UNIT 9 – METALS AND THEIR COMPOUNDS

UNIT 9

METALS AND THEIR COMPOUNDS

Teacher Version

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Contents
a) Classification – Metals, Non-metals and Metalloids
b) Physical Properties of Metals
c) Chemical Properties of Metals
d) Extraction of Metals
e) Uses of Metals and Metalloids
f) Properties and Reactions of Metal Compounds
g) Qualitative Analysis Part 4

Key words: metal, non-metal, metalloid, malleability, ductility, lustre, sonority, alloy, bronze, brass, steel, solder, malleable, ductile, sonorous, lustrous, ferromagnetic, s-block metal, p-block metal, d-block metal, transition metal, rusting, reactivity series, metal displacement reaction, blast furnace, basic oxygen process, complex ion, ligand

Units which must be completed before this unit can be attempted:
Unit 1 – Atoms and the Periodic Table
Unit 2 – Particles, Bonding and Structure
Unit 3 – Amount of Substance and Measurement
Unit 4 – Introduction to Physical Chemistry
Unit 5 – Acids, Bases and Salts
Unit 6 – Redox Reactions
Unit 8 – Solubility and Precipitation Reactions

Estimated Teaching Time: 10 hours
**UNIT 9 – METALS AND THEIR COMPOUNDS**

**UNIT 9 SUMMARY AND SYLLABUS REFERENCE**

<table>
<thead>
<tr>
<th>Lesson</th>
<th>Title and Syllabus Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Classification of Metals</td>
</tr>
<tr>
<td></td>
<td><strong>CA3ci metallic Bonding - factors influencing its formation</strong> (Factors should include: atomic radius, ionization potential, and number of valence electrons, type of specific packing not required); <strong>CA6bii alloys; ISA4.1 particulate nature of matter</strong> (elements - metals and non-metals (1st to 20th elements in the periodic table)); <strong>ISA10.1 classification of materials</strong> (classification of materials into metals, semi-metals (metaloids), and non-metals); <strong>CC8dii copper – properties</strong>; <strong>ISA10.2 uses of metals and semi-metals</strong></td>
</tr>
</tbody>
</table>

| 2      | Physical Properties of Metals |
|        | **CA2c elements of the first transition series** (physical properties of the elements - physical properties should include: physical states, metallic properties and magnetic properties); **CA3cii properties of metals** (typical properties including conductivity - malleability, ductility demonstrated using metals like Mg, Zn, Sn, Fe); **CC8a properties of sodium**; **CC8b properties of calcium**; **CC8dii copper – properties**; **ISA10.2 uses of metals and semi-metals** (physical properties of metals and semi-metals under conductivity, luster, malleability, ductility, sonority, density, melting point and tensile strength) |

| 3      | Chemical Properties of Metals (s and p block) |
|        | **CA7a definitions of acids and bases** (effects of acids and bases on metals); **CA7b chemical properties of acids and bases** (balanced chemical equations of all reactions); **CC8a properties of sodium**; **CC8b properties of calcium**; **CC8c reactivity of aluminium with air, water and acids**; **ISA7.2 chemical properties of acids and bases** |

| 4      | Chemical Properties of Metals (d block) |
|        | **CA2c elements of the first transition series** (chemical reactivities of the elements, other properties should include variable oxidation states, catalytic abilities); **CA7a definitions of acids and bases** (effects of acids and bases on metals); **CA7b chemical properties of acids and bases** (balanced chemical equations of all reactions); **CC8b reactivity of iron with air, water and acids**; **CC8dii copper – properties**; **ISA7.2 chemical properties of acids and bases** |

| 5      | Reactivity Series of Metals |
|        | **CA2c elements of the first transition series** (chemical reactivities of the elements); **CA4aii chemical equations** (displacement of metal ions); **CC8a properties of sodium**; **CC8b properties of calcium**; **CC8c reactivities of aluminium and iron with air, water and acids**; **CC8dii copper – properties**; **ISA7.2 chemical properties of acids and bases** |

| 6      | Extraction of Metals |
|        | **CA12bii extraction of metals - Al, Fe, Au and Sn** (raw materials, processing, main products, byproducts, recycling) |

| 7      | Uses of Metals; Compounds of Sodium and Calcium |
|        | **CA12bii extraction of metals** (uses of the metals); **CA12bii alloys** (common alloys of Cu, Al, Pb, and Fe and their uses); **CC8a properties and uses of compounds of sodium** (NaCl, NaOH, NaNO₃ and Na₂SO₄); **CC8b properties and uses of compounds of calcium** (CaO, CaSO₄, CaCl₂, Ca(OH)₂); **CC8dii copper – uses**; **ISA10.2 uses of metals and semi-metals** (uses of the following elements: Al, Cu, Fe, Au, application of semi-metals); **ISA10.3 alloys** (examples of alloys and their constituent elements (steel, bronze, brass), uses of alloys, advantages of alloys in manufacture of certain household items) |

| 8      | Compounds of D-block Metals |
|        | **CA2c elements of the first transition series** (electronic configuration, chemical reactivities of the compounds, other properties should include formation of coloured compounds, complex ion formation); **CA13cii characteristic tests of the following cations with dilute NH₃(aq): Ca²⁺, Pb²⁺, Cu²⁺, Fe³⁺; Fe⁺³, Al³⁺ and Zn²⁺; CC7b water and solution** (test for water will be required); **CC8dii copper – uses of compounds** (the compounds must be limited to CuSO₄, CuO and CuCl₂) |

| 9      | Qualitative Analysis – Complex Formation Reactions |
|        | **CA2c elements of the first transition series** (chemical reactivities of the compounds, other properties should include complex ion formation); **CA13cii characteristic tests of the following cations with dilute NH₃(aq) and NaOH: Ca²⁺, Pb²⁺, Cu²⁺, Fe³⁺, Fe⁺³, Al³⁺ and Zn²⁺, confirmatory tests for these cations** |

| 10     | Unit 9 Revision and Summary |
UNIT 9 – METALS AND THEIR COMPOUNDS

Lesson 1 – What are metals?

a) Classification – Metals, Non-metals and Metalloids

Summary Activity 1.1: What can you remember about metals?
- What is a metal?
- Describe the structure and bonding in a typical metal
- What are the properties of a typical metal?
- What is a non-metal?
- Describe the structure and bonding in a typical non-metal
- Why are some elements metals and others non-metals?
- What is an alloy? How many alloys can you name?

- A metal is a substance (usually an element) which contains metallic bonding
- Metallic bonding is the attraction between a lattice of cations and a sea of delocalised electrons
- Metallic bonding is quite strong so metals often have high melting points; the delocalised electrons make them good conductors of electricity; the metal ions can move past each other without disrupting the metallic bonding, so metals tend to be malleable and ductile
- A non-metal is an element which contains covalent bonding
- Non-metals either have simple molecular structures - small groups of atoms held together by covalent bonds (called molecules) and weak Van der Waal’s forces between the molecules, or giant covalent structures (lattice of atoms held together by covalent bonds
- Electropositive atoms do not hold on to their electrons strongly and allow their valence electrons to be delocalised; electronegative atoms hold on to their electrons strongly and form covalent bonds instead
- An alloy is a mixture of atoms held together by metallic bonds; the major component of the mixture must be a metal (eg brass, bronze, steel, solder)

(i) Metals

• Metals are elements which contain metallic bonding

• In metallic bonding, the atoms lose some or all of their outer shell (valence) electrons to form a lattice of cations and a sea of delocalised electrons; the delocalised electrons and the cations are attracted to each other and this holds the structure together

• Metallic bonding is preferred when all of the bonding atoms are electropositive, and hence do not hold on tightly to their valence electrons; it is therefore most commonly found in the elements of atoms on the left-hand side, and at the bottom, of the Periodic Table (bonding in elements becomes less metallic and more covalent from left to right across a Period, and more metallic and less covalent from top to bottom in a Group)

• In some cases, two or more different atoms can form a metallic bond; usually if both atoms are electropositive (ie come from metals; eg Cu and Zn); electropositive atoms can also form metallic bonds with atoms which are not electropositive (ie come from non-metals), provided that the electropositive atom is more abundant (eg Fe and C); mixtures which contain metallic bonds are known as alloys; alloys are not compounds because they do not have a fixed composition
(ii) **Non-metals**

- **Non-metals** are elements which do not contain metallic bonds; the vast majority of non-metals contain covalent bonding.

- In elements, covalent bonding is preferred to metallic bonding if the atoms are not electropositive enough to give up their valence electrons; it is therefore most commonly found in the elements of atoms on the right-hand side, and at the top of the Periodic Table (bonding in elements becomes less metallic and more covalent from left to right across a Period, and more metallic and less covalent from top to bottom in a Group); elements containing covalent bonds are called **non-metals**

(iii) **Metalloids**

- **Metalloids** or **semi-metals** are substances which contain bonding intermediate in character between metallic and covalent; metalloids can be elements, compounds or mixtures, but the term is most commonly used to describe elements; often these substances display metallic properties at higher temperatures and non-metallic properties at lower temperatures; they are found in between the metals and the non-metals in the Periodic Table.

- The following diagram shows the location of the metals, non-metals and metalloids in the Periodic Table; note the trend from metal to metalloid to non-metal across a Period from right to left and up a Group from bottom to top: (white = metal, grey = metalloid; black = non-metal)

<table>
<thead>
<tr>
<th>I</th>
<th>II</th>
</tr>
</thead>
<tbody>
<tr>
<td>Li</td>
<td>Be</td>
</tr>
<tr>
<td>Na</td>
<td>Mg</td>
</tr>
<tr>
<td>K</td>
<td>Ca</td>
</tr>
<tr>
<td>Rb</td>
<td>Sr</td>
</tr>
<tr>
<td>Cs</td>
<td>Ba</td>
</tr>
<tr>
<td>Fr</td>
<td>Ra</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>III</th>
<th>IV</th>
<th>V</th>
<th>VI</th>
<th>VII</th>
<th>0</th>
</tr>
</thead>
<tbody>
<tr>
<td>B</td>
<td>C</td>
<td>N</td>
<td>O</td>
<td>F</td>
<td>Ne</td>
</tr>
<tr>
<td>Al</td>
<td>Si</td>
<td>P</td>
<td>S</td>
<td>Cl</td>
<td>Ar</td>
</tr>
<tr>
<td>Ti</td>
<td>V</td>
<td>Cr</td>
<td>Mn</td>
<td>Fe</td>
<td>Co</td>
</tr>
<tr>
<td>Zr</td>
<td>Nb</td>
<td>Mo</td>
<td>Tc</td>
<td>Ru</td>
<td>Rh</td>
</tr>
<tr>
<td>Hf</td>
<td>Ta</td>
<td>W</td>
<td>Re</td>
<td>Os</td>
<td>Ir</td>
</tr>
</tbody>
</table>

(iv) **Further classification of metals**

- Metals can be further classified into s-block, p-block, d-block or f-block metals based on their electronic configuration and consequent position in the Periodic Table:

- Metals made from atoms containing valence electrons in s-orbitals only are called s-block metals (eg Na, Ca)

- Metals made from atoms containing valence electrons in s and d-orbitals only are called d-block metals (eg Fe, Cu, Au)

- Metals made from atoms containing valence electrons in s and p-orbitals only are called p-block metals (eg Al)
Test your knowledge 1.2: Classifying metals, non-metals and metalloids

(a) Name two s-block metals
(b) Name two p-block metals
(c) Name two d-block metals
(d) Name two metalloids
(e) Name two non-metals
(f) Why does metallic character decrease across a Period?
(g) Why does metallic character increase down a Group?

(a) Eg sodium, calcium, magnesium, potassium
(b) Eg aluminium, tin, lead
(c) Eg copper, iron, zinc
(d) Eg boron, silicon
(e) Eg oxygen, bromine, neon
(f) Electronegativity increases across a Period, so the attraction to bonding electrons increases, so atoms become less likely to allow bonding electrons to delocalise
(g) Electronegativity decreases down a Period, so the attraction to bonding electrons decreases, so atoms become more likely to allow bonding electrons to delocalise

**Lesson 2 – What are the physical properties of metals?**

**b) Physical Properties of Metals**

- Most physical properties of metals are caused by the metallic bonding; in most cases, variations in the physical properties of the metals are caused by variations in the strength of metallic bonding:

(i) **Strength of Metallic Bonding**

- The strength of metallic bonding depends on:
  - The charge on the cations: the higher the charge on the cations, the stronger the metallic bonding
  Eg Mg$^{2+}$ is smaller than Na$^+$ so the metallic bonding is stronger in Mg than in Na
  - The size of the cations: the smaller the size of the cations, the stronger the metallic bonding
  Eg Na$^+$ is smaller than K$^+$ so the metallic bonding is stronger in Na than in K
  - The number of delocalised electrons per atom: the greater the number of delocalised electrons, the stronger the metallic bonding; the number of delocalised electrons per atom is equal to the charge on the cation
    Eg Mg gives up two electrons, Na gives up only one, so the metallic bonding is stronger in Mg than in Na
  - The block of the Periodic Table; the bonding in d-block metals is much stronger than the bonding in s and p-block metals (the reasons for this will not be discussed in this course)

- Generally speaking, the further to the right, and the higher up, the element is found in the Periodic Table, the stronger the metallic bonding
(ii) Physical Properties of Metals

- Some physical properties are common to all metals:
  - **Electrical conductivity**: all metals contain delocalised electrons, which can move when a potential difference is applied; all metals are therefore good conductors of electricity
  - **Thermal conductivity**: the delocalised electrons carry heat energy with them, which can therefore move rapidly through the structure as the electrons move; all metals are therefore good conductors of heat
  - **Malleability**: in metals, the attraction between the delocalised electrons and the cations does not depend on the position of the cations; it is therefore possible to move the cations around without having a significant impact on the bonding; this means that metals can be easily bent into different shapes; all metals are therefore malleable
  - **Ductility**: because metals can be bent into different shapes, they can also be turned into wires; this property is known as ductility; all metals are therefore ductile, especially those with stronger metallic bonding
  - **Lustre**: the delocalised electrons in metals reflect light back towards its source; this causes metals to have a characteristic shiny appearance known as lustre
  - **Sonority**: the delocalised electrons in metals can also transport sound energy efficiently through the solid, which means that the sound wave can travel through the metal with little loss of energy; the ability of a solid to make a ringing, resonant sound when hit is known as sonority; most metals are sonorous (although high density metals are less sonorous)

- Some other physical properties of metals depend on the strength of the metallic bonding, which varies significantly from metal to metal:
  - **Melting point**: the melting point of metals depends on the strength of the metallic bonding; it is generally high as metallic bonding is generally strong, so a lot of energy is needed to break up the metallic lattice; however the strength of the metallic bonding varies widely between metals and so does the melting point. Group I metals have weak metallic bonding and a low melting point; d-block metals have strong metallic bonding and a high melting point
  - **Tensile strength**: because metals are malleable, they can absorb large quantities of energy without break and so tend to have a high tensile strength; however the tensile strength depends on the strength of the metallic bonding, which varies widely between metals; Group I metals have weak metallic bonding and a tensile strength; d-block metals have strong metallic bonding and a high tensile strength

- Some properties of metals do not result from either the delocalised electrons or the strength of the metallic bonding:
  - **Magnetic Properties**: some metals have their delocalised electrons arranged in such a way that they can align themselves to a magnetic field and attract a magnet; the ability to do this is known as ferromagnetism; only Fe, Co and Ni are ferromagnetic; ferromagnetism is caused by the presence of unpaired electrons in the atoms: Fe, Co and Ni have electronic configurations d⁶, d⁷ and d⁸ respectively, which means that they have four, three and two unpaired electrons respectively; however not all atoms with unpaired electrons are ferromagnetic and the full causes of ferromagnetism will not be discussed in this course
  - **Density**: density is mass per unit volume; the density of a metal therefore depends on the atomic mass (the higher the atomic mass the higher the density) and the atomic size (the larger the atomic size the lower the density); the density also depends to an extent on the type of crystal structure adopted by the metal, as some crystal structures are more compact than others; atomic mass increases across a Period and atomic size decrease across a period; the density of metals therefore increases across a Period; atomic mass increases down a group but so does atomic size; generally, the increase in atomic mass down a group is more significant so density tends to increase down a group; the metallic elements with the highest density are therefore to be found at the bottom right of the Periodic Table
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- **Summary of properties of metals:**

<table>
<thead>
<tr>
<th>Property</th>
<th>s-block and p-block</th>
<th>d-block</th>
</tr>
</thead>
<tbody>
<tr>
<td>Electrical and Thermal Conductivity</td>
<td>High</td>
<td>High</td>
</tr>
<tr>
<td>Melting Point</td>
<td>Low-medium</td>
<td>High</td>
</tr>
<tr>
<td>Tensile strength</td>
<td>Low-medium</td>
<td>High</td>
</tr>
<tr>
<td>Malleability</td>
<td>High</td>
<td>High</td>
</tr>
<tr>
<td>Ductility</td>
<td>Medium</td>
<td>High</td>
</tr>
<tr>
<td>Lustre</td>
<td>High</td>
<td>High</td>
</tr>
<tr>
<td>Sonority</td>
<td>High</td>
<td>High</td>
</tr>
<tr>
<td>Magnetism</td>
<td>Not-magnetic</td>
<td>Fe, Co, Ni magnetic</td>
</tr>
<tr>
<td>Density</td>
<td>Low-medium</td>
<td>Medium-high</td>
</tr>
</tbody>
</table>

Test your knowledge 2.1: Describing physical properties of metals

(a) Why do all metals conduct electricity?
(b) Why are all metals malleable?
(c) Why does magnesium have a higher melting point than sodium?
(d) Why does potassium have a lower melting point than potassium?
(e) Why is aluminium stronger than magnesium?
(f) Why is iron denser than aluminium?
(g) Why is iron more magnetic than copper?

(a) Delocalised electrons are free to move
(b) Cations can move around without breaking metallic bonds
(c) Mg\(^{2+}\) is smaller than Na\(^+\) and is more highly charged, so it attracts delocalised electrons more strongly, so the metallic bonds are stronger and more energy is needed to break them
(d) K\(^+\) is larger than Na\(^+\), so it attracts delocalised electrons less strongly, so the metallic bonds are weaker and less energy is needed to break them
(e) Al\(^{3+}\) is smaller than Mg\(^{2+}\) and is more highly charged, so it attracts delocalised electrons more strongly, so the metallic bonding is stronger
(f) Iron has a larger atomic mass than aluminium
(g) Iron has more unpaired electrons than copper

Lesson 3 – How do s and p-block metals react with air, water and acids?

Summary Activity 3.1: What can you remember about redox reactions of metals?

- Give one example of a reaction between a metal and oxygen
- Give one example of a reaction between a metal and an acid
- Give one example of a metal displacement reaction
- In each reaction, what is oxidised and what is reduced?

- Eg 2Mg + O\(_2\) → 2MgO or 4Al + 3O\(_2\) → 2Al\(_2\)O\(_3\) (metal oxidised, O reduced)
- Eg Mg + 2H\(^+\) → Mg\(^{2+}\) + H\(_2\) or Zn + 2H\(^+\) → Zn\(^{2+}\) + H\(_2\) (metal oxidised, H\(^+\) reduced)
- Eg Zn + Cu\(^{2+}\) → Zn\(^{2+}\) + Cu (Zn oxidised, Cu\(^{2+}\) reduced)
c) Chemical Properties of Metals

(i) s-block metals (Na and Ca)

- s-block metals are electropositive and do not hold on to their electrons strongly; the valence electrons are easily lost during chemical reactions and the metal is oxidised; most s-block metals react rapidly with air, water and acids as a result

- s-block metals only form one stable cation, corresponding to the loss of all valence electrons; Group 1 metals form +1 ions only (eg Na\(^+\)) and Group 2 metals form +2 ions only (eg Ca\(^{2+}\))

- Chemical reactivity increases as electropositivity increases; reactivity therefore decreases down a Group and Group 1 metals are more reactive than Group 2 metals

- With air, the elements will react to form oxides (containing O\(^2-\)) (eg CaO)
  - although barium and all Group 1 metals except lithium can also form peroxides (containing O\(^2-\)) (eg Na\(_2\)O\(_2\))
  - these reactions are violent and exothermic once the metal is ignited
  - the more reactive s-block metals can ignite spontaneously, most s-block metals tarnish rapidly on exposure to air (they form an oxide layer at the surface of the metal which makes it less shiny)
  - 4Na(s) + O\(_2\)(g) \rightarrow 2Na\(_2\)O(s); 2Na(s) + O\(_2\)(g) \rightarrow Na\(_2\)O\(_2\)(s); 2Ca(s) + O\(_2\)(g) \rightarrow 2CaO(s)
  - these reactions are redox reactions; the metal is oxidised and the oxygen is reduced; most s-block metals are stored under oil to prevent this reaction from taking place

- With water, the elements will react to form hydroxides (containing OH\(^-\)) and hydrogen (H\(_2\))
  - all s-block elements except Be and Mg react rapidly with water
  - with the most reactive metals the reaction is very explosive; this is another reason why the s-block metals are stored under oil; the metal fizzes rapidly as hydrogen is given off
  - 2Na(s) + 2H\(_2\)O(l) \rightarrow 2NaOH(aq) + H\(_2\)(g); Ca(s) + 2H\(_2\)O(l) \rightarrow Ca(OH)\(_2\)(aq) + H\(_2\)(g)
  - these reactions are redox reactions; the metal is oxidised and the hydrogen in the water is reduced
  - watch these reactions on: [www.youtube.com/watch?v=fAFrzlyrqCU](https://www.youtube.com/watch?v=fAFrzlyrqCU)
  - with calcium, the solution quickly becomes cloudy; Ca(OH)\(_2\) is only sparingly soluble in water and so the solution becomes saturated quickly; any more Ca(OH)\(_2\) formed cannot dissolve and will form as a solid

- With acids, the reaction is even more vigorous and is dangerously explosive, especially with the Group 1 metals; the metal is oxidised and the H\(^+\) ion in the acid is reduced (to hydrogen gas); a salt is also produced (see Topic 6 – Redox Reactions)
  - 2Na(s) + 2HCl(aq) \rightarrow 2NaCl(aq) + H\(_2\)(g); Ca(s) + 2HCl(aq) \rightarrow CaCl\(_2\)(aq) + H\(_2\)(g)

(ii) p-block metals (Al, Sn and Pb)

- p-block metals are much less reactive than s-block metals; they are less electropositive and hold onto their electrons more strongly than s-block elements; of the common p-block metals, Al is the most reactive, followed by Sn and then lead

- aluminium only forms Al\(^{3+}\) ions, corresponding to the loss of all valence electrons; tin can lose all of its electrons to form Sn\(^{4+}\); tin can also lose its p-electrons only (not its s-electrons) to form Sn\(^{2+}\); lead generally loses its p-electrons only to form Pb\(^{2+}\)
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• Although Al is considered a reactive metal, it shows almost no reaction with air or water; the reaction is much slower than expected because a small quantity of aluminium at the surface can react to form \( \text{Al}_2\text{O}_3 \); this oxide forms a very stable layer on the surface of the metal which prevents any further oxygen or water from reacting with the aluminium.

• Aluminium powder can react with air if heated, and aluminium can react with steam if heated strongly:

\[
\begin{align*}
4\text{Al}(s) + 3\text{O}_2(g) & \rightarrow 2\text{Al}_2\text{O}_3(s) \\
4\text{Al}(s) + 6\text{H}_2\text{O}(g) & \rightarrow 2\text{Al}_2\text{O}_3(s) + 3\text{H}_2(g)
\end{align*}
\]

The aluminium is oxidised; with oxygen, the O is reduced; with water, the H is reduced.

• Most p-block metals, including Al, will react slowly with acids to give a salt and hydrogen:

\[
\begin{align*}
2\text{Al}(s) + 6\text{HCl}(aq) & \rightarrow \text{AlCl}_3(aq) + 3\text{H}_2(g)
\end{align*}
\]

The Al is oxidised and the \( \text{H}^+ \) in the acid is reduced (see Topic 6 – Redox Reactions).

Test your knowledge 3.2: Describing chemical properties of s and p-block metals

(a) Write equations for the reactions of potassium and calcium with water. Explain what you would observe, state the type of reaction taking place and explain why potassium reacts with water more quickly than calcium.

(b) Write equations for the reactions of sodium and calcium with oxygen. What is the difference between the two products?

(c) Why are the alkali metals stored under oil?

(d) Write equations for the reactions of magnesium and aluminium with hydrochloric acid. Explain the type of reaction taking place and explain why magnesium reacts with acids more quickly than aluminium.

(e) Explain why aluminium shows little reaction with air or water despite being considered a reactive metal.

Lesson 4 - How do d-block metals react with air, water and acids?

Summary Activity 4.1: What can you remember about d-block metals?

- Why does iron rust?
- How can rusting be prevented?
- How is copper purified?
- What are the electronic configurations of Cu and Fe?
- What are the electronic configurations of \( \text{Cu}^{2+} \), \( \text{Fe}^{2+} \) and \( \text{Fe}^{3+} \)?
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- Iron rusts when is is oxidised by O₂ and H₂O to Fe(OH)₃; the rust does not stick to the surface to the iron but flakes off, exposing the iron underneath to further reaction
- By painting, greasing, galvanising, sacrificial protection with a more reactive metal
- By electrolysis of CuSO₄ using copper electrodes; the copper on the impure anode dissolves (Cu → Cu²⁺ + 2e and pure copper is deposited at the cathode: Cu²⁺ + 2e → Cu
- Cu: 1s²2s²2p⁶3s²3p⁶4s²3d¹⁰; Fe: 1s²2s²2p⁶3s²3p⁶4s²3d⁶
- Cu²⁺: 1s²2s²2p⁶3s²3p⁶3d³; Fe²⁺: 1s²2s²2p⁶3s²3p⁶3d⁶; Fe³⁺: 1s²2s²2p⁶3s²3p⁶3d⁵

(iii) d-block metals (Fe, Cu and Au)

- d-block metals are generally less reactive than p-block metals, although their electropositivity and reactivity does vary considerably; of the common d-block metals, Zn is the most reactive, followed by Fe; Cu is unreactive; Ag and Au are very unreactive
- d-block metals always lose their valence s-electrons; most d-block metals have two s-electrons so form 2+ ions (eg Fe²⁺) although atoms in columns 4 and 9 of the d-block only have one s-electron (eg Cu has the electronic configuration [Ar]3d¹⁰4s¹) so can form +1 ions (eg Cu⁺, Au⁺); the d-electrons in the outer shell, however, can behave as valence electrons in certain circumstances; this means that many atoms in the d-block can form several different ions, depending on how many d-electrons they lose:
  - Cu has the electronic configuration [Ar]3d¹⁰4s¹; it loses its valence s-electron to form Cu⁺; ([Ar]3d¹⁰); it usually also loses one valence d-electron to form Cu²⁺ ([Ar]3d⁹)
  - Fe has the electronic configuration [Ar]3d⁶⁴s²; it loses its valence s-electrons to form Fe²⁺ ([Ar]3d⁶); it can also lose one valence d-electron to form Fe³⁺ ([Ar]3d⁵)

Transition metals can therefore have several stable oxidation states; it is common for d-block metals to form stable ions in which the d-orbitals are partially filled (ie ions which have 1 – 9 electrons in their d-orbitals); a metal which forms at least one stable ion with a partially filled d-orbital is called a transition metal; Cu, Fe and Au are all transition metals

- The ability of the d-electrons in transition metals to be lost and gained again results in transition metals (and their compounds) being good catalysts:
  - Fe is used as a catalyst in the Haber Process (the manufacture of ammonia: N₂ + 3H₂ → 2NH₃)
  - Ni is used as a catalyst in the hydrogenation of alkenes (C₂H₄ + H₂ → C₂H₆)

- Reaction with air:
  - Most d-block metals show almost no reaction with air or water at normal temperatures, either because they are unreactive or because they form a stable oxide layer
  - Iron can react with air, when heated, to form iron (II) oxide and iron (III) oxide; copper can also react with air when heated to form copper (II) oxide; 2Fe(s) + O₂ → 2FeO; 4Fe(s) + 3O₂ → 2Fe₂O₃; 2Cu + O₂ → 2CuO
  - Gold does not readily react with oxygen; this is why gold remains shiny for a long time
UNIT 9 – METALS AND THEIR COMPOUNDS

• Reaction with water:
  - Iron does not readily react with water in the absence of air but in the presence of air, iron can react with both oxygen and water to form rust: \(4\text{Fe} + 6\text{H}_2\text{O} + 3\text{O}_2 \rightarrow 4\text{Fe(OH)}_3\); unlike most other oxides and hydroxides of d-block elements, rust does not provide protect the iron surface and so iron will continue to rust if not protected.
  - Copper is less reactive than iron and also forms a protective oxide layer; it does not react with water at all at room temperature but if it exposed to hot water and air for a period of time it can form a green layer (copper carbonate) on its surface; this layer also provides protection to the copper.
  - Gold does not readily react with water.

• Reaction with acids:
  - Some d-block metals, including Fe and Zn will react slowly with acids to give a salt and hydrogen: \(\text{Fe(s)} + 2\text{HCl(aq)} \rightarrow \text{FeCl}_2(\text{aq}) + \text{H}_2(\text{g})\); the Fe is oxidised and the \(\text{H}^+\) in the acid is reduced; this can be a useful way of preparing salts of d-block metals.
  - Cu and Au do not react readily with dilute acids.

Test your knowledge 4.2: Describing chemical properties of d-block metals
(a) Explain why d-block metals often have more than one stable oxidation state
(b) What is the difference between a d-block metal and a transition metal?
(c) Explain why transition metals are good catalysts and give an example of a reaction catalysed by a transition metal.
(d) Write the equation for the rusting of iron
(e) Explain why copper is used in water pipes and why gold makes good jewellery.
(f) Write an equation for the reaction between iron and sulphuric acid.

(a) because they do not always lose all of their valence d-electrons; the number of electrons they lose can vary.
(b) d-block metal: has valence s electrons and d electrons but no p-electrons; transition metal: forms at least one stable ion with a partially filled d-orbital (not all d-block metals are transition metals).
(c) they can change their oxidation state so can accept and donate electrons; eg Fe in Haber process.
(d) \(4\text{Fe} + 6\text{H}_2\text{O} + 3\text{O}_2 \rightarrow 4\text{Fe(OH)}_3\)
(e) Copper does not react with water; gold does not react with air or water so stays shiny for a long time.
(f) \(\text{Fe} + \text{H}_2\text{SO}_4 \rightarrow \text{FeSO}_4 + \text{H}_2\)
Lesson 5 – How can we compare the reactivity of different metals?

**Summary Activity 5.1: Which types of redox reaction involve metals?**

- Write an equation to show how sodium reacts with water; indicate what is oxidised and what is reduced; why does zinc not react with water?
- Write an equation to show how zinc reacts with hydrochloric acid; indicate what is oxidised and what is reduced; why does copper not react with hydrochloric acid?
- Write an equation to show how zinc reacts with copper sulphate; indicate what is oxidised and what is reduced; why does copper not react with zinc sulphate?
- Write an equation to show how zinc oxide reacts with carbon; indicate what is oxidised and what is reduced; why does aluminium oxide not react with carbon?
- Write an equation to show how copper oxide reacts with hydrogen; indicate what is oxidised and what is reduced; why does zinc oxide not react with hydrogen?
- What is the general name given to all of these reactions?

\[
2\text{Na} + 2\text{H}_2\text{O} \rightarrow 2\text{NaOH} + \text{H}_2; \text{Na oxidised from 0 to } +1; \text{H reduced from } +1 \text{ to } 0; \text{zinc less reactive than sodium and cannot displace hydrogen from water}
\]

\[
\text{Zn} + 2\text{HCl} \rightarrow \text{ZnCl} + \text{H}_2; \text{Zn oxidised from 0 to } +2; \text{H reduced from } +1 \text{ to } 0; \text{copper less reactive than zinc and cannot displace hydrogen from acids}
\]

\[
\text{Zn} + \text{CuSO}_4 \rightarrow \text{Cu} + \text{ZnSO}_4; \text{Zn oxidised from 0 to } +2; \text{Cu reduced from } +2 \text{ to } 0; \text{copper less reactive than zinc so cannot displace zinc from its compounds}
\]

\[
\text{ZnO} + \text{C} \rightarrow \text{Zn} + \text{CO}; \text{C oxidised from 0 to } +2; \text{Zn reduced from } +2 \text{ to } 0; \text{aluminium more reactive than carbon so cannot be displaced from its compounds by carbon}
\]

\[
\text{CuO} + \text{H}_2 \rightarrow \text{Cu} + \text{H}_2\text{O}; \text{H oxidised from 0 to } +1; \text{Cu reduced from } +2 \text{ to } 0; \text{zinc more reactive than hydrogen so cannot be displaced from its compounds by hydrogen}
\]

- These are all examples of metal displacement reactions

(iv) **The Reactivity Series of Metals**

- The reactivity of different metals to oxygen, metals and water varies widely; it is possible to rank the reactivity of the different metals (and some non-metals); this is known as the reactivity series; the metals at the top are the most reactive:

<table>
<thead>
<tr>
<th>Metal</th>
<th>Reactivity</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sodium</td>
<td>Reacts with cold water</td>
</tr>
<tr>
<td>calcium</td>
<td>Reacts with cold water</td>
</tr>
<tr>
<td>magnesium</td>
<td>Reacts with acids but not water</td>
</tr>
<tr>
<td>aluminium (carbon)</td>
<td>Reacts with acids but not water</td>
</tr>
<tr>
<td>zinc</td>
<td>Reacts with acids but not water</td>
</tr>
<tr>
<td>iron</td>
<td>Reacts with acids but not water</td>
</tr>
<tr>
<td>tin</td>
<td>Reacts with acids but not water</td>
</tr>
<tr>
<td>lead</td>
<td>No reaction with acids or water</td>
</tr>
<tr>
<td>copper</td>
<td>No reaction with acids or water</td>
</tr>
<tr>
<td>gold</td>
<td></td>
</tr>
</tbody>
</table>

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UNIT 9 – METALS AND THEIR COMPOUNDS

• The higher up the reactivity series a metal is, the more reactive it is and the more vigorously and quickly it will react with water, acids, oxygen and other non-metals; in addition, the more reactive the metal is, the harder it is to extract the metal from its compounds.

• More reactive metals can displace less reactive metals from their compounds; for example, zinc will displace copper from copper sulphate solution: \( \text{Zn} + \text{CuSO}_4 \rightarrow \text{ZnSO}_4 + \text{Cu} \); these reactions are called metal displacement reactions; they also work in the solid state; for example, aluminium will reduce iron oxide to iron: \( 2\text{Al} + \text{Fe}_2\text{O}_3 \rightarrow 2\text{Fe} + \text{Al}_2\text{O}_3 \).

• These reactions are all examples of redox reactions (see Topic 6 – Redox Reactions); the reactivity of the metals can be deduced directly from the standard electrode potentials, the most common of which are shown below:

<table>
<thead>
<tr>
<th>Half-reaction</th>
<th>Conventional representation</th>
<th>( E^0 / \text{V} )</th>
</tr>
</thead>
<tbody>
<tr>
<td>( \text{Mg}^{2+}(aq) + 2e^- \rightarrow \text{Mg(s)} )</td>
<td>( \text{Mg</td>
<td>Mg}^{2+} )</td>
</tr>
<tr>
<td>( \text{Al}^{3+}(aq) + 3e^- \rightarrow \text{Al(s)} )</td>
<td>( \text{Al</td>
<td>Al}^{3+} )</td>
</tr>
<tr>
<td>( \text{Zn}^{2+}(aq) + 2e^- \rightarrow \text{Zn(s)} )</td>
<td>( \text{Zn</td>
<td>Zn}^{2+} )</td>
</tr>
<tr>
<td>( \text{Fe}^{2+}(aq) + 2e^- \rightarrow \text{Fe(s)} )</td>
<td>( \text{Fe</td>
<td>Fe}^{2+} )</td>
</tr>
<tr>
<td>( \text{Pb}^{2+}(aq) + 2e^- \rightarrow \text{Pb(s)} )</td>
<td>( \text{Pb</td>
<td>Pb}^{2+} )</td>
</tr>
<tr>
<td>( 2\text{H}^+(aq) + 2e^- \rightarrow \text{H}_2(g) )</td>
<td>( \text{Pt</td>
<td>H}_2, 2\text{H}^+ )</td>
</tr>
<tr>
<td>( \text{Cu}^{2+}(aq) + 2e^- \rightarrow \text{Cu(s)} )</td>
<td>( \text{Cu</td>
<td>Cu}^{2+} )</td>
</tr>
<tr>
<td>( \text{Ag}^+(aq) + e^- \rightarrow \text{Ag(s)} )</td>
<td>( \text{Ag</td>
<td>Ag}^+ )</td>
</tr>
</tbody>
</table>

- the more negative the standard electrode potential, the more reactive the metal; the above table shows clearly that the metals can be ranked in decreasing order of reactivity as follows: magnesium, aluminium, zinc, iron, lead, (hydrogen), copper, gold (see Unit 6 – Redox Reactions).
Practical 5.2: Compare the reactivity of different metals

1) Using a measuring cylinder, pour 5 cm$^3$ of 0.5 moldm$^{-3}$ ZnSO$_4$ into each of three different test tubes
2) Use a thermometer to measure the temperature of each solution
3) Add a small spatula measure of magnesium powder to one test tube
4) Observe what happens and note any temperature change
5) Repeat steps (c) to (d) but this time adding copper powder instead of magnesium powder
6) Repeat steps (c) to (d) but this time adding iron powder instead of magnesium powder
7) Repeat steps (a) to (f) but this time use 5 cm$^3$ of 0.5 moldm$^{-3}$ CuSO$_4$, and add zinc powder to the first test tube, magnesium powder to the second test-tube and iron powder to the third test tube
8) Repeat steps (a) to (f) but this time use 5 cm$^3$ of 0.5 moldm$^{-3}$ FeSO$_4$, and add zinc powder to the first test tube, magnesium powder to the second test-tube and copper powder to the third test tube
9) Repeat steps (a) to (f) but this time use 5 cm$^3$ of 0.5 moldm$^{-3}$ MgSO$_4$, and add zinc powder to the first test tube, copper powder to the second test-tube and iron powder to the third test tube
10) Record your results in a table similar to the one below:

<table>
<thead>
<tr>
<th>Metal powder</th>
<th>salt solution</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>ZnSO$_4$</td>
</tr>
<tr>
<td>Zn</td>
<td></td>
</tr>
<tr>
<td>Cu</td>
<td></td>
</tr>
<tr>
<td>Fe</td>
<td></td>
</tr>
<tr>
<td>Mg</td>
<td></td>
</tr>
</tbody>
</table>

In which mixtures does a reaction take place? Write an equation for each reaction.
Can you list the four metals copper, magnesium, zinc and iron in order of reactivity?
UNIT 9 – METALS AND THEIR COMPOUNDS

Chemicals needed per group: 12 test tubes, three 10 cm³ measuring cylinders, one test tube rack, one thermometer access to 0.5 moldm⁻³ solutions of CuSO₄, ZnSO₄, MgSO₄ and FeSO₄ (5 cm³ per group), each bottle with its own dropping pipette; access to powdered samples of Zn, Fe, Cu and Mg, each with its own spatula.

Signs of reaction will include: a temperature rise which can be large, bubbles, a change in colour of the solution or of the powder; the largest temperature change will be with Mg and CuSO₄; the reactions of Zn with CuSO₄ and Mg with FeSO₄ may also be vigorous.

<table>
<thead>
<tr>
<th>Metal powder</th>
<th>ZnSO₄</th>
<th>CuSO₄</th>
<th>FeSO₄</th>
</tr>
</thead>
<tbody>
<tr>
<td>Zn</td>
<td>Vigorous reaction – large temperature change, orange solid produced</td>
<td>Reaction – small temperature change</td>
<td>No visible reaction /temperature change</td>
</tr>
<tr>
<td>Cu</td>
<td>No visible reaction /temperature change</td>
<td>No visible reaction /temperature change</td>
<td>No visible reaction /temperature change</td>
</tr>
<tr>
<td>Fe</td>
<td>No visible reaction /temperature change</td>
<td>Reaction – small temperature change, orange solid produced</td>
<td>No visible reaction /temperature change</td>
</tr>
<tr>
<td>Mg</td>
<td>Reaction – small temperature change</td>
<td>Very vigorous reaction – very large temperature change, orange solid produced</td>
<td>Vigorous reaction – large temperature change</td>
</tr>
</tbody>
</table>

CuSO₄ with Mg, Zn and Fe: CuSO₄ + Mg → MgSO₄ + Cu; CuSO₄ + Zn → ZnSO₄ + Cu; CuSO₄ + Fe → FeSO₄ + Cu
FeSO₄ with Mg and Zn: FeSO₄ + Mg → MgSO₄ + Fe; FeSO₄ + Zn → ZnSO₄ + Fe
ZnSO₄ with Mg: ZnSO₄ + Mg → MgSO₄ + Zn

Mg most reactive as it displaces Cu, Fe and Zn from their salts; then Zn which displaces Fe and Cu but not Mg from their salts; then Fe which can only displace Cu from their salts, then Cu which cannot displace any of the other metals from their salts.

Test your knowledge 5.3: Understanding metal displacement reactions

Predict whether a chemical reaction will take place when the following substances are mixed; write the symbol equation for any reaction occurring:

(a) copper and magnesium sulphate
    (b) magnesium and copper sulphate
    (c) magnesium sulphate and zinc
    (d) zinc sulphate and copper
    (e) zinc and copper sulphate
    (f) aluminium oxide and carbon
    (g) iron (III) oxide and carbon
    (h) tin (IV) oxide and carbon

(a) no reaction; (b) Mg + CuSO₄ → MgSO₄ + Cu; (c) no reaction; (d) no reaction; (e) Zn + CuSO₄ → ZnSO₄ + Cu; (f) no reaction; (g) Fe₂O₃ + 3C → Fe₂O₃ + 3CO; (h) SnO₂ + 2C → Sn + 2CO
Lesson 6 - How are metals extracted from their ores?

Summary Activity 6.1: Electrolytic Processes
- Give the half-equations for the reactions occurring at each electrode when Al is extracted from Al₂O₃
- Give the half-equations for the reactions occurring at each electrode during the purification of copper

<table>
<thead>
<tr>
<th>Electrode</th>
<th>Reaction</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cathode: Al³⁺ + 3e⁻ → Al</td>
<td>2O²⁻ → O₂ + 4e⁻</td>
</tr>
<tr>
<td>Cathode: Cu²⁺ + 2e⁻ → Cu</td>
<td>Anode: Cu → Cu²⁺ + 2e⁻</td>
</tr>
</tbody>
</table>

**d) Extraction of Metals**

(i) General Principles of Extraction

- A mineral containing a sufficient quantity of a metal for its extraction to be commercially viable is called an *ore*.
- Because metals are so useful, the extraction of metals from their ores is very commercially important; there are a number of different processes by which metals can be extracted from their ores; the choice of extraction process depends on:
  - the position of the metal in the reactivity series
  - the cost of the process
  - the required purity of the metal

- Metals can be extracted by electrolysis:
  - this works for all ores in which the bonding is ionic
  - it produces a pure form of the metal
  - but electrolysis is very expensive and only used for the most reactive metals
  - *aluminium*, for example, is extracted by the electrolysis of purified bauxite (aluminium oxide)

- Metals can also be extracted by reacting them with another element which is more reactive:
  - carbon is often used for this purpose as the process is very cheap
  - *iron*, *tin* and *copper* are extracted by reducing their ores (e.g. Fe₂O₃ and SnO₂) with carbon
  - this method cannot be used if the metal is above carbon in the reactivity series (like aluminium) or if the metal is not useful when contaminated with carbon (like titanium)
  - in such cases hydrogen or a reactive metal such as calcium should be used

- Metals which are present only in low concentrations require a more complex extraction process, and this is usually only commercially viable for very precious metals (like *gold*).

(ii) Extraction of aluminium

- Al₂O₃ is purified and then electrolysed (see Unit 6 – Redox Reactions):
  - the Al₂O₃ is dissolved in molten cryolite (Na₃AlF₆) as it has a lower melting point than Al₂O₃
  - the electrode reactions are: Al³⁺ + 3e⁻ → Al (at the cathode) and 2O²⁻ → O₂ + 4e⁻ (at the anode)
  - graphite electrodes are used
  - the graphite anodes react with the oxygen produced and so they need to be regularly replaced
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• This process uses a lot of energy and so is expensive; for this reason, recycling aluminium is very important

(iii) Extraction of iron

• Iron is extracted from its main ore haematite (Fe₂O₃) by reaction with carbon in a blast furnace; limestone is also added to remove the main impurity (SiO₂)
  - The carbon is added in the form of coke and it reacts as follows: C + O₂ → CO₂; C + CO₂ → 2CO
  - the CO reduces the iron oxide to iron: Fe₂O₃ + 3CO → 2Fe + 3CO₂
  - Some Fe₂O₃ may be reduced by C directly: Fe₂O₃ + 3C → 2Fe + 3CO
  - limestone (CaCO₃) is also added; it decomposes on heating to give CaO: CaCO₃ → CaO + CO₂
  - the CaO (which is basic) reacts with the main impurity SiO₂ (which is acidic): CaO + SiO₂ → CaSiO₃
  - the CaSiO₃ is known as “slag”; it has a low density and can be separated from the reaction mixture; CaSiO₃ is used in road surfacing; it is laid underneath the tar

• The iron produced by this method contains a large quantity of carbon (up to 4%); some carbon is desirable (pure iron combined with a small quantity of iron makes the alloy steel) but 4% is too high; it is therefore necessary to blow oxygen through the molten iron in order to remove some of the C as CO and CO₂; other impurities such as sulphur can also be removed in this way; this is called the basic oxygen process

• The amount of carbon to be removed depends on the amount of carbon wanted in the steel; low-carbon steel is soft and used to make decorative items such as railings; medium-carbon steel is harder but still flexible and mainly used in construction; high-carbon steel is hard and brittle and used in tools

• iron is easy to recycle; it can be separated from other metals due to its magnetic properties

(iv) Extraction of tin

• Tin is reduced in a similar method to iron; its ore SnO₂ is heated with iron in a furnace: SnO₂ + 2C → Sn + 2CO

(v) Extraction of gold

• Gold exists naturally as an element; its extraction consists of concentrating the gold by separating it from the surrounding rock

• Gold is unreactive but it can be converted into a soluble compound in the presence of oxygen and cyanide ions:

• This compound can be converted back into gold by reaction with carbon; the gold is formed on the surface of the carbon
### Test your knowledge 6.2: Understanding the Extraction of Metals

(a) State three factors which determine how a metal is extracted from its ore.
(b) Explain why the extraction of aluminium is expensive.
(c) Explain why aluminium oxide is dissolved in molten cryolite.
(d) Write the equations for the electrode reactions taking place during the electrolysis of aluminium oxide.
(e) Explain why the anodes have to be regularly replaced.
(f) Write two equations to show how carbon monoxide is produce in a blast furnace.
(g) Write an equation for the reaction of tin (IV) oxide and carbon monoxide.
(h) Write an equation for the reaction of iron (III) oxide and carbon monoxide.
(i) Explain why limestone is added to the blast furnace.
(j) Explain how excess carbon is removed from the iron produced in the blast furnace.
(k) Explain why aluminium cannot be produced in the blast furnace.
(l) State one useful by-product of the blast furnace and explain why it is useful.
(m) Explain why iron is easier to recycle than aluminium or tin.
(n) Explain why the extraction of gold is difficult, even though it occurs naturally.
(o) Outline the stages in the extraction of gold (no equations required).

| (a) | Reactivity, required purity, cost of process |
| (b) | Lots of energy needed to melt the cryolite and for the electricity |
| (c) | The melting point of cryolite is lower than the melting point of pure aluminium oxide |
| (d) | $\text{Al}^{1+} + 3\text{e}^- \rightarrow \text{Al}$ (at the cathode) and $2\text{O}^{2+} \rightarrow \text{O}_2 + 4\text{e}^-$ (at the anode) |
| (e) | Anodes react with oxygen $\text{C} + \text{O}_2 \rightarrow \text{CO}_2$ |
| (f) | $\text{C} + \text{O}_2 \rightarrow \text{CO}_2$; $\text{C} + 2\text{CO}_2 \rightarrow 2\text{CO}$ |
| (g) | $\text{SnO}_2 + 2\text{CO} \rightarrow \text{Sn} + 2\text{CO}_2$ |
| (h) | $\text{Fe}_2\text{O}_3 + 3\text{CO} \rightarrow 2\text{Fe} + 3\text{CO}_2$ |
| (i) | It helps remove the main impurity $\text{SiO}_2$; $\text{CaCO}_3$ decomposes to produce $\text{CaO}$, which reacts with $\text{SiO}_2$ to produce $\text{CaSiO}_3$, which can be removed. |
| (j) | Oxygen is bubbled through the molten iron; the oxygen removes the C as $\text{CO}_2$: $\text{C} + \text{O}_2 \rightarrow \text{CO}_2$ |
| (k) | $\text{Al}$ is more reactive than $\text{C}$ and cannot be reduced from its oxide by $\text{C}$ or $\text{CO}$ |
| (l) | $\text{CaSiO}_3$ is used in road-building |
| (m) | $\text{Fe}$ is magnetic so can be separated from other scrap using a magnet |
| (n) | It is present in low concentrations and is difficult to obtain in pure form |
| (o) | It reacts with oxygen in the presence of cyanide ions to form a soluble compound; this compound is converted back to gold by reaction with carbon |
Lesson 7 – Why are metals and their compounds useful (part I)?

e) Uses of Metals and Metalloids

- The uses and applications of different metals depends on their physical and chemical properties:
  - **Sodium** is too reactive to be useful in elemental form; due to its reactivity, it can be used to produce titanium from titanium chloride; this is used in the extraction of titanium from its ore: \( \text{TiCl}_4 + 4\text{Na} \rightarrow 4\text{NaCl} + \text{Ti} \)
  - **Calcium** is also too reactive to be useful in elemental form; like sodium, its reactivity makes it useful in extracting other elements from their ores; calcium is used in the extraction of uranium from its ore
  - **Aluminium** is very useful in elemental form due to its strength, low density and resistance to corrosion (due to its protective oxide layer); it has a lower density than iron and doesn’t corrode as easily; it is also non-toxic so can be used in food containers; aluminium is used in aeroplane parts, cans, foils and kitchen utensils; aluminium is often combined with other elements to make alloys which improve its properties (silicon is added to make it harder and stronger; magnesium is added to reduce its density)
  - **Tin** forms a protective oxide layer and is often used to coat other metals to prevent corrosion; tin combines with copper to make the alloy bronze; this is used to make statues and ornaments, as well as coins; tin combines with lead to make the alloy solder; this has a low melting point and is useful for welding electrical components together
  - **Iron** is very strong, especially when small amounts of carbon are added to make steel; steel is used to make steel girders for construction and is used in reinforced concrete; it is also used in bridges, pylons, bicycles, cars and pipes; because of its magnetic properties it is also used to make magnets
  - **Gold** is very unreactive and also soft; it is used in jewellery as it stays shiny for a long time; it is also stored as currency
  - **Copper** is unreactive and very malleable so is used to make coins and pipes; it is a good conductor of electricity so it is used to make electrical wires; when mixed with zinc it makes the alloy brass; brass has antibacterial properties and it used in taps and door handles, especially in bathrooms; it is shiny and resists corrosion and is often used to coat ornaments; it is also highly sonorous and so used to make musical instruments (trumpets, trombones, saxophones and horns are known as “brass” instruments); when copper is mixed with tin it makes the alloy bronze; bronze is very malleable and used in coins and sculptures; brass and bronze are both unreactive, like copper

- **Metalloids** (eg silicon and boron) are semi-conductors; they conduct poorly at low temperatures but better at high temperatures; for this reason they are very useful in making electronic components; they are also added to metals to make them harder (less metallic)

---

Test your knowledge 7.1: Describing uses of metals and metalloids

(a) Give the name of the alloys formed by mixing the following elements:
   (i) copper and tin; (ii) copper and zinc; (iii) tin and lead; (iv) iron and carbon

(b) Complete the following table to show the at least one common use of the following metals and metalloids; in each case, identify the property which makes the material suitable for its use

<table>
<thead>
<tr>
<th>Material</th>
<th>Use</th>
<th>Property</th>
</tr>
</thead>
<tbody>
<tr>
<td>tin</td>
<td></td>
<td></td>
</tr>
<tr>
<td>solder</td>
<td></td>
<td></td>
</tr>
<tr>
<td>steel</td>
<td></td>
<td></td>
</tr>
<tr>
<td>gold</td>
<td></td>
<td></td>
</tr>
<tr>
<td>copper</td>
<td></td>
<td></td>
</tr>
<tr>
<td>brass</td>
<td></td>
<td></td>
</tr>
<tr>
<td>silicon</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
(a) (i) bronze; (ii) brass; (iii) solder; (iv) steel
(b) Properties and Reactions of Metal Compounds

(i) Compounds of sodium

- All compounds of sodium are soluble in water; like most metals, it forms basic oxides and hydroxides and it can also form salts
- Sodium ions (Na⁺) are colourless; they form colourless solutions and white solids, unless the anion in the compound is coloured; this is true of all s and p-block metal ions
- The most common salts of sodium are:
  - **sodium chloride (NaCl)** is a salt of a strong acid and a strong base; it is therefore neutral; it is commonly known as salt, common salt or table salt; it is added as flavouring to food and as a source of Na⁺ ions; it is also placed on roads in cold weather to stop water freezing (all dissolved solutes lower the freezing point of water) as it is very cheap and non-toxic
  - **sodium nitrate (NaNO₃)** is a salt of a strong acid and a strong base; it is therefore neutral; it is commonly known as Chile saltpetre; it is an important source of nitrate ions which are important in fertilisers
  - **sodium sulphate (Na₂SO₄)** is a salt of a strong acid and a strong base; it is therefore neutral; it is mainly used in detergents
- **sodium hydroxide (NaOH)** is a strong base due to the OH⁻ ion; it is also known as caustic soda; it is mainly used to neutralise fatty acids to make soaps; it is also used in the manufacture of paper

(ii) Compounds of calcium

- Like most metals, calcium forms basic oxides and hydroxides and it can also form salts
- Calcium ions (Ca²⁺) are colourless; they form colourless solutions and white solids, unless the anion in the compound is coloured; this is true of all s and p-block metal ions
UNIT 9 – METALS AND THEIR COMPOUNDS

- The most common salts of calcium are:
  - **Calcium chloride (CaCl₂)** is a salt of a strong acid and a strong base; it is therefore neutral; it is highly soluble in water even at low temperatures and is very effective at reducing the freezing point of water, hence preventing freezing; it is a very useful de-icer; it is also hygroscopic and so is used as a drying agent and is also placed on dusty roads so the road holds water better and gets less dusty, which preserves the road; it is also a water softener because the presence of Ca²⁺ ions stops CO₃²⁻ and SO₄²⁻ ions dissolving in water.
  - **Calcium sulphate (CaSO₄)** is a salt of a strong acid and a strong base; it is therefore neutral; it is insoluble in water but is most stable in its hydrated form: CaSO₄.2H₂O – known as gypsum; the anhydrous form (CaSO₄) and another hydrated form CaSO₄.0.5H₂O are both hygroscopic, which makes them useful as drying agents; CaSO₄.0.5H₂O is known as plaster of Paris – it forms a mouldable paste with water which hardens into CaSO₄.2H₂O when it dries; this is the most useful application of CaSO₄.

- **Calcium oxide (CaO)** is a base due to the O²⁻ ion; the O²⁻ ion is a strong base but CaO is only slightly soluble in water so is considered a weak base; CaO absorbs water readily to form Ca(OH)₂: CaO + H₂O → Ca(OH)₂; it is therefore hygroscopic and a good drying agent; CaO is used to remove impurities in the manufacture of steel; it reacts with acidic oxides left in the iron (mainly SiO₂) to produce salts: eg CaO + SiO₂ → CaSiO₃.

- **Calcium hydroxide (Ca(OH)₂)** is a base due to the OH⁻ ion; it is only slightly soluble in water and a saturated solution of Ca(OH)₂ in water is known as limewater and is used to test for carbon dioxide; it is used in water treatment – it is added to water to reduce its acidity and the precipitate of Ca(OH)₂ also acts as an effective water filter; Ca(OH)₂ is used in the production of cement and in the production of paper; it can also be heated to convert it into CaO: Ca(OH)₂ → CaO + H₂O.

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**Test your knowledge 7.2: Describing uses of the hydroxides, oxides and salts of sodium and calcium**

(a) Name the three most common sodium salts and state a use for each one
(b) Name the two most common calcium salts and state a use for each one
(c) State two uses for each of sodium hydroxide, calcium oxide and calcium hydroxide

| (a) Sodium chloride – flavouring food and de-icing roads; sodium nitrate – fertiliser; sodium sulphate - detergent |
| (b) Calcium chloride – drying agent, making roads less dusty |
| (c) NaOH: make soap, make paper; CaO – drying agent, remove SiO₂ from iron; Ca(OH)₂ – water treatment and cement production |

**Lesson 8 – Why are metals and their compounds useful (part II)?**

(iii) **Compounds of d-block metals (Cu, Zn and Fe)**

- Most compounds of copper, zinc, iron and other d-block metals are bases or salts
- **Copper chloride (CuCl₂)** is a salt of a strong acid and a relatively weak base
  - it is therefore very slightly acidic due to the hydrolysis of the Cu²⁺ ion
  - it is a green solid and dissolves in water for form a blue-green solution
  - it is used as a catalyst in a variety of industrial processes
• **Copper sulphate (CuSO$_4$)** is a salt of a strong acid and a relatively weak base
  - it is therefore very slightly acidic due to the hydrolysis of the Cu$^{2+}$ ion
  - it is a blue solid in hydrated form and dissolves in water for form a blue solution
  - it is slightly toxic and is mainly used as a fungicide
  - it is also present in a number of chemical reagents, mainly Benedict’s solution and Fehling’s solution, which are used to test for reducing sugars in organic chemistry; Cu$^{2+}$ is a mild oxidising agent as it can be reduced to copper (I) oxide (Cu$_2$O), which is an orange solid
  - it is also used in Biuret’s solution as a test for amines

• **Copper oxide (CuO)** is a base due to the O$^{2-}$ ion
  - the O$^{2-}$ ion is a strong base but CuO is insoluble in water so is considered a weak base
  - it is a black solid and it is used in pigments
  - it is also used in the disposal of some toxic compounds; it is a weak oxidising agent as it can be reduced to copper metal and can oxidise some toxic chemicals into less harmful ones

• d-block ions have the ability to form dative covalent bonds with other molecules or anions and form complex ions:
  - any molecule or anion which can form a dative covalent bond with a metal ion is called a ligand
  - a complex ion consists of a metal ion datively bonded to one or more ligands; the ligand provides both of the electrons needed for the covalent bond
  - two examples of complex ions are [Cu(NH$_3$)$_4$]$^{2+}$ and [Zn(NH$_3$)$_6$]$^{2+}$; they are formed when a solution containing Cu$^{2+}$ or Zn$^{2+}$ ions reacts with excess ammonia; these reactions take place in two stages:
    - ammonia solution is an alkali and contains hydroxide ions (NH$_3$ + H$_2$O $\rightleftharpoons$ NH$_4^+$ + OH$^-$); so when ammonia solution is added to solutions containing Cu$^{2+}$ or Zn$^{2+}$ ions, a blue or white precipitate is formed:
      - Cu$^{2+}$(aq) + 2OH$^-$ (aq) $\rightarrow$ Cu(OH)$_2$(s) (blue)
      - Zn$^{2+}$(aq) + 2OH$^-$ (aq) $\rightarrow$ Zn(OH)$_2$(s) (white)
    - if more ammonia is added, a complex ion is formed and the precipitate redissolves to form a deep blue or a colourless solution:
      - Cu(OH)$_2$(s) + 4NH$_3$(aq) $\rightarrow$ [Cu(NH$_3$)$_4$]$^{2+}$(aq) + 2OH$^-$ (aq) (deep blue)
      - Zn(OH)$_2$(s) + 6NH$_3$(aq) $\rightarrow$ [Zn(NH$_3$)$_6$]$^{2+}$(aq) + 2OH$^-$ (aq) (colourless)
  - the reactions of Cu$^{2+}$ and Zn$^{2+}$ with aqueous ammonia can be used to confirm their presence in aqueous solution (qualitative analysis)

• The formation of complex ions is the reason why many compounds of transition metals are coloured
  - copper ions (Cu$^{2+}$), iron (II) ions (Fe$^{2+}$) and iron (III) ions are all coloured
  - complex ions in which the central metal ion has partially filled d-orbitals (ie 1 – 9 electrons in a d-subshell) have the ability to absorb visible light
    - most compounds containing Cu$^{2+}$ are blue or turquoise
    - most compounds containing Fe$^{2+}$ are green
    - most compounds containing Fe$^{3+}$ are orange or brown
    - the exact colour depends on the anion and on whether the compound is in solid or aqueous form
  - zinc ions (Zn$^{2+}$) are colourless, like the ions of s and p-block metals; this is because the 3d subshell in Zn$^{2+}$ is full (electronic configuration = [Ar]3d$^{10}$); they form colourless solutions and white solids, unless the anion in the compound is coloured; for this reason zinc is not considered a transition metal (although it is a d-block metal)
UNIT 9 – METALS AND THEIR COMPOUNDS

- Water can also behave as a ligand and most d-block metal ions in solution exist in the form of hydrated complex ions: eg $[\text{Fe(H}_2\text{O)}_6]^{2+}$ and $[\text{Cu(H}_2\text{O)}_4]^{2+}$
  - these hydrated complex ions also exist in the solid state in the form of hydrated salts: eg $\text{CuSO}_4.5\text{H}_2\text{O}$ and $\text{FeSO}_4.6\text{H}_2\text{O}$
  - the presence of water as a ligand allows the electrons in partially filled d-orbitals to absorb visible light and be coloured; anhydrous compounds of transition metals contain no complex ions and so are generally not coloured
    - Eg $\text{CuSO}_4.5\text{H}_2\text{O}$ (hydrated copper sulphate) is blue; $\text{CuSO}_4$ (anhydrous copper sulphate) is white
- Anhydrous copper sulphate turns from white to blue when water is added to it:
  \[
  \text{CuSO}_4(s) + 5\text{H}_2\text{O(l)} \rightarrow \text{CuSO}_4.5\text{H}_2\text{O(s)}
  \]
  white \quad blue

This reaction can be used as a test for water: add a few drops of the liquid to anhydrous copper sulphate; if water is present, the white solid should turn blue

Practical 8.1: Form complex ions by reacting metal ions with excess ammonia

You are given seven solutions: lead (II) nitrate, calcium chloride, iron (II) sulphate, iron (III) sulphate, zinc sulphate, aluminium sulphate and copper sulphate; you are also given a solution of ammonia

1) Pour 2 cm$^3$ of one of the above solutions into a test tube
2) Add a few drops of NH$_3$, then add NH$_3$ until the test tube is almost full; note your observations
3) Repeat steps (a) and (b) using the other solutions
4) Record all of your observations in the following table:

<table>
<thead>
<tr>
<th>Cation present in solution</th>
<th>Observation on adding a few drops of ammonia</th>
<th>Observation on adding excess ammonia</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pb$^{2+}$</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Ca$^{2+}$</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Fe$^{2+}$</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Fe$^{3+}$</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Zn$^{2+}$</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Al$^{3+}$</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Cu$^{2+}$</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Write ionic equations for the precipitation reactions taking place when a few drops of ammonia are added.

Write ionic equations for any complex formation reactions taking place when excess ammonia is added.

Which cations have different reactions with excess NH$_3$ and excess NaOH?
Equipment needed: 0.1 mol dm$^{-3}$ solutions of any soluble salt of Pb$^{2+}$, Ca$^{2+}$, Fe$^{2+}$, Fe$^{3+}$, Zn$^{2+}$, Al$^{3+}$ and Cu$^{2+}$ - one bottle of each is sufficient - each bottle should come with its own dropping pipette - 2 cm$^3$ per group; 1 – 2 mol dm$^{-3}$ ammonia solution - one bottle per group - 100 cm$^3$ per group needed; 7 test tubes and one test tube rack per group

Expected observations:

<table>
<thead>
<tr>
<th>Cation present in solution</th>
<th>Observation on adding a few drops of ammonia</th>
<th>Observation on adding excess ammonia</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pb$^{2+}$</td>
<td>white precipitate</td>
<td>no change</td>
</tr>
<tr>
<td>Ca$^{2+}$</td>
<td>white precipitate</td>
<td>no change</td>
</tr>
<tr>
<td>Fe$^{2+}$</td>
<td>dark green precipitate</td>
<td>no change</td>
</tr>
<tr>
<td>Fe$^{3+}$</td>
<td>orange/brown precipitate</td>
<td>no change</td>
</tr>
<tr>
<td>Zn$^{2+}$</td>
<td>white precipitate</td>
<td>precipitate dissolves; colourless solution formed</td>
</tr>
<tr>
<td>Al$^{3+}$</td>
<td>white precipitate</td>
<td>no change</td>
</tr>
<tr>
<td>Cu$^{2+}$</td>
<td>pale blue precipitate</td>
<td>precipitate dissolves; dark blue solution formed</td>
</tr>
</tbody>
</table>

Pb$^{2+}$(aq) + 2OH$^-$ (aq) → Pb(OH)$_2$(s); Ca$^{2+}$(aq) + 2OH$^-$ (aq) → Ca(OH)$_2$(s); Fe$^{2+}$(aq) + 2OH$^-$ (aq) → Fe(OH)$_2$(s); Fe$^{3+}$(aq) + 3OH$^-$ (aq) → Fe(OH)$_3$(s); Zn$^{2+}$(aq) + 2OH$^-$ (aq) → Zn(OH)$_2$(s); Al$^{3+}$(aq) + 3OH$^-$ (aq) → Al(OH)$_3$(s); Cu$^{2+}$(aq) + 2OH$^-$ (aq) → Cu(OH)$_2$(s)

Zn(OH)$_2$(s) + 6NH$_3$(aq) → [Zn(NH$_3$)$_6$]$^{2+}$(aq) + 2OH$^-$ (aq); Cu(OH)$_2$(s) + 4NH$_3$(aq) → [Cu(NH$_3$)$_4$]$^{2+}$(aq) + 2OH$^-$ (aq); Pb(OH)$_2$ and Al(OH)$_3$ dissolve in excess NaOH but not excess NH$_3$; Cu(OH)$_2$ dissolves in excess NH$_3$ but not excess NaOH

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**Practical 8.2: React anhydrous copper sulphate with water**

1) Using a spatula, place 2 – 3 g of anhydrous copper sulphate on a watch glass
2) Add a few drops of water; note any change in colour
3) Repeat steps (a) and (b) but add ethanol and paraffin instead of water; note any change in colour

**What type of reaction is taking place when water is added to anhydrous copper sulphate?**

**Chemicals needed:** anhydrous CuSO$_4$ (5 g per group); one bottle per class each of paraffin and ethanol, each with its own dropping pipette

**Apparatus needed per group:** three watch glasses and one spatula, access to mass balance

The water will turn anhydrous copper sulphate blue; the paraffin should not; the ethanol might turn the copper sulphate slightly blue if it also contains water

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**Test your knowledge 8.3: Describing properties and reactions of compounds of d-block metals**

(a) Give the formula of the two most common copper salts and give one use of each
(b) Give the formula of the most common base of copper and state one use for it
(c) Define the terms “complex ion” and “ligand”; give one example of a complex ion
(d) Explain why complex ions of d-block metals are often coloured
(e) Explain why zinc compounds are colourless
(f) Explain why anhydrous copper sulphate is colourless
(g) Write an equation for the reaction between Cu$^{2+}$ and excess ammonia and state what you would see
(h) Write an equation for the reaction between Zn$^{2+}$ and excess ammonia and state what you would see
(i) Describe a chemical test for water and write an equation for the reaction taking place
Lesson 9 - How can we use complex formation reactions in qualitative analysis?

g) Qualitative Analysis Part 4 – Complex Formation Reactions

Summary Activity 9.1: What can you remember about qualitative analysis?
- What is qualitative analysis?
- Which cations form precipitates with aqueous sodium hydroxide? What are the colours of these precipitates?
- Which of these precipitates dissolve in excess aqueous sodium hydroxide?

- The experimental identification of a substance of species present in a substance
- Fe\(^{2+}\) (dark green), Fe\(^{3+}\) (orange); Ca\(^{2+}\) (white); Al\(^{3+}\) (white); Pb\(^{2+}\) (white); Cu\(^{2+}\) (pale blue); Zn\(^{2+}\) (white)
- Al(OH)\(_3\), Pb(OH)\(_2\) and Zn(OH)\(_2\)

- Some simple qualitative tests for cations, anions and gases were covered in Unit 5 – Acids, Bases and Salts, Unit 6 – Redox Reactions and Unit 8 – Solubility and Precipitation

- Complex formation reactions are useful in qualitative analysis; Fe\(^{2+}\), Fe\(^{3+}\), Ca\(^{2+}\), Al\(^{3+}\), Pb\(^{2+}\), Cu\(^{2+}\) and Zn\(^{2+}\) all give insoluble hydroxide precipitates when added to ammonia but only Cu(OH)\(_2\) and Zn(OH)\(_2\) dissolve in excess ammonia to give a complex ion
**Practical 9.2: Use complex formation reactions to identify cations in solution**

You are given four colourless solutions, labelled A, B, C and D; each solution contains one cation from the following: \( \text{Ca}^{2+}, \text{Zn}^{2+}, \text{Pb}^{2+} \) and \( \text{Al}^{3+} \).

1) Pour 2 cm\(^3\) of solution A into a test tube
2) Add a few drops of NaOH, then adding NaOH until the test tube is almost full; note your observations
3) Repeat steps (a) to (b) using solutions B – D
4) Pour 2 cm\(^3\) of solution A into a test tube
5) Add a few drops of \( \text{NH}_3 \), then adding \( \text{NH}_3 \) until the test tube is almost full; note your observations
6) Repeat steps (a) to (b) using solutions B – D

Use your observations to identify which cation is in which solution; present your answers in a table similar to the one below:

<table>
<thead>
<tr>
<th>Solution</th>
<th>Few drops NaOH</th>
<th>Excess NaOH</th>
<th>Few drops ( \text{NH}_3 )</th>
<th>Excess ( \text{NH}_3 )</th>
<th>Cation Present</th>
</tr>
</thead>
<tbody>
<tr>
<td>A</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>B</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>C</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>D</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Which two cations cannot be distinguished by this combination of tests? How would you distinguish between solutions containing these cations?

Equipment needed: 0.1 moldm\(^{-3}\) solutions of any soluble salt of \( \text{Pb}^{2+}, \text{Ca}^{2+}, \text{Zn}^{2+} \) and \( \text{Al}^{3+} \) - one bottle of each is sufficient; they should be labelled A, B, C and D - each bottle should come with its own dropping pipette - 5 cm\(^3\) per group; 1 – 2 moldm\(^{-3}\) ammonia solution - one bottle per group - 50 cm\(^3\) per group needed; 0.5 - 1 moldm\(^{-3}\) NaOH solution – one bottle per group - 50 cm\(^3\) per group needed; 8 test tubes and one test tube rack per group

Expected observations and results:

<table>
<thead>
<tr>
<th>Solution</th>
<th>Few drops NaOH</th>
<th>Excess NaOH</th>
<th>Few drops ( \text{NH}_3 )</th>
<th>Excess ( \text{NH}_3 )</th>
<th>Cation Present</th>
</tr>
</thead>
<tbody>
<tr>
<td>A</td>
<td>white precipitate</td>
<td>dissolves - colourless solution</td>
<td>white precipitate</td>
<td>no change</td>
<td>( \text{Pb}^{2+} ) or ( \text{Al}^{3+} )</td>
</tr>
<tr>
<td>B</td>
<td>white precipitate</td>
<td>no change</td>
<td>white precipitate</td>
<td>no change</td>
<td>( \text{Ca}^{2+} )</td>
</tr>
<tr>
<td>C</td>
<td>white precipitate</td>
<td>dissolves - colourless solution</td>
<td>white precipitate</td>
<td>dissolves - colourless solution</td>
<td>( \text{Zn}^{2+} )</td>
</tr>
<tr>
<td>D</td>
<td>white precipitate</td>
<td>dissolves - colourless solution</td>
<td>white precipitate</td>
<td>no change</td>
<td>( \text{Pb}^{2+} ) or ( \text{Al}^{3+} )</td>
</tr>
</tbody>
</table>

\( \text{Pb}^{2+} \) and \( \text{Al}^{3+} \) cannot be distinguished by these tests; \( \text{Pb}^{2+} \) gives a precipitate with \( \text{Cl}^- \) ions but \( \text{Al}^{3+} \) does not, so the addition of a few drops of hydrochloric acid will give a white precipitate with the solution containing \( \text{Pb}^{2+} \) but not the solution containing \( \text{Al}^{3+} \).
Test your knowledge 9.3: Understanding qualitative analysis by complex formation reactions
(a) Describe a chemical test to distinguish between solutions of aluminium sulphate and zinc sulphate

Answer: add dilute ammonia dropwise until in excess; both solutions will give a white precipitate; the precipitate formed from the solution of zinc sulphate will dissolve in excess ammonia but the precipitate formed from the solution of aluminium sulphate will not.

Lesson 10 – What have I learned about metals and their compounds?

1. Describe the bonding in metals and hence explain why metals are malleable and ductile.

2. State two other physical properties of metals and explain why metals have these properties.

3. Write equations for the following reactions and state the type of reaction occurring:
   (a) Magnesium with dilute nitric acid
   (b) Sodium with water
   (c) Zinc with copper sulphate

4. Explain why zinc reacts with copper sulphate but copper does not react with zinc sulphate.

5. Describe how aluminium is extracted from its ore.

6. State why brass and solder are useful alloys and state which pure metals they are made from.

7. Explain why aqueous solutions of zinc salts are colourless but aqueous solutions of copper salts are coloured.

8. Explain the difference between a d-block metal and a transition metal.

9. Explain why transition metals can form variable oxidation states.

10. State what you would see when dilute ammonia solution is added gradually to a solution containing Zn²⁺ ions. Write equations for any reactions occurring.

11. Explain how you would distinguish between aqueous solutions of zinc nitrate and aluminium nitrate.

12. Describe a chemical test for water.
UNIT 9 – METALS AND THEIR COMPOUNDS

1. Lattice of cations, held together by a sea of delocalised electrons; cations can move past each other without breaking the attraction between cations and electrons
2. Electrical conductors (delocalised electrons); sonorous (sound waves can travel through with little loss of energy); lustrous (electrons reflect light back to its source)
3. (a) Mg + 2HNO\(_3\) → Mg(NO\(_3\))\(_2\) + H\(_2\); (b) 2Na + 2H\(_2\)O → 2NaOH + H\(_2\); (c) Zn + CuSO → ZnSO\(_4\) + Cu; redox reactions
4. Zinc is more reactive than copper so zinc can displace copper from its compounds; copper is less reactive than zinc so copper cannot displace zinc from its compounds
5. Purified Al\(_2\)O\(_3\) is dissolved in molten cryolite and electrolysed using graphite anodes; molten aluminium is produced at the cathode
6. Brass – used in taps due to its anti-bacterial properties; made from copper and zinc; solder is used to weld electrical components together due to its low melting point; made from tin and lead
7. Hydrated Cu\(^{2+}\) ions have a d\(^9\) configuration so can absorb visible light; hydrated Zn\(^{2+}\) ions have a d\(^{10}\) configuration so cannot
8. d-block metals have s and d electrons but no p-electrons in their outer shell; transition metals can form at least one stable ion with partially filled d-orbitals; all transition metals come from the d-block but not all d-block metals are transition metals
9. the number of d-electrons lost by transition metals can vary depending on the reaction; the energy required to remove the d-electrons is sometimes but not always recovered in bonding
10. with Zn\(^{2+}\); white precipitate, which dissolves in excess ammonia to give a colourless solution: Zn\(^{2+}\)(aq) + 2OH\(^-\)(aq) → Zn(OH)\(_2\)(s); Zn(OH)\(_2\)(s) + 6NH\(_3\)(aq) → [Zn(NH\(_3\))\(_6\)]\(^{2+}\)(aq) + 2OH\(^-\)(aq); with Cu\(^{2+}\); pale blue precipitate, which dissolves in excess ammonia to give a deep blue solution: Cu\(^{2+}\)(aq) + 2OH\(^-\)(aq) → Cu(OH)\(_2\)(s); Cu(OH)\(_2\)(s) + 4NH\(_3\)(aq) + 2H\(_2\)O(l) → [Zn(NH\(_3\))\(_4\)(H\(_2\)O)\(_2\)]\(^{2+}\)(aq) + 2OH\(^-\)(aq)
11. add aqueous NH\(_3\) to both gradually until in excess; with Zn\(^{2+}\), a white precipitate will form which dissolves in excess NaOH; with Al\(^{3+}\), a white precipitate will form which is insoluble in excess NaOH
12. Add a few drops of the liquid to anhydrous copper sulphate; if it turns blue, water is present.