

13.1 Transition Metals

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13.1.1 Transition Elements

Transition Elements

- Transition metals are elements with an incomplete d-subshell or that can form at least one stable cation with an incomplete d-subshell
- \blacksquare This definition distinguishes them from d-block elements, because scandium and zinc do not fit the definition
	- Scandium only forms the ion $\texttt{Sc}^{\texttt{3+}}$, configuration **[Ar] 3d⁰**
	- Zinc only forms the ion $\mathsf{Zn^{2+}}$, configuration **[Ar] 3d¹⁰**
- \blacksquare The elements of the first transition series are therefore titanium to copper

The transition elements and the d-block elements

Electron Configuration

- The full electronic configuration of the first d-series transition metals is shown in the table below
- **Following the Aufbau Principle electrons occupy the lowest energy subshells first**
- The 4s overlaps with the 3d subshell so the 4s is filled first
- Remember that you can abbreviate the first five subshells, 1s-3p, as $[Ar]$ representing the configuration of argon (known as the argon core)

Table showing the electronic configuration of the first d-series transition elements

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- From your previous studies in chemistry, you should recall there are two exceptions to the Aufbau Principle in the first row of d-block, chromium and copper
- \blacksquare In both cases an electron is promoted from the 4s to the 3d to achieve a half full and full d-subshell, respectively
- **EXTER THE FOLLO FIGHT AND FORM CONFIGURATION** Chromic Chromium and controller and you may expect:
	- Cr is [Ar] 3d⁵ 4s¹ not [Ar] 3d⁴ 4s²
	- Cu is [Ar] 3d¹⁰ 4s¹ not [Ar] 3d⁹ 4s²
- This is because the [Ar] 3d⁵ 4s¹ and [Ar] 3d¹⁰ 4s¹ configurations are **energetically more stable** and are preferred configurations
- **When forming cations, remove the 4s electrons first**

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- A complex ion is a molecule or ion, consisting of a central metal atom or ion, with a number of molecules or ions surrounding it
- A molecule or ion surrounding the central metal atom or ion is called a ligand
- Due to the different oxidation states of the central metal ions, a different number and wide variety of ligands can form bonds with the transition element
	- For example, the chromium(III) ion can form [Cr(NH₃₎₆]³⁺, [Cr(OH)₆]^{3–} and [Cr(H₂O)₆]³⁺ complex ions

Forming coloured compounds

- Another characteristic property of transition elements is that their compounds are often coloured
	- For example, the colour of the [Cr(OH) $_\mathrm{6}$]³⁻ complex (where oxidation state of Cr is +3) is **dark green**
	- Whereas the colour of the [Cr(NH₃₎₆]³⁺ complex (oxidation state of Cr is still +3) is **purple**

Transition elements as catalysts

- Since transition elements can have variable oxidation states, they make excellent catalysts
- **During catalysis, the transition element can change to various oxidation states by gaining electrons** from or donating electrons to reagents within the reaction
- Substances can also be adsorbed onto their surface and activated in the process

Magnetic Properties

- Materials are classified as diamagnetic, paramagnetic or ferromagnetic according to their behaviour when placed in an external magnetic field
- Transition metals exhibit these properties depending on their electronic configurations

13.1.2 Variable Oxidation States

Variable Oxidation States

- When transition elements forms ions they lose electrons from the 4s subshell first
- \blacksquare This is because when the orbitals are occupied, the repulsion between electrons pushes the 4s into a higher energy state so that it now becomes slightly higher in energy than the 3d subshell
	- \blacksquare The 4s is now the outer shell and loses electrons first
- \blacksquare The loss of the 4s electrons means that $+2$ is a common oxidation state in transition metals
- **The reason why the transition metals have variable oxidation states all comes down to energy**

Table showing the the common oxidation states of transition elements

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13.1.3 Complex Ions

Complex Ions

- Transition element ions can form complexes which consist of a central metal ion and ligands
- A ligand is a molecule or ion that forms a co-ordinate bond with a transition metal by donating a pair of electrons to the bond
	- \blacksquare This is also the definition of a **Lewis base**
- This means ligands have a negative charge or a lone pair of electrons capable of being donated This definition may seem familiar: a ligand is the same as a nucleophile
- Different ligands can form different numbers of dative bonds to the central metal ion in a complex
	- Some ligands can form one co-ordinate bond to the central metal ion
	- Other ligands can form two co-ordinate bonds, and some can form multiple dative bonds
- Co-ordination number is number of co-ordinate bonds to the central metal atom or ion

Common Ligands

- Water molecules frequently act as ligands. Each water molecule makes a single bond with the metal ion using one of the lone pairs on the oxygen atom
- The lone pair is donated to the partially filled d-subshell ofthe transition metal ion

Table showing examples of common monodentate ligands

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Representing complex ions

- Square brackets are used to group together the ligands and metal ion in a representation of the geometrical arrangement
- The overall charge on the complex ion is the sum of the oxidation states of all the species present

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If the ligands are neutral then the overall charge will be the same as the oxidation state of the metal ion

Examples of complexes with monodentate ligands

Co-ordination number

- The coordination number is the number of coordinate bonds to the metal ions
- This number can be the same as the number of ligands if they are monodentate, but it could be differentif bi- or multidentate ligands are present

Naming complexes

- **Complexes are named in the following way**
- \blacksquare If the overall ion is a cation then the nomenclature is:

Prefix for number of ligands/ligand name/element/oxidation number

- The prefixes are the same ones used in organic chemistry: di, tetra, hexa for 2, 4 & 6 respectively (3 & 5 are rarely encountered exceptin mixed ligand complexes)
- If the overall ion is an anion, the name of element is modified to have the name ending 'ate' and sometimes Latin word stems are used
- \blacksquare Using the examples in the illustration above, the names are:
	- tetrachlorcuprate(II)

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- hexaaquairon(II)
- hexaamminecobalt(II)
- tetracyanonickelate(II)
- Notice in these examples that
	- cuprate(Latin cuprum) and nickelate are used in place of copper and nickel as they are anions
	- Ammonia takes the prefix ammine as a ligand, which is spelt with a double 'm' unlike the functional group amine

Bidentate Ligands

- **Bidentate** ligands can each form two co-ordinate bonds to the central metal ion
- This is because each ligand contains two atoms with lone pairs of electrons
- **Examples of bidentate ligands are:**
	- 1,2-diaminoethane ($H_2NCH_2CH_2NH_2$) which is also written as 'en'
	- Ethanedioate ion (C₂O₄²⁻) which is sometimes written as **'ox'** (coming from the common name of oxalate)

Examples of complexes with bidentate ligands

Multidentate Ligands

- Some ligands contain more than two atoms with lone pairs of electrons
- These ligands can form more than two dative bonds and are said to be multidentate or polydentate ligands
- An example of a multidentate ligand is EDTA⁴⁻, which is a **hexadentate** ligand as it forms 6 dative covalent bonds to the central metal ion
- **EDTA comes from ethylenediaminetetraac**etic acid, which is rather a mouthful so EDTA is easier!

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Example of a polydentate ligand complex

Deducing the Charge on a Complex Ion

You can deduce the charge on a complex ion if you know the charges on the ligands and the oxidation state of the transition metal ion

Worked example

The three formulas shown are compounds of chromium(III). What are the charges on the complex ions shown?

 $I [Cr(H₂O)₆]Cl₃$

 $II [CrCl(H₂O)₅]Cl₂.H₂O$

 III [CrCl₂(H₂O)₄]Cl.2H₂O

A 0 , 0 , 0

 B 1+, 2+, 3+

 C 2+, 3+, 1+

 D 3+, 2+, 1+

Answer

The correct option is D

- **The water molecules do not contribute to the charge**
- The chloride ion, CI⁻, outside the square brackets must balance against the charge on the complex
- $[Cr(H₂O)₆]Cl₃ contains three chloride ions, so the charge on the complex is$ 3+
- $[CrCl(H₂O)₅]Cl₂H₂O contains two chloride ions outside the square bracket, so$ the charge on the complex is 2+
- $[CrCl₂(H₂O)₄]Cl.2H₂O contains one chloride ion outside the square bracket, so$ the charge on the complex is 1+

13.1.4 Catalytic & Magnetic Properties

Catalytic Properties

- **Transition metals are often used as catalysts in the elemental form or as compounds**
- \blacksquare The ability of transition metals to form more than one stable oxidation state means that they can accept and lose electrons easily
- This enables them to catalyse certain redox reactions. They can be readily oxidised and reduced again, or reduced and then oxidised again, as a consequence of having a number of different oxidation states of similar stability
- **There are two types of catalyst:**
	- A heterogeneous catalyst is in a different physical state (phase) from the reactants
		- The reaction occurs at active sites on the surface of the catalyst
		- An example is the use ofiron, Fe, in the Haber process for making ammonia

$$
N_2(g) + 3H_2(g) \approx 2NH_3(g)
$$

A homogeneous catalyst is in the same physical state (phase) as the reactants

Further Examples of transition metal catalysts

The hydrogenation or reduction of alkenes makes use of a nickel catalyst

$$
CH_2=CH_2(g)+H_2(g)\rightarrow CH_3CH_3(g)
$$

- The same reaction is used in the hydrogenation of vegetable oils to form polyunsaturated fats
- The decomposition of hydrogen peroxide is a common reaction in the study of chemical kinetics and uses manganese(IV) oxide as the catalyst

$$
2H_2O_2(g) \to 2H_2O(aq) + O_2(g)
$$

Catalytic converters

Catalytic converters are used in car exhaust boxes to reduce air pollution. They usually consist of a mixture of finely divided platinum and rhodium supported on a ceramic base

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Diagram of a catalyst on an inert support medium in a vehicle catalytic converter

- Carbon monoxide, nitrogen dioxide and unburnt hydrocarbons are sources of pollution in car exhaust \blacksquare
- \blacksquare The transition metal catalysts facilitate the conversion of these pollutants into less harmful products

 $2NO(g) + 2CO(g) \rightarrow N_2(g) + 2CO_2(g)$

$$
CH_3CH_2CH_3(g) + 5 O_2(g) \rightarrow 3CO_2(g) + 4H_2O(g)
$$

- Some ofthe transition metals are precious metals so they can be very expensive
- In order to minimise the cost and maximise the efficiency of the catalyst the following measures can be taken:
	- **Increasing the surface area of the catalyst**
	- **Coating an inert surface medium with the catalyst to avoid using large amounts of the catalyst**
- \blacksquare This is achieved by spreading the catalyst over a hollow matrix such as a honeycomb-like structure

Biological catalysts

- \blacksquare Many of the enzyme catalysed reactions in the body make use of homogeneous transition metal catalysts
- An example ofthis is haemoglobin, abbreviated to Hb, which transports oxygen around the blood

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Your notes

The structure of haemoglobin

The structure of haem

- The iron(II) ion is in the centre of a large heterocyclic ring called a **porphyrin**
- \blacksquare The iron has a coordination number of four, is square planar and can bind to one oxygen molecule
- \blacksquare The Hb molecule contains four porphyrin rings so each Hb can transport four oxygen molecules

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Magnetic Properties

- Materials are classified as diamagnetic, paramagnetic or ferromagnetic according to their behaviour when placed in an external magnetic field
- Diamagnetism is a property of all materials and produces a very weak opposition to an applied magnetic field
	- It arises from the repulsion of electrons to the applied magnetic field
- **Paramagnetism** only occurs in substances which have unpaired electrons
	- It produces magnetisation proportional to the applied field and in the same direction
	- **Transition metal complexes show paramagnetism**
- **Ferromagnetism** has the largest effect and produces magnetisation greater than the applied field

Diamagnetism

- The atoms of diamagnetic materials have paired electrons
- **Spinning electrons create a tiny magnetic dipole**
- \blacksquare The paired electrons orientate themselves so that the magnetic field they create opposes the external field
- **F** This result is a very weak repulsion force

Argon is diamagnetic with the electron configuration 1s² 2s² 2p⁶ 3s² 3p⁶

- Many molecules are diamagnetic since all the electrons are paired up in bonds
- It is very hard to demonstrate diamagnetism but it is possible by suspending a sample of the material from a sensitive force meter and lowering it into a strong horseshoe magnet - a slight change in the force should be seen

Paramagnetism

- Paramagnetic materials are attracted to an external magnetic field
- The unpaired electrons can be temporarily aligned to the magnetic field causing attraction into the field

The electrons in titanium are arranged in their orbitals as shown. The unpaired electrons can be temporarily aligned in an external field.

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- \blacksquare Most of the transition metals and their ions are paramagnetic as they have unpaired electrons
- Paramagnetism increases with the number of unpaired electrons, so it generally increases across the d-block up to a maximum with chromium and then decreases
- **EXTER** Zinc has no unpaired electrons so is not paramagnetic

3d 4s $[Ar]$ Sc Ti [Ar] ∧ ∧ V [Ar] ∧ ∧ Cr [Ar] Mn $[Ar]$ ∧ ∧ Fe [Ar] л Co $[Ar]$ ∧ Ni $[Ar]$ Cu $[Ar]$ Zn $[Ar]$

Orbital diagrams for the first row d-block showing the number of unpaired electrons increasing up to chromium

Ferromagnetism

- The metals iron, cobalt and nickel show the unusual property of ferromagnetism
- The alignment ofthe unpaired electrons in an external field in ferromagnetic materials can be retained so the material becomes permanently magnetised
- If ferromagnetic materials are heated and cooled in a magnetic field, the magnetic field of the electrons remains
- Magnetic regions within the metal that are aligned magnetically are know as **domains**
- **Banging or heating a permanent magnet will weaken the magnetism**

Ferromagnetic materials are used to make permanent magnets which produce characteristic magnetic field lines

Your notes

13.1.5 Colourin Transition Metals

Crystal Field Theory

Perception of colour

- **Most transition metal compounds appear coloured. This is because they absorb energy** corresponding to certain parts of the visible electromagnetic spectrum
- The colour that is seen is made up of the parts of the visible spectrum that aren't absorbed
- For example, a green compound will absorb all frequencies of the spectrum apart from green light, which is transmitted
- The colours absorbed are complementary to the colour observed

The colour wheel showing complementary colours in the visible light region of the electromagnetic spectrum

- Complementary colours are any two colours which are directly opposite each other in the colour wheel
	- For example, the complementary colour of red is green and the complementary colours of redviolet are yellow-green

Crystal Field Theory (CFT)

The crystal field theory is a model based on electrostatic point charges and is used to explain colour in transition metal compounds

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- \blacksquare In a transition metal atom, the five orbitals that make up the d-subshell all have the same energy. The term for this is degenerate
- However, when ligands are attached to a transition metal ion, the electric field formed by the lone pairs of electrons on the ligands repelthe electrons in the d-subshell causing the d-orbitals to splitin energy
- **The dative bonding from the ligands causes the five d orbitals to split into two sets**
- These two sets are not equal in energy and are described as being non-degenerate orbitals

- The lone pairs of the ligands repel the electrons in the x^2 -y² and z^2 orbitals of the metal ion **more** than they repel the electrons in the 3d_{vz}, 3d_{xz,} and 3d_{xv} orbitals
- This is because the 3d_{x2-v2} and 3d_{z2} orbitals line up with the dative bonds in the complex's octahedral shape
- The incoming ligands are attached to or approaching the central metal ion along the x, y and z axes, and the 3d $_{x2-y2}$ and 3d $_{z2}$ orbitals have lobes along these axes
- The electrons in these two orbitals are closer to the bonding electrons, so there is more repulsion
- This means that when the d orbitals split, the 3d $_{\rm X2-V2}$ and 3d $_{\rm z2}$ orbitals are at a slightly higher energy level than the other three
- The difference in energy between the non-degenerate d orbitals is labelled as *Δ*E

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Diagram showing the shapes and orientation of the five d-orbitals

Absorption of light

- When white light passes through a solution of aqueous nickel(II) sulfate an electron in the lower energy d-orbitals is excited and jumps up into the higher energy d-orbitals
- A photon of red light is absorbed and light of the complementary colour (green) is transmitted
- **This is why nickel(II) sulfate solution appears green**
- \blacksquare The energy of the separation is ΔE corresponding to a wavelength of about 630-700 nm

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Your notes

Q Examiner Tip

The colour wheel is given in Section 17 of the Data booklet, so there is no need to learn it. There are different splitting patterns possible but you are only required to know the octahedral splitting pattern discussed above.

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13.1.6 Factors Affecting Colour

Factors Affecting Colour

- The size of the splitting energy ΔE in the d-orbitals is influenced by the following four factors:
	- The size and type of ligands
	- **FIGE** The nuclear charge and identity of the metal ion
	- The oxidation state of the metal
	- The shape of the complex

The large variety of coloured compounds is a defining characteristic of transition metals

Size and type of ligand

- The nature of the ligand influences the strength of the interaction between ligand and central metal ion
- Ligands vary in their charge density \blacksquare
- The greater the charge density; the more strongly the ligand interacts with the metal ion causing greater splitting of the d-orbitals
- The further it is then shifted towards the region of the spectrum where it absorbs higher energy
- **As a result, a different colour of light is absorbed by the complex solution and a** different complementary colour is observed

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- This means that complexes with the same transition elements ions, but different ligands, can have different colours
	- For example, the $\rm [Cu(H_2O)_6]^{2+}$ complex has a **light blue** colour
	- Whereas the [Cu(NH₃)₄ (H₂O)₂]²⁺ has a **dark blue** colour despite the copper(II) ion having an oxidation state of +2 in both complexes

Ligand exchange of the waterligands by ammonia ligands causes a change in colour of the copper(II) complex solution

Ammonia has a greater charge density than water and so produces a larger splitin the d-orbitals

Graph showing the replacement of the water molecules with four ammonia molecules causes a shift in maximum absorbance towards shorter wavelength

The nuclear charge

- The strength of the attraction between the metal ion and lone pairs of electrons from the ligand can vary depending on the **effective nuclear charge** on the metal ion
- For example, aqueous Mn(II) and Fe(III) have the same electronic configuration:

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$[Ar]$ 3d⁵

- $Mn(II)$ (Z=25) absorbs in the green region of the spectrum so appears pink
- The higher effective nuclear charge on Fe(III) (Z= 26) causes a stronger interaction with the ligands, so it absorbs in the higher energy blue part of the spectrum and appears yellow/orange in colour

Oxidation state

- When the same metal is in a higher oxidation state that will also create a stronger interaction with the ligands
- If you compare iron(II) and iron (III):
	- $[Fe(H₂O)₆]²⁺$ absorbs in the red region and appears green
	- But, [Fe(H₂O)₆]³⁺ absorbs in blue region and appears orange

Shape

- The change of colour in a complex is also partly due to the change in coordination number and geometry of the complex ion
- The splitting energy, ΔE, ofthe d-orbitals is affected by the relative orientation ofthe ligand as well as the d-orbitals
- **EXT** Changing the coordination number generally involves changing the ligand as well, so it is a combination of these factors that alters the strength of the interactions

The Spectrochemical Series

- The Japanese chemist, R. Tuschida, proposed a ranking of ligands base on their ability to separate the two sets of d-orbitals
- This is known as the spectrochemical series
- \blacksquare The higher the charge density of the ligand; the more strongly it influences the splitting of the dorbitals so the greater the energy difference between them

Table showing the spectrochemical series for common ligands

- You can see at either end of the series lies jodide jons, I⁻, and carbon monoxide, CO
- \blacksquare Iodide ions are large (think how many electron shells they have) so they have a relative low charge density and produce the weakest electric field so the separation energy ofthe d-orbitals is the smallestin the series
- Chloride ions, CI⁻, on the other hand are smaller, have a higher charge density and consequently produce a large separation energy
- \blacksquare However, size is not the only factor as carbon monoxide and cyanide ions produce a larger splitting due to complex interactions involving the pi bonds present in those molecules; those interactions occur when electrons in the d-orbitals of the transition metal interact with electrons in the p-orbitals of the ligands
- You should be able to see why adding ammonia to aqueous copper(II) ions results in a darker blue complex
	- Ammonia is a stronger ligand than water so the separation energy is larger and the wavelength of absorbed light shorter
	- Shorter wavelength means moving towards to bluer higher energy end ofthe visible spectrum

Q Examiner Tip

You do not need to learn the spectrochemical series as it is given in the data booklet in section 15. A list of polydentate ligands is also given in the data booklet in section 16.

