for statement-1
- (C) Statement-1 is true, Statement-2 is false
- (D) Statement-1 is false, Statement-2 is true
- **Sol.** (A) $Ag_2O + C_6H_{12}O_6 \rightarrow 2Ag + C_6H_{12}O_7$.

Statement-1: The number of unpaired electrons in the following gaseous ions Mn³⁺, Cr³⁺, V³⁺ and Ti³⁺ 13. are 4, 3, 2 and 1 respectively. Statement-2 : Cr³⁺ is most stable in aqueous solution among these ions. (A) Statement-1 is True, Statement-2 is True; Statement-2 is a correct explanation for Statement-1. (B) Statement-1 is True, Statement-2 is True; Statement-2 is NOT a correct explanation for Statement-1 (C) Statement-1 is True, Statement-2 is False (D) Statement-1 is False, Statement-2 is True $Mn^{3+} = [Ar]^{18} 3d^4$, $Cr^{+3} = [Ar]^{18} 3d^3$, $V^{3+} = [Ar]^{18} 3d^2$, $Ti^{3+} = [Ar]^{18} 3d^1$ Sol. **(B)** Cr^{3+} is most stable in aqueous solution because it has half filled t^{3}_{2g} energy level of 3d orbitals in octahedral crystal field spliting and according to crystal field theory (CFT) it has highest value of CFSE i.e. 1.2 Δ_0 . 14. S_1 : Interstitial compounds have high melting points, higher than those of pure metals. **S**₂ : Permanganate titrations in presence of hydrochloric acid are unsatisfactory. S₃: KMnO₄ does not act as an oxidising agent in strong alkaline medium. S_4 : KMnO₄ on heating in a current of H₂ gives MnO. (A) T T F T(B) T F F T (C) T F T T (D) F F T F Sol. **S**₁ : Due to strong interatomic forces. S₂: Some of the hydrochloric acid is oxidised to chlorine and thus we get less volume of KMnO₄ than the actual one. $S_3: MnO_4^- + e^- \xrightarrow{OH^-} MnO_4^{2-}$ $S_4: 2KMnO_4 + 5H_2 \longrightarrow 2KOH + 2MnO + H_2O.$ Therefore, (A) option is correct. 15. Match the reactions in Column I with nature of the reactions/type of the products in Column II. Column I Column II (A) AgNO₃(aq) + I_2 (excess) + $H_2O \longrightarrow$ (p) Disproportionation (B) $K_2MnO_4(aq) + CO_2(q) \longrightarrow$ (q) Comproportionation (C) Na₂Cr₂O₇ + C $\xrightarrow{\Delta}$ (r) Redox (D) $CuCl_2(aq) + Cu(s) \longrightarrow$ (s) One of the products is insoluble in water $\begin{array}{l} (A \rightarrow \ p, r, s) ; (B \rightarrow \ p, r, s) ; (C \rightarrow \ r, s) : (D \rightarrow q, r, s) \\ (A) 5 AgNO_3(aq) + 3 \stackrel{0}{I}_2 (excess) + 3H_2O \longrightarrow \stackrel{+V}{HIO_3} + 5Ag\stackrel{-1}{SAgI} + 5HNO_3 \end{array}$ Ans. Sol. So it is redox and disproportionation reaction. Agl insoluble in water. (B) $3K_2MnO_4$ (aq) + 2CO₂(g) \longrightarrow 2KMnO₄ + MnO₂ + 2K₂CO₃ So it is redox and disproportionation reaction. MnO₂ insoluble in water. (C) $2\overset{0}{C}$ + Na₂^{+VI}Cr₂O₇ $\xrightarrow{\Delta}$ Cr₂O₃ + Na₂^{+IV}CO₃ + $\overset{+II}{CO}$ \uparrow So it is redox reaction. Cr₂O₃ (green pigment) is insoluble in water. (D) $\overset{+\text{II}}{\text{CuCl}_2}$ (aq) + $\overset{0}{\text{Cu}}(s) \longrightarrow \overset{+\text{I}}{\text{Cu}_2\text{Cl}_2}(s)$ So it is redox and comproportionation reaction. Cu₂Cl₂ is insoluble in water. 16. What is the composition of mischmetal alloy and what are it's uses ? Mischmetal consists of lanthanoid metal (~95%) and iron (~5%) and traces of S, C, Ca and AI. Ans. Mischmetal is used in Mg based alloy to produce bullets, shell and lighter flint.