

THE D AND F BLOCK ELEMENTS

Element		Sc	Ti	V	Cr	Mn	Fe	Co	Ni	Cu	Zn
Atomic number		21	22	23	24	25	26	27	28	29	30
Electronic configuration											
	M	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 ¹ 4s ¹	3d ² 4s ¹	3d ³ 4s ¹	3d ⁵	3d ⁵ 4s ¹	3d ⁶ 4s ¹	3d ⁷ 4s ¹	3d ⁸ 4s ¹	3d ¹⁰	3d ¹⁰ 4s ¹
	M ²⁺	3d ¹	3d ²	3d ³	3d ⁴	3d ⁵	3d ⁶	3d ⁷	3d ⁸	3d ⁹	3d ¹⁰
	M ³⁺	[Ar]	3d ¹	3d ²	3d ³	3d ⁴	3d ⁵	3d ⁶	3d ⁷	–	–
Enthalpy of atomisation, Δ _a H [⊕] / kJ mol ⁻¹											
		326	473	515	397	281	416	425	430	339	126
Ionisation enthalpy / Δ _i H ⁻ / kJ mol ⁻¹											
Δ _i H [⊕]	I	631	656	650	653	717	762	758	736	745	906
Δ _i H [⊕]	II	1235	1309	1414	1592	1509	1561	1644	1752	1958	1734
Δ _i H [⊕]	III	2393	2657	2833	2990	3260	2962	3243	3402	3556	3829
Metallic/ionic	M	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		–	-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	–	–	–
Density/g cm ⁻³		3.43	4.1	6.07	7.19	7.21	7.8	8.7	8.9	8.9	7.1

d-block Elements										
Last electron enters in $(n - 1)d$ subshell. d-block elements lie on middle of periodic table.										
General Introduction										
1.	Occurrence	Three of transition metals are very abundant in the earth's crust.								
		<p>Fe \rightarrow 4th most abundant elements in earth's crust by weight.</p> <p>Ti \rightarrow 5th most abundant elements in earth's crust by weight.</p> <p>Mn \rightarrow 12th most abundant elements in earth's crust by weight.</p>								
	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	<p>Less abundant than 1st d- series elements.</p> <p>Tc \rightarrow does not occur in nature.</p> <p>Last six members of 2nd and 3rd row occur less than 0.16 ppm in earth's crust.</p>								
	Coinage metals	Cu, Ag and Au are called coinage metals.								
	Noble metals	Cu, Ag, Au, Hg and Pt.								
2.	No. of elements	<p>Total d-block elements – (40)</p> <p>Total transition elements – (36). If 112 elements are included in periodic table. (Except Zn, Cd, Hg and Uub)</p>								
3.	Groups	IIIB to VIIB, VIII, IB, IIB or group 3 to 12 (IUPAC).								
4.	Period	<p>4th to 7th</p> <table border="1" style="margin-left: auto; margin-right: auto;"> <tbody> <tr> <td>4th period</td> <td>3d- series (Sc - Zn)</td> </tr> <tr> <td>5th period</td> <td>4d- series (Y - Cd)</td> </tr> <tr> <td>6th period</td> <td>5d- series (La – Hg)</td> </tr> <tr> <td>7th period</td> <td>6d- series (Ac – Uub)</td> </tr> </tbody> </table>	4 th period	3d- series (Sc - Zn)	5 th period	4d- series (Y - Cd)	6 th period	5d- series (La – Hg)	7 th period	6d- series (Ac – Uub)
4 th period	3d- series (Sc - Zn)									
5 th period	4d- series (Y - Cd)									
6 th period	5d- series (La – Hg)									
7 th period	6d- series (Ac – Uub)									
Periodic trends and chemical properties										
S.N.	Properties	DISCRIPTION								
1	Metallic character	<p>All the transition elements are metals, therefore they are showing all the metallic properties.</p> <p>Good conductor of heat and electricity.</p> <p>Metallic lustre, strong and ductile.</p>								

With the exception of Zn, Cd, Hg, Mn all the transition metals have or more regular metallic structure.

The transition metals are hard and have low volatility (except Zn, Cd and Hg).

Lattice Structures of Transition Metals (Table-1)

Sc	Ti	V	Cr	Mn	Fe	Co	Ni	Cu	Zn
hcp	hcp	bcc	bcc	X	bcc	ccp	ccp	ccp	X
(bcc)	(bcc)		(bcc,ccp)	(hcp)	(hcp)				(hcp)
Y	Zr	Nb	Mo	Tc	Ru	Rh	Pd	Ag	Cd
Hcp	hcp	bcc	bcc	hcp	hcp	ccp	ccp	ccp	X
(bcc)	(bcc)								(hcp)
La	Hf	Ta	W	Re	Os	Ir	Pt	Au	Hg
hcp	hcp	bcc	bcc	hcp	hcp	ccp	ccp	ccp	X
(ccp, bcc)	(bcc)								

bcc = body centred cubic ; hcp = hexagonal close packed

ccp = cubic close packed ; X = a typical metal structure

2 Electronic configuration $(n - 1)d^{1-10} ns^{1 \text{ or } 2}$

	3 IIIB	4 IVB	5 VB	6 VIB	7 VIIB	8 VIII	9 VIII	10 VIII	11 IB	12 IIB
3d Series	21 Sc $3d^1 4s^2$	22 Ti $3d^2 4s^2$	23 V $3d^3 4s^2$	24 Cr $3d^5 4s^1$	25 Mn $3d^5 4s^2$	26 Fe $3d^6 4s^2$	27 Co $3d^7 4s^2$	28 Ni $3d^8 4s^2$	29 Cu $3d^{10} 4s^1$	30 Zn $3d^{10} 4s^2$
4d Series	39 Y $4d^1 5s^2$	40 Zr $4d^2 5s^2$	41 Nb $4d^4 5s^1$	42 Mo $4d^5 5s^1$	43 Tc $4d^5 5s^2$	44 Ru $4d^7 5s^1$	45 Rh $4d^8 5s^1$	46 Pd $4d^{10}$	47 Ag $4d^{10} 4s^1$	48 Cd $4d^{10} 4s^2$
5d Series	57 La* $5d^1 6s^2$	72 Hf $4f^{14} 5d^2 6s^2$	73 Ta $5d^3 6s^2$	74 W $5d^4 6s^2$	75 Re $5d^5 6s^2$	76 Os $5d^6 6s^2$	77 Ir $5d^7 6s^2$	78 Pt $5d^9 6s^1$	79 Au $5d^{10} 6s^1$	80 Hg $5d^{10} 6s^2$
6d Series	89 Ac** $6d^1 7s^1$	104 Rf	105 Db	106 Sg	107 Bh	108 Hs	109 Mt	110 Ds	111 Rg	112 Cn

Electrons are being filled penultimate shells, thus outer electronic configuration remains same , with exception in some cases, e.g.Cr and Cu.

Cr and Cu attain extra stability of half-filled and full-filled electronic configuration by changing the expected pattern of electronic configuration.

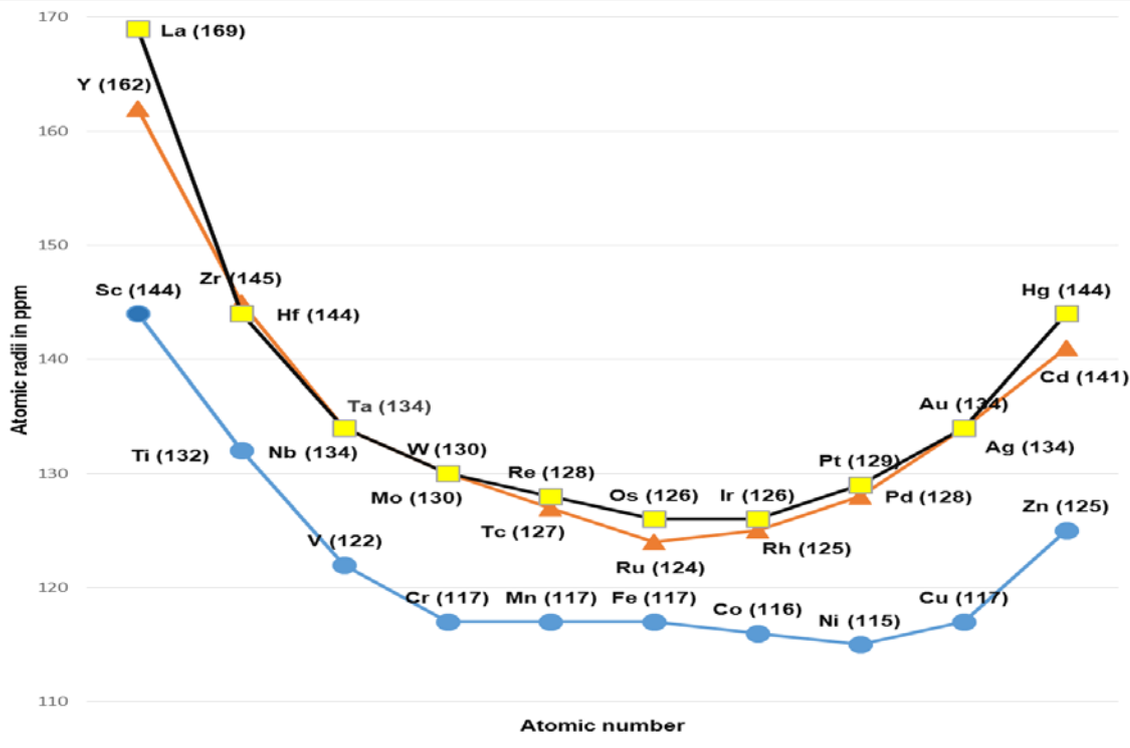
The electronic structure of 2nd and 3rd rows do not always follow the pattern of the first row, e.g. Group 10

Ni	$3d^8 4s^2$
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			Pd	$4d^{10}5s^0$																							
			Pt	$5d^9 6s^1$																							
		<p>d-orbitals are complete at Cu, Pd, Ag, Au.</p> <table border="1" style="margin-left: auto; margin-right: auto;"> <tr> <td>Ni ($3d^8 4s^2$)</td> <td>Cu($3d^{10}4s^1$)</td> <td>Zn($3d^{10}4s^2$)</td> </tr> <tr> <td>Pd($4d^{10}5s^0$)</td> <td>Ag($4d^{10}5s^1$)</td> <td>Cd($4d^{10}5s^2$)</td> </tr> </table> <p>Even though ground state of the atom has a d^{10} configuration Pd and coinage metals behaves as typical transition elements, as in their most common oxidation states they have incomplete d-orbital.</p>				Ni ($3d^8 4s^2$)	Cu($3d^{10}4s^1$)	Zn($3d^{10}4s^2$)	Pd($4d^{10}5s^0$)	Ag($4d^{10}5s^1$)	Cd($4d^{10}5s^2$)																
Ni ($3d^8 4s^2$)	Cu($3d^{10}4s^1$)	Zn($3d^{10}4s^2$)																									
Pd($4d^{10}5s^0$)	Ag($4d^{10}5s^1$)	Cd($4d^{10}5s^2$)																									
	Transition elements	<p>A transition element is defined as the one which has incompletely filled <i>d</i> orbitals in its ground state or in any one of its oxidation states.</p> <p>Zinc, Cadmium and Mercury of group 12 have full d^{10} configuration in their ground state as well as in their common oxidation states and hence, they are not regarded as transition metals.</p> <p>All transition elements are d-block elements but all d-block elements are not transition elements. (as per NCERT)</p>																									
3	Size of atoms and ions	<p>In the transition series (e.g. in the first transition series), the covalent radii of the elements decrease from left to right across a row until near the end when the size increases slightly.</p>																									
	d-orbital contraction	<p>On moving from left to right, extra protons are placed in the nucleus and the extra electron are added. The orbital electron shield the nuclear charge incompletely. Thus the nuclear charge attracts all the electrons more strongly, hence a contraction in size occurs. The radii of the elements from Cr to Cu, are very close to one another because the extra electron being added increases the repulsion between the electrons and counter balances the increased nuclear charge on the outer electrons (4s). As a result of this, the size of the atom does not change much in moving from Cr to Cu and for zinc this repulsion even dominates the nuclear charge so size slightly increases.</p> <table border="1" style="margin-left: auto; margin-right: auto;"> <thead> <tr> <th>Element</th> <th>Sc</th> <th>Ti</th> <th>V</th> <th>Cr</th> <th>Mn</th> <th>Fe</th> <th>Co</th> <th>Ni</th> <th>Cu</th> <th>Zn</th> </tr> </thead> <tbody> <tr> <td>Atomic radius (pm)</td> <td>144</td> <td>132</td> <td>122</td> <td>117</td> <td>117</td> <td>117</td> <td>116</td> <td>115</td> <td>117</td> <td>125</td> </tr> </tbody> </table>				Element	Sc	Ti	V	Cr	Mn	Fe	Co	Ni	Cu	Zn	Atomic radius (pm)	144	132	122	117	117	117	116	115	117	125
Element	Sc	Ti	V	Cr	Mn	Fe	Co	Ni	Cu	Zn																	
Atomic radius (pm)	144	132	122	117	117	117	116	115	117	125																	
	Lanthanide Contraction	<p>As we move along the lanthanide series, the nuclear charge increases by one unit at each successive element. The new electron is added into the same subshell (4f). As a result, the attraction on the electrons by the nucleus increases and this tends to decrease the size. Further, as the new electron is added into the f-subshell, there is imperfect shielding of one electron by another in this subshell due to the shapes of these f-orbitals. This imperfect shielding is unable to counter balance the effect of the increased nuclear charge. Hence, the net result is a contraction in the size though the decrease is very small.</p>																									

1. **Silver nitrate, $AgNO_3$ (Lunar Caustic)**

Thus covalent and ionic radii of Nb (5th period) and Ta (6th period) are almost same due to poor shielding of f-orbitals electrons.



Ti < Zr ≈ Hf (Lanthanide Contraction)

Sc < Y < La

Due to Lanthanide contraction elements of 4d and 5d series belong to a particular group have almost same atomic radii. Ex. Zr ≈ Hf, Tc ≈ Re, Nb ≈ Ta, Ru ≈ Os etc.

4

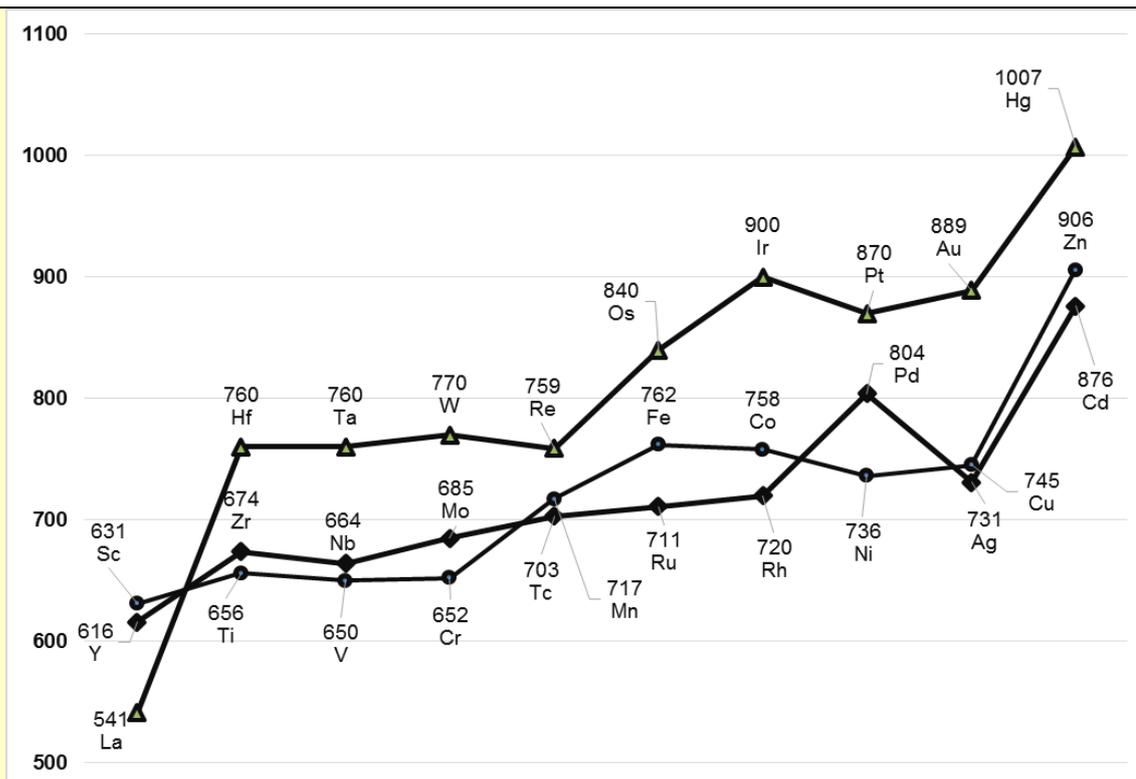
Ionization energies of transition metals

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 metallic silver, reduced by organic tissue. It is therefore, stored in colored Bottles.
- (ii) Solutions of halides phosphates, sulfides chromates thiocyanates, sulphates and thiosulphates salt with silver nitrate solution.



First ionization energies of the d-block elements (Numerical values are given in kJ mol^{-1}).

- (i) The first ionisation potential for the first four 3d-block elements (Sc, Ti, V & Cr) differ only slightly from one another.
- (ii) Similarly the value first ionisation energy of Fe, Co, Ni & Cu also are fairly close to one another.
- (iii) The value of first ionisation energy for Zn is considerably higher due to the extra stability of $3d^{10}4s^2$ level which is completely filled in Zn atom.
- (iv) The first ionization energy of Mn is slightly higher than adjacent elements in 3d series due to half-filled stable configuration.
- (v) Second ionisation energy value for Cr & Cu are higher as the Cr^+ & Cu^+ ion are extra stable due to $3d^5$ and $3d^{10}$ Configuration.
- (vi) In vertical columns, the ionisation energy decrease from first member to the second member.
- (vii) However the third member (5d series) has higher value than second member (4d-series) due to lanthanide contraction. I.E. (3^{rd} T.S.) > I.E. (2^{nd} T.S.) > I.E. (1^{st} T.S.)

5

Oxidation-State

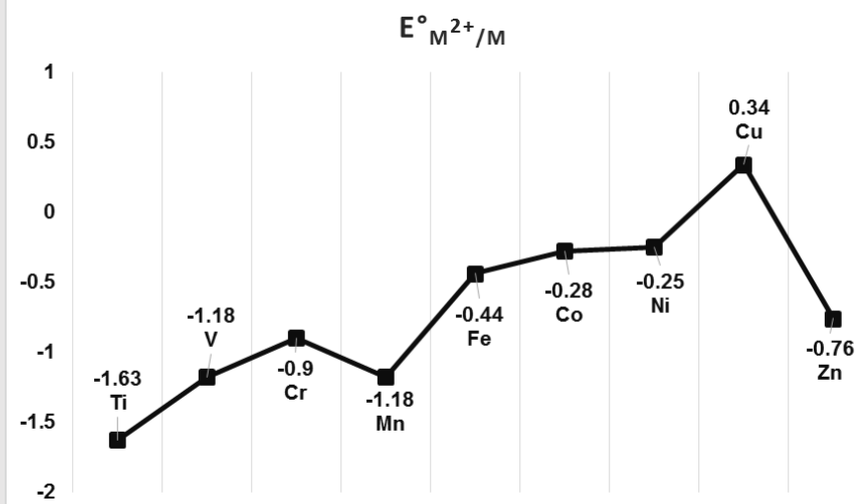
They show variable oxidation-state.

Sc	Ti	V	Cr	Mn	Fe	Co	Ni	Cu	Zn
			+1					+1	
	+2	+2	+2	+2 ^(a)	+2	+2	+2	+2	+2

			<u>+3</u>	+3	<u>+3</u>	<u>+3</u>	+3	+3	<u>+3</u>	<u>+3</u>		
				<u>+4</u>	+4	+4	<u>+4^(b)</u>	+4	+4	+4		
					+5	+5	+5					
						+6	+6	+6				
							+7					
		(a) Acidic medium	(b) Basic medium									
		* Shaded states are popular ones.										
		* Underline states are the most stable ones.										
		The oxidation states change in units of 1, unlike that in p-block elements, where generally oxidation state differs by units of 2.										
		<p>Trends of Oxidation state</p> <p>Highest oxidation state of 3d-series elements can be calculated by $n + 2$ (where n = number of unpaired electrons) It is not applied for Cr and Cu.</p> <p>The transition metal ions having stable configuration like d^0, d^5 or d^{10} are more stable. Ex. Sc^{+3}, Ti^{+4}, V^{+5}, Fe^{+3}, Mn^{+2}, Zn^{+2} etc.</p> <p>Most common oxidation state among the transition elements is +2.</p> <p>Highest oxidation state shown by transition elements of '4d' and '5d' series is +8 by Ru (44) and Os (76).</p> <p>The highest oxidation states (+4, +5, +6, +7, +8) are found in compounds of oxide and oxyfluoride because fluorine and oxygen are most electronegative elements and good oxidants.</p> <p>e.g. MnO_4^-, $Cr_2O_7^{2-}$, CrO_4^{2-}, MnO_2, MnO_4^{2-}, CrO_2F_2</p> <p>Lower oxidation states even negative (-2, -1, 0, +1) is stabilised by π-bonding ligands such as carbon monoxide and dipyridine due to phenomena of synergic bonding $Ni(CO)_4$ and $Fe(CO)_5$, oxidation state of Ni and Fe is zero.</p> <p>They form ionic compounds in low oxidation states and covalent compounds in Higher oxidation state.</p>										
		Formulas of Halides of 3d Metals										
		Key: $X = F \rightarrow I$; $X^I = F \rightarrow Br$; $X^{II} = F, Cl$; $X^{III} = Cl \rightarrow I$										
		OXIDATION NUMBER	Sc	Ti	V	Cr	Mn	Fe	Co	Ni	Cu	Zn
		+6				CrF_6						

	+5			VF ₅	CrF ₅						
	+4		TiX ₄	VX ₄ ¹	CrX ₄	MnF ₄					
	+3		TiX ₃	VX ₃	CrX ₃	MnF ₃	FeX ₃ ¹	CoF ₃			
	+2		TiX ₂ ^{III}	VX ₂	CrX ₂	MnX ₂	FeX ₂	CoX ₂	NiX ₂	CuX ₂ ^{II}	ZnX ₂
	+1									CuX ^{III}	
Formulas of Oxides of 3d Metals (* mixed oxides)											
OXIDATION NUMBER	Sc	Ti	V	Cr	Mn	Fe	Co	Ni	Cu	Zn	
+7					Mn ₂ O ₇						
+6					CrO ₃						
+5			V ₂ O ₅								
+4		TiO ₂	V ₂ O ₄	CrO ₂	MnO ₂						
+3	Sc ₂ O ₃	Ti ₂ O ₃	V ₂ O ₃	Cr ₂ O ₃	Mn ₂ O ₃	Fe ₂ O ₃					
					Mn ₃ O ₄ [*]	Fe ₃ O ₄ [*]	Co ₃ O ₄ [*]				
+2		TiO	VO	(CrO)	MnO	FeO	CoO	NiO	CuO	ZnO	
									Cu ₂ O		
6	Colour	<p>Ionic and covalent compounds of many transition elements are coloured. They form different coloured compounds. The compounds in which metal ion has an unpaired electron that compound may be coloured. When an electron from a lower energy d-orbital is excited to a higher energy d-orbital, if the energy of excitation lies in visible range, complimentary colour is observed.</p> <p>MnO₄⁻ is coloured inspite of d⁰ configuration of Mn. It is not due to d-d transition but it occurs due to charge transfer between Mn and O.</p>									
	Oxidation State	Sc	Ti	V	Cr	Mn	Fe	Co	Ni	Cu	Zn
	(I)									Colourless	
	(II)				Blue	Pink	Green	Pink	Green	Blue	Colourless
	(III)	Colourless	Purple	Green	Green	Violet	Yellow	Blue			
	(IV)		Colourless	Blue							
7	Electrode potential										
	E ⁰ _{M²⁺/M}	<p>There is no regular trend in these values. This is attributed to the irregular variation of ionisation enthalpies (IE₁ + IE₂), hydration energies and the sublimation energies in the period.</p>									

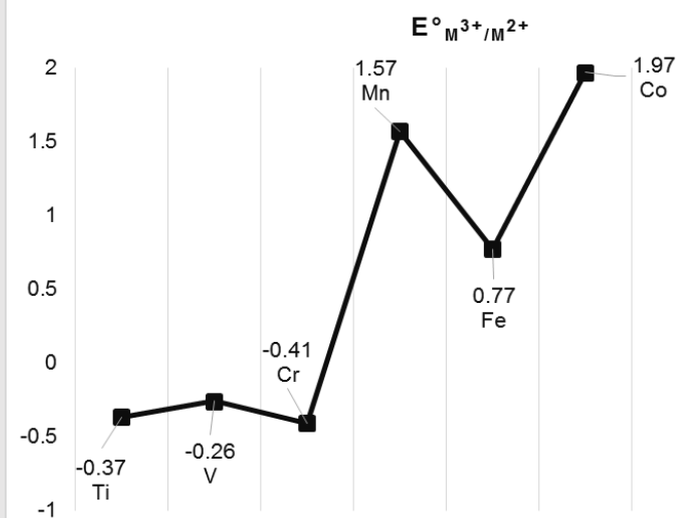
Element (M)	$\Delta_a H_q$ (M)	$\Delta_f H_1^\theta$	$\Delta_f H_2^\theta$	$\Delta_{\text{hyd}} H^\theta$ (M^{2+})	E^θ /V
Ti	469	661	1310	-1866	-1.63
V	515	648	1370	-1895	-1.18
Cr	398	653	1590	-1925	-0.90
Mn	279	716	1510	-1862	-1.18
Fe	418	762	1560	-1998	-0.44
Co	427	757	1640	-2079	-0.28
Ni	431	736	1750	-2121	-0.25
Cu	339	745	1960	-2121	0.34
Zn	130	908	1730	-2059	-0.76



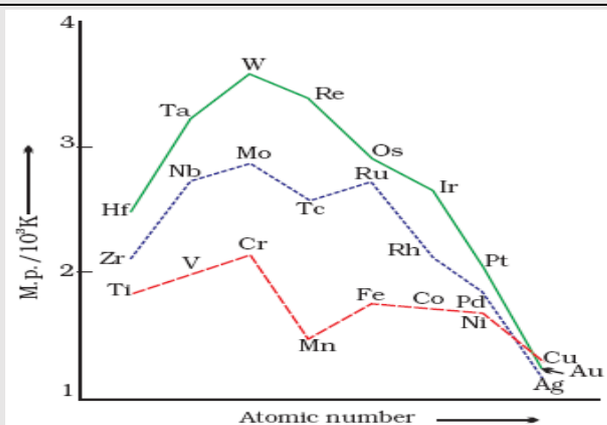
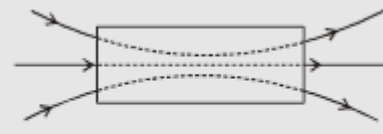
The stability of the half-filled d sub-shell in Mn^{2+} and the completely filled d^{10} configuration in Zn^{2+} are related to their E^θ values, whereas E^θ for Ni is related to the highest negative hydration energy.

It may be noted that the electrode potentials of transition metals are low in comparison to elements of group 2 (e.g., $Ca = -2.87$ V, the transition elements have fairly large ionisation enthalpies and very large enthalpies of atomisation). Compared to group 2 elements. These reduce their electrode potentials though their hydration enthalpies are large.

$E^\theta_{M^{3+}/M^{2+}}$



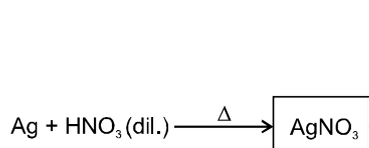
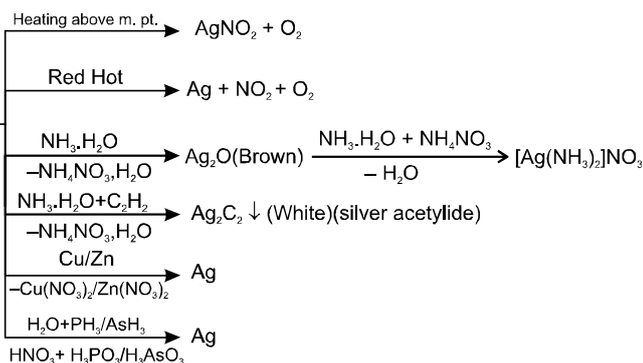
	<p>Comparative graph of $E_{M^{2+}/M}^0$ and $E_{M^{3+}/M^{2+}}^0$</p>	
	<p>Graph of E_{Red}^0</p>	
		<p>The low value for Sc reflects the stability of Sc^{3+} which has a noble gas configuration.</p> <p>The highest value for Zn is due to the removal of an electron from the stable d^{10} configuration of Zn^{2+}.</p> <p>The comparatively high value for Mn shows that Mn^{2+} (d^5) is particularly stable, whereas comparatively low value for Fe shows the extra stability of Fe^{3+} (d^5).</p>
<p>8</p>	<p>Density</p>	<p>Across a period from left to right atomic volumes decreases and atomic masses increases. Hence densities also increases across a period. The last element zinc (Zn) is an exception, having large atomic volume and hence lower density.</p> <p>Elements with highest densities are</p> <p>Osmium (Os) = 22.51 g/cm^3, Iridium (Ir) = 22.61 g/cm^3</p>
<p>9</p>	<p>Melting and Boiling point</p>	<p>High melting points are attributed to involvement of greater number of electrons from $(n-1)d$ in addition to the ns electrons in inter atomic metallic bonding thus highest for d^5, except for anomalous value for Mn, and Tc fall regularly.</p>

		<p>m.p. and b.p. are generally very high exceptions</p> <p>→ Zn (420°C)</p> <p>→ Cd (321°C)</p> <p>→ Hg liquid at room temperature.</p> <p>last of each d-series element behave a typically because the d-shell is complete and d-electrons do not participate in metallic bonding.</p>						
		<p>They have high enthalpies of atomization, which are maximum at middle of each series, indicates that one unpaired e⁻ per d-orbital is particularly favourable for strong interatomic interaction.</p> <p>Metals of 2nd and 3rd transition series has higher melting pt. bonding pt. as well as enthalpy of atomisation, than corresponding elements of the 1st series this is due to metal-metal bonding of heavy transition metals.</p>						
								
10	Magnetic character	<p>Metals, which have unpaired electrons show paramagnetism.</p> <p>Spin only magnetic moment $\mu = \sqrt{n(n+2)}$ here n = no. of unpaired electron.</p> <p>Diamagnetic substance is one which is slightly repelled by a magnetic field.</p> <p>A paramagnetic substance is one which is attracted into a magnetic field.</p> <p>In paramagnetic substance the magnetic field lines of force travel easier than they travel in vacuum. Thus it can be seen that a paramagnetic material attracts lines of force, if it is free to move, a paramagnetic material will move from a weaker to a stronger part of the field.</p>  <p style="text-align: center;">Paramagnetic material</p>						
	Ferromagnetism	<p>Ferromagnetic materials may be regarded as special case of paramagnetism in which the moments of individual domains becomes aligned and all points in the same direction. Ferromagnetic materials ⇒ Fe, Co, Ni.</p>						
11	Catalytic Properties	<p>The transition metals and their compounds are known for their catalytic activity. This is due to ability to adopt multiple oxidation states and to form complexes.</p>						
		<table border="1"> <thead> <tr> <th>Catalyst</th> <th>Used</th> </tr> </thead> <tbody> <tr> <td>Fe</td> <td>Haber's process for manufacture of NH₃.</td> </tr> <tr> <td>V₂O₅</td> <td>Contact process for H₂SO₄ manufacture.</td> </tr> </tbody> </table>	Catalyst	Used	Fe	Haber's process for manufacture of NH ₃ .	V ₂ O ₅	Contact process for H ₂ SO ₄ manufacture.
Catalyst	Used							
Fe	Haber's process for manufacture of NH ₃ .							
V ₂ O ₅	Contact process for H ₂ SO ₄ manufacture.							

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	SOME IMPORTANT ALLOY	<p><i>* Not to be memories</i></p> <table border="1"> <tbody> <tr> <td>(a)</td> <td>Bronze</td> <td>Cu (75 - 90 %) + Sn (10 - 25 %)</td> </tr> <tr> <td>(b)</td> <td>Brass</td> <td>Cu (60 - 80 %) +Zn (20 - 40 %)</td> </tr> <tr> <td>(c)</td> <td>Gun metal</td> <td>(Cu + Zn + Sn) (87 : 3 : 10)</td> </tr> <tr> <td>(d)</td> <td>German Silver</td> <td>Cu + Zn + Ni (2 : 1 : 1)</td> </tr> <tr> <td>(e)</td> <td>Bell metal</td> <td>Cu (80 %) + Sn(20 %)</td> </tr> <tr> <td>(f)</td> <td>Nichrome</td> <td>(Ni + Cr + Fe)</td> </tr> <tr> <td>(g)</td> <td>Alnico</td> <td>(Al, Ni,Co)</td> </tr> <tr> <td>(h)</td> <td>Type Metal</td> <td>Pb + Sn + Sb</td> </tr> <tr> <td>(i)</td> <td>Alloys of steel</td> <td></td> </tr> <tr> <td></td> <td>• Vanadium steel</td> <td>V (0.2 - 1 %)</td> </tr> <tr> <td></td> <td>• Chromium steel</td> <td>Cr (2 - 4 %)</td> </tr> <tr> <td></td> <td>• Nickel steel</td> <td>Ni (3 -5 %)</td> </tr> <tr> <td></td> <td>• Manganese steel</td> <td>Mn (10 -18 %)</td> </tr> <tr> <td></td> <td>• Stainless steel</td> <td>Cr (12 - 14 %)& Ni (2 - 4 %)</td> </tr> <tr> <td></td> <td>• Tunguston steel</td> <td>W (10 - 20 %)</td> </tr> <tr> <td>(o)</td> <td>14 Carat Gold</td> <td>54 % Au + Ag (14 to 30 %) + Cu (12 - 28 %)</td> </tr> </tbody> </table>	(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 %)		• Stainless steel	Cr (12 - 14 %)& Ni (2 - 4 %)		• Tunguston steel	W (10 - 20 %)	(o)	14 Carat Gold	54 % Au + Ag (14 to 30 %) + Cu (12 - 28 %)
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(k)	24 Carat Gold	100 %Au
(i)	Solder	Pb + Sn
(m)	Magnellium	Mg (10%) + Al (90%)
(n)	Duralumin	(Al + Mn + Cu)
(o)	Artificial Gold	Cu (90 %) + Al (10%)
(p)	Constantan	Cu(60%) + Ni (40%)
% of Carbon 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	2.6 to 4.3
(d)	Pig Iron	2.3 to 4.6

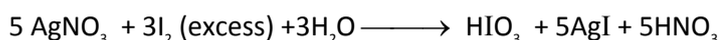
(iii) Ammonical silver nitrate is called as Tollen's reagent and used to identify reducing sugars and aldehydes.

PREPARATION**CHEMICAL PROPERTIES**

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



(iv) Reactions with Iodine :



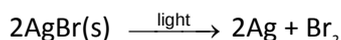
Uses : It is used

- (i) as a laboratory reagent for the identification of various acids 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.

2. Photography :

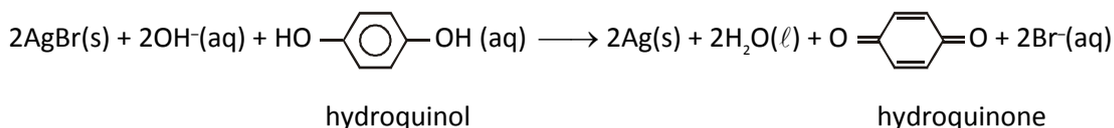
(i) A photographic film consists of a light sensitive emulsion of fine particles (grains) of silver salts in gelatine spread on a clear celluloid strip or a glass plate. AgBr is mainly used as the light sensitive material.

(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.

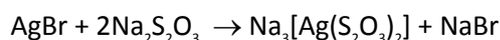


(iii) In modern photography only a short exposure of perhaps 1/100th of a second is used. In this short time, only a few atoms of silver (perhaps 10–50) are produced in each grain exposed to light. Parts of the film which have been exposed to the bright parts of the subject contain a lot of grains with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

(iv) Next the film is placed in 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.



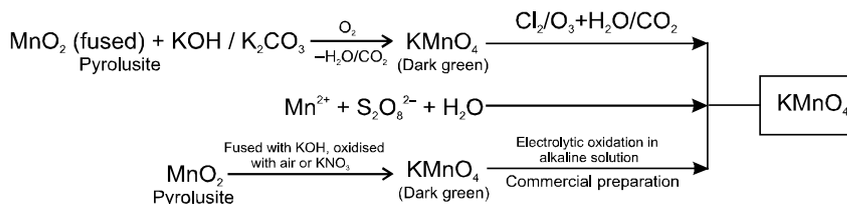
(v) If the film was brought out into daylight at this stage, the unexposed parts of the emulsion would turn black and thus destroy the picture. To prevent this happening any unchanged silver halides are removed by placing the film in a fixer solution. A solution of sodium thiosulphate is used as fixer.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



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

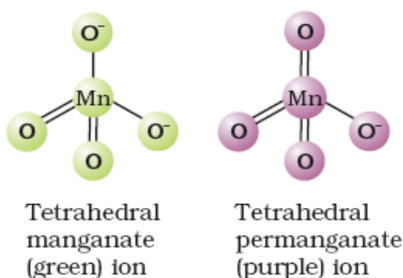
Light is passed through the negative onto a piece of paper coated with AgBr emulsion. This is then developed and fixed in the same way as before.

3. Potassium Permanganate(KMnO₄)

PREPARATION**Physical Properties :**

Purple coloured crystalline compound.

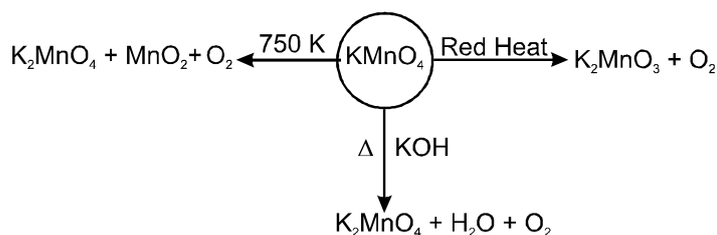
Moderately soluble in water at room temperature.



Structure of manganate and permanganate ion.

Chemical Properties

(i) Heating effect



MnO_4^{2-} in dilute alkaline, water and acidic solutions is unstable and disproportionates to give MnO_4^- and MnO_2 .



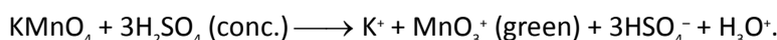
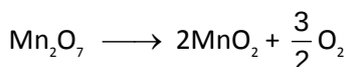
or



On heating in current of H_2 , solid KMnO_4 gives MnO



- (ii) On treatment with concentrated H_2SO_4 (KMnO_4 is taken in excess), it forms manganese heptoxide via permanganyl sulphate which decomposes explosively on heating.



- (iii) Potassium permanganate is a powerful oxidising agent.

Potassium permanganate acts as an oxidising agent in alkaline, neutral or acidic solutions.

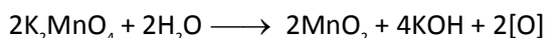
A mixture of sulphur, charcoal and KMnO_4 forms an explosive powder. A mixture of oxalic acid and KMnO_4 catches fire spontaneously after a few seconds. The same thing happens when glycerine is poured over powdered KMnO_4 .

In alkaline & neutral medium :

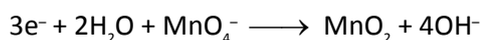
In strongly alkaline medium KMnO_4 is reduced to manganate.



However if solution is dilute then K_2MnO_4 is converted into MnO_2 which appears as a brownish precipitate.

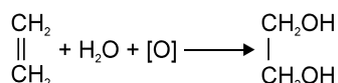


This type of behaviour is shown by KMnO_4 itself in neutral medium.



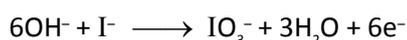
In alkaline or neutral medium KMnO_4 shows following oxidising properties.

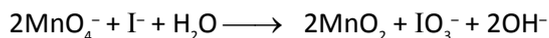
- (a) It oxidises ethene to glycol.



In alkaline medium KMnO_4 solution is also known as Bayer's reagent (1% alkaline KMnO_4 solution).

- (b) It oxidises iodide into iodate.

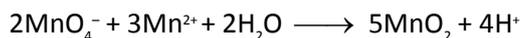




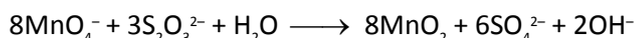
(c) H_2S is oxidised into sulphur :



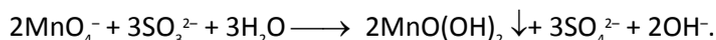
(d) Manganous salt is oxidised to MnO_2 ; the presence of zinc sulphate or zinc oxide catalyses the oxidation :



(e) In neutral/ faintly alkaline solution thiosulphate is quantitatively oxidised to sulphate.

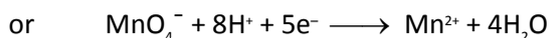


(f) In the presence of sodium hydroxide, sodium sulphite is oxidised in to sodium sulphate.



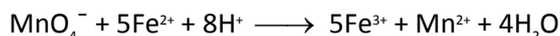
In acidic medium (in presence of dilute H_2SO_4) :

Manganous sulphate is formed. The solution becomes colourless.



This medium is used in quantitative (volumetric) estimations. The equivalent mass of KMnO_4 in acidic medium is = $\frac{\text{Molecular mass}}{5}$.

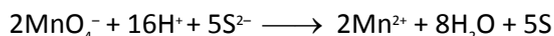
(a) Ferrous salts are oxidised to ferric salts.



(b) Iodine is evolved from potassium iodide.



(c) H_2S is oxidised to sulphur :



(d) SO_2 is oxidised to H_2SO_4 :



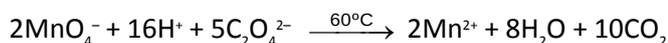
(e) Nitrites are oxidised to nitrates :



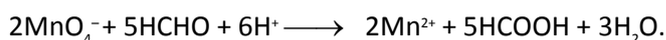
(f) Oxalic acid is oxidised to CO_2 :

This reaction is slow at room temperature, but is rapid at 60°C .

Mn(II) ions produced catalyse the reaction; thus the reaction is autocatalytic.



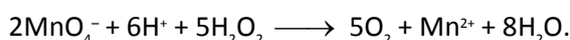
(g) HCHO is oxidised to HCOOH



(h) It oxidises hydrogen halides (HCl, HBr or HI) into X_2 (halogen)



(i) H_2O_2 is oxidised to O_2 .



Uses : It is used

(i) KMnO_4 is used as an oxidising agent in laboratory and industry.

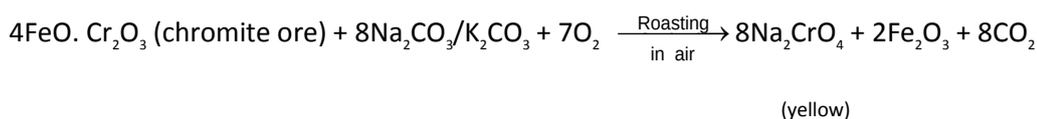
(ii) Alkaline potassium permanganate is called Bayer's reagent. This reagent is used in organic chemistry for the test of unsaturation. KMnO_4 is used in the manufacture of saccharin, benzoic acid, acetaldehyde etc.

(iii) KMnO_4 is used in qualitative analysis for detecting halides, sulphites, oxalates, etc.

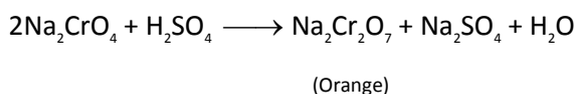
4. POTASSIUM DICHROMATE ($\text{K}_2\text{Cr}_2\text{O}_7$) :

Preparation :

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



The roasted mass is extracted with water when Na_2CrO_4 goes into the solution leaving behind insoluble Fe_2O_3 . The solution is then treated with calculated amount of H_2SO_4 .



The solution is concentrated when less soluble Na_2SO_4 crystallises out. The solution is further concentrated when crystals of $\text{Na}_2\text{Cr}_2\text{O}_7$ are obtained. Hot saturated solution of $\text{Na}_2\text{Cr}_2\text{O}_7$ is then treated with KCl when orange red crystals of $\text{K}_2\text{Cr}_2\text{O}_7$ are obtained on crystallisation. Sodium dichromate is more soluble than potassium dichromate.



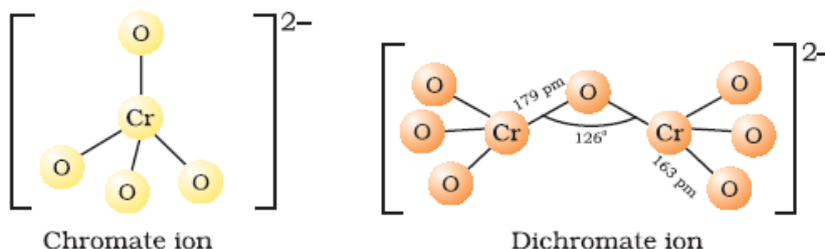
$\text{K}_2\text{Cr}_2\text{O}_7$ is preferred over $\text{Na}_2\text{Cr}_2\text{O}_7$ as a primary standard in volumetric estimation because $\text{Na}_2\text{Cr}_2\text{O}_7$ is hygroscopic in nature but $\text{K}_2\text{Cr}_2\text{O}_7$ is not.

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 .

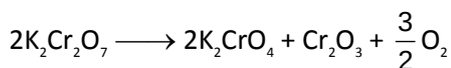
Structure of Chromate and Dichromate ion



(b) Chemical :

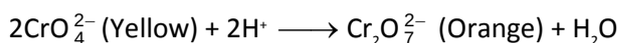
(i) Effect of heating :

On heating strongly, it decomposes liberating oxygen.



On heating with alkalis, it is converted to chromate, i.e., the colour changes from orange to yellow. On acidifying, yellow colour again changes to orange.

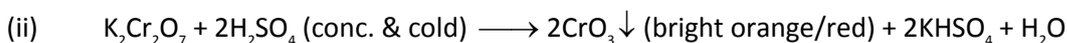




Thus CrO_4^{2-} and $\text{Cr}_2\text{O}_7^{2-}$ exist in equilibrium and are interconvertable by altering the pH of solution.



In alkaline solution, chromate ions are present while in acidic solution, dichromate ions are present.



(iii) Acidified $\text{K}_2\text{Cr}_2\text{O}_7$ solution reacts with H_2O_2 to give a deep blue solution due to the formation of CrO_5 .

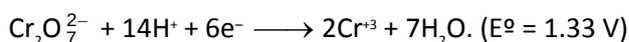


Blue colour in aqueous solution fades away slowly due to the decomposition of CrO_5 to Cr^{3+} ions and oxygen. In less acidic solution $\text{K}_2\text{Cr}_2\text{O}_7$ and H_2O_2 give salt which is violet coloured and diamagnetic due to the formation of $[\text{CrO}(\text{O}_2)(\text{OH})]^-$.

(iv) Potassium dichromate reacts with hydrochloric acid and evolves chlorine gas.

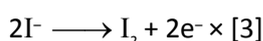
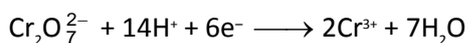


(v) It acts as a powerful oxidising agent in acidic medium (dilute H_2SO_4)

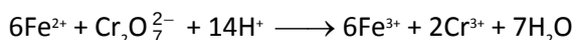


The oxidation state of Cr changes from +6 to +3.

(a) Iodine is liberated from potassium iodide :



(b) Ferrous salts are oxidised to ferric salts :



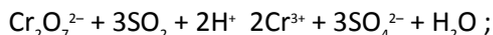
(c) Sulphites are oxidised to sulphates :



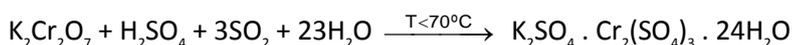
(d) H_2S is oxidised to sulphur :



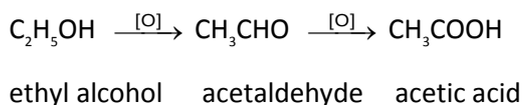
(e) SO_2 is oxidised to H_2SO_4 :



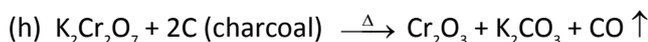
Chrome alum is obtained when acidified $\text{K}_2\text{Cr}_2\text{O}_7$ solution is saturated with SO_2 .



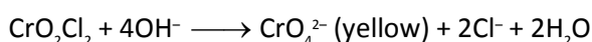
(f) It oxidises ethylalcohol to acetaldehyde and acetaldehyde to acetic acid



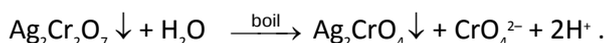
(g) It also oxidises nitrites to nitrates, arsenites to arsenates, HBr to Br_2 , HI to I_2 , etc.



(vi) Chromyl chloride test : $4\text{Cl}^- + \text{Cr}_2\text{O}_7^{2-} + 6\text{H}^+ \longrightarrow 2\text{CrO}_2\text{Cl}_2 \uparrow$ (deep red) + $3\text{H}_2\text{O}$



(vii) $\text{Cr}_2\text{O}_7^{2-}$ (concentrated solution) + $2\text{Ag}^+ \longrightarrow \text{Ag}_2\text{Cr}_2\text{O}_7 \downarrow$ (reddish brown)



(viii) $\text{Cr}_2\text{O}_7^{2-} + \text{Ba}^{2+} + \text{H}_2\text{O} \rightleftharpoons 2\text{BaCrO}_4 \downarrow + 2\text{H}^+$

As strong acid is produced, the precipitation is only partial. But if NaOH or CH_3COONa 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.

f-block elements					
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	All the f-block elements are heavy metals. It shows high melting and boiling point. The most common oxidation state of these elements is +3.			
2.	No. of elements	Total number of f-block elements – (28)			
3.	Groups	IIIB/3rd group is called longest group having 32 elements including 14 Lanthanides and 14 Actinides. <table border="1" style="margin-left: auto; margin-right: auto;"> <tr> <td>III B/ 3rd</td> </tr> <tr> <td>Sc</td> </tr> <tr> <td>Y</td> </tr> </table>	III B/ 3rd	Sc	Y
III B/ 3rd					
Sc					
Y					

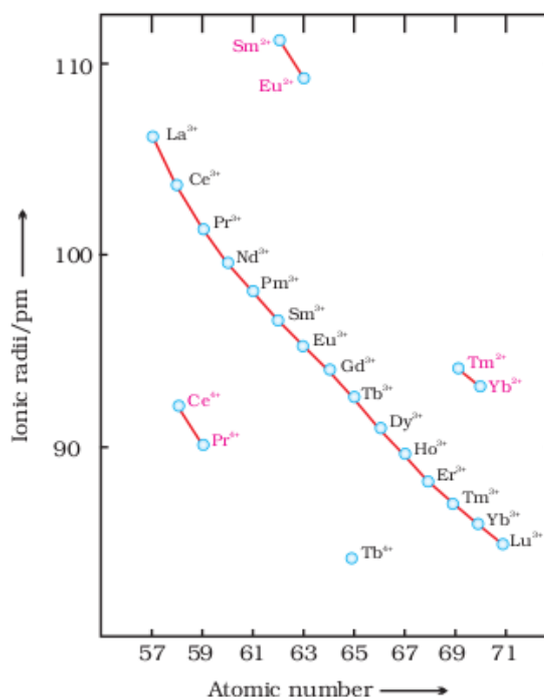
			La	Lanthanides (14) Ce ₅₈ – Lu ₇₁	
			Ac	Actinides (14) Th ₉₀ – Lr ₁₀₃	
4.	E.C.	Lanthanide series $4f^1 - 14 5d^0$ or $1 6s^2$ Actinide series $5f^1 - 14 6d^0$ or $1 7s^2$			
5.	Period		Period	III B/ 3rd	
				Sc	
				Y	
			6 th period	La	Lanthanides (14) Ce ₅₈ – Lu ₇₁
			7 th period	Ac	Actinides (14) Th ₉₀ – Lr ₁₀₃
6.	Inner transition elements	<p>The elements in which all the three shells that is ultimate (n) penultimate (n-1) and pre or antipenultimate (n-2) shell are incomplete are called inner transition elements. Ce₅₈ = [Xe] 6s², 5d¹, 4f¹</p> <p>Inner transition elements are divided into two series.</p>			
	(i) Lanthanide series or Rare earth elements or Lanthanones	<p>Ce₅₈ – Lu₇₁ 14 elements</p> <p>Lanthanides are found rarely on earth so these are called rare earth metals.</p> <p>The first element of this series is Cerium and not Lanthanum.</p> <p>In these elements, last electron enters into 4f subshell.</p> <p>They are present in IIIB group and 6th period of the periodic table.</p> <p>Promethium (₆₁Pm) is the only lanthanide which is synthetic and radioactive in nature.</p>			
	(ii) Actinide series or Man made elements or Actinones	<p>Th₉₀ – Lr₁₀₃ 14 elements.</p> <p>All the actinides are radioactive elements.</p> <p>The first element of this series is Thorium and not Actinium.</p> <p>In these elements, last electron enters into 5f subshell.</p> <p>They are present in IIIB group and 7th period of the periodic table.</p> <p>All the actinides are radioactive in nature.</p> <p>First three elements (Th, Pa, U) are found in nature while others are synthetic in nature.</p> <p>Transuranic actinides are man-made elements (Np₉₃ – Lw₁₀₃)</p>			

		<p>After U_{92} i.e. from Np_{93} onwards elements are called transuranic elements because</p> <p>(i) They are heavier than uranium.</p> <p>(ii) They are derived from uranium by nuclear reactions.</p>
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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 4f 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^{\ominus} value for Ce^{4+}/Ce^{3+} 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 f^7 configuration accounts for the formation of this ion.

However, Eu^{2+} is a strong reducing agent changing to the common +3 state. Similarly Yb^{2+} 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 Number	Name	symbol	Electronic configurations*				Radii/pm	
			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	4f 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.

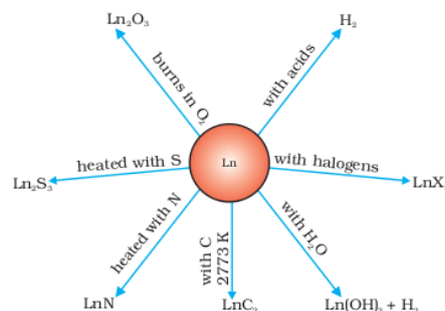
They have typical metallic structure and are good conductors of heat and electricity. Density and other properties change smoothly except for Eu and Yb and occasionally for Sm and Tm.

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^0 type (La^{3+} and Ce^{4+}) and the f^{14} type (Yb^{2+} and Lu^{3+}) 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:

$\text{Ln}^{3+}(\text{aq}) + 3e^- \rightarrow \text{Ln}(\text{s})$ 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 M_2O_3 and hydroxides $\text{M}(\text{OH})_3$. 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 ($\sim 95\%$) and iron ($\sim 5\%$) and traces of S, C, Ca and Al. 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.

Atomic Number	Name	symbol	Electronic configurations*			Radii/pm	
			M	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	Proactinium	Pa	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	5f ⁵	5f ⁴	100	90
95	Americium	Am	5f ⁷ 7s ²	5f ⁶	5f ⁵	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	Einsteinium	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^0 , f^7 and f^{14} occupancies of the 5f orbitals. Thus, the configurations of Am and Cm are $[Rn] 5f^7 7s^2$ and $[Rn] 5f^7 6d^1 7s^2$.

Although the 5f orbitals resemble the 4f orbitals in their angular part of the wave-function, they are not as buried as 4f orbitals and hence 5f electrons can participate in bonding to a far greater extent.

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.

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.

