Most metals are found in the ground as compounds (mineral ores). An ore is a rock that contains enough metal to make it worthwhile extracting it. Many exist as **metal oxides**.

e.g. Iron (III) oxide

A common ore of iron is haematite (heema-tite), which contains iron oxide: Iron + Oxygen \rightarrow Iron oxide 4Fe + $3O_2 \rightarrow 2Fe_2O_3$



This is an **oxidation** reaction because the metal gains oxygen.

Extracting iron from iron oxide is a **reduction** reaction, because it involves the loss oxygen.

 $Fe_2O_3 + 3CO \longrightarrow 2Fe + 3CO_2$ $+ 3 \bigcirc \longrightarrow 2 \bigcirc + 3 \bigcirc (1 + 3 \bigcirc (1 + 3))$

Aluminium oxide

Bauxite is aluminium oxide.

Aluminium + Oxygen \rightarrow Aluminium oxide This prevents aluminium corroding. Aluminium is too reactive to be obtained by reduction by carbon. Electrolysis must be used.

Oxides of Group 1 and Group 2

- Sodium oxide is formed when sodium is burned in oxygen
- It is a white powder and is soluble
- During the reaction, Na loses electrons and O gains electrons.

 $2Na - 2e^{-} \rightarrow 2Na^{+}$ $O_2 + 2e^{-} \rightarrow O^{2-}$

- This forms a basic oxide
- When dissolved in water, it makes a hydroxide solution (alkaline) and turns universal indicator purple

```
Na_2O + H_2O \rightarrow 2NaOH
```

The Reactivity Series

Most reactive



Least reactive

The differences in reactivity is because of the different **tendency** of each metal to form positive ions.

electron in outer shell is further from the nucleus, there is less attractive force, so the electron is more easily lost



A more reactive metal can displace a less reactive metal from a solution of a compound of the less reactive metal. This is a **displacement** reaction.

Iron sulphate + Magnesium \rightarrow Magnesium sulphate + Iron



Ion formation Positive ions are formed by a loss of electrons. $(K - e^- \rightarrow K^+)$ Displacement would be shown by: $Mg + Fe^{2+} \rightarrow Mg^{2+} + Fe$

Extraction of metals

Most metals are found as compounds as they react with other elements

Metals more reactive than carbon need to be extracted by electrolysis.	For example:	potassium, sodium, calcium, magnesium, aluminium
Metals less reactive than carbon can be extracted from their oxides by reduction with carbon .	For example:	zinc iron lead

Reduction with carbon usually refers to 'smelting' and involves the **loss of oxygen** eg. zinc oxide + carbon → zinc + carbon dioxide

Reaction of metals with acids Metal + Acid → Salt + Hydrogen To make a crystallised salt:

a. excess magnesium needs to be added to the acidb. the solution needs to be filtered into a crystallising dishc. the solution needs to be concentrated by evaporationd. and the solution then left to evaporate, to crystallise.

		a		
powdered metal			filter pape and funn	er unres el solid
	spatula warm acid	unreacted	salt solution	salt

Forming lons

Metal	5	Acids			
nagnesium	Mg ²⁺	sulfuric acid	H_2SO_4	$2H^+$	SO ₄ 2-
ron	Fe ²⁺	hydrochloric acid	HCl	H^{+}	C1 ⁻
zinc	Zn^{2^+}				
The charaes must cancel each					

The charges must cancel each other out and the equation must be balanced

 $2H^+ + 2e^- \rightarrow H_2$

Electron gain is

reduction

Using half equations Mq + 2H⁺ \rightarrow Mg²⁺ + H₂

 $Mg - 2e^- \rightarrow Mg^{2+}$

Electron loss is oxidation

As both these reactions take place at the same time this is a *redox* reaction

Alkalis and Bases

- A base is any substance that neutralises an acid.
- Metal oxides and metal hydroxides are **bases**.
- A few bases are soluble in water, these are called **alkalis**.

NEUTRALISATION happens when an acid and base react

Acid + Base → Salt + Water



Metal **carbonates** react with acids to make a metal salt and water, but **carbon dioxide** is also produced.

Acid + Metal Carbonate \rightarrow Salt + Water + CO2The salt madeCommon acidThe salt madenitricnitrateThe second part ofnitricnitratethe salt name issulfuricsulfateThe first part is takenhydrochloricchloridefrom the base.							
Ions from alk: carbonates	lis and	Ions fi carboi	om bas 1ates	ses and	1	Ions fro acids	om
sodium	Na ⁺	magne	sium	Mg	2+	chloride	e Cl⁻
potassium	K^+	zinc		Zn ²	+	nitrate	NO3-
calcium	Ca ²⁺	copper		Cu ²	+	sulfate	SO4 ²⁻
eg. MgSO ₄ , K ₂ SO ₄ , KCl, CaCl ₂ > BALANCED CHARGES							
Soluble S Stage 1 Mix the reactan Ensure the copy in excess excess copper oxide	Salts ts and stir. er oxide is struct acid	Stage 2 F	ilter off the xcess copper	oxide opper sulfat olution and xcess coppe xide funnel paper ate per sulfate ion)	te ar	stage 3 Leav evap cryst dish sultate solution copper	e the filtrate to orate in an orating basin c allising dish allising per sulfate rystals
Single charg	e Cl⁻	NO3-	он⁻	Na ⁺	K+	H^{+}	
Doub charg	e SO4 ²⁻	Ca ²⁺	Mg ²⁺	Zn ²⁺	Fe ²⁺	Cu ²⁺	
CuO	+ 2HCI		 CuCl 	2	+	H ₂ O	

Acids and alkalis Acids are substances that produce hydrogen ions in aqueous solution.				
Examples of a	acids are:	Exa	amples of all	kalis are:
hydrochloric acid	HCl		sodium hydroxide	NaOH
nitric acid	HNO3		potassium hydroxide	кон
sulfuric acid	H_2SO_4		ammonia	NH ₃ (aq
ethanoic acid	CH ₃ COOH			
citric acid	$C_6H_8O_7$			

The hydrogen ions they produce have the symbol H⁺. They are ions with a positive charge.

The pH Scale

When universal indicator (UI) is added to solutions, it changes colour.

1 2 3 4 5 6 7 8 9 10 11 12 13 14

1 = very acidic, 7 = neutral, 14 = very alkaline

The higher the concentration of acid, the lower the pH

Neutralisation

If an acid is added to an alkali, neutralisation takes place

An acid solution has a low pH. If an alkali is added slowly to an acid, the pH number of the acid will gradually increase. When it gets to pH 7 the acid is neutralised.

An alkaline solution has a high pH. If acid is slowly added to an alkali, the pH number will gradually decrease. When it gets to pH 7 the

Alkalis are substances

that make hydroxide

These hydroxide ions

have the symbol OH .

They are ions with a

negative charge.

ions in aqueous solution.

alkali has been neutralised.

 $H^+_{(aa)} + OH^-_{(aa)} \rightarrow H_2O_{(l)}$

Neutralisation leaves no free H⁺ ions.

(HT) Strong and We	Electrol	ysis is the proc	ess of	
Strong: In water all of the acid molecules, HA, become ions (H ⁺ and A ⁻).	Weak: In water only a few of the acid molecules, HA, become ions (H^+ and A^-), most stay as molecules.	ionic co so brea dischar	k the compound to m	ove ic nd do e elem
Strong acids ionise completely in water.	Weak acids do not ionise fully. The equilibrium lies to the left.	electro	des.	
HCl → H ⁺ + Cl ⁻ A high concentration of H ⁺ means that the pH is low.	CH ₃ COOH ≓H ⁺ + CH ₃ COO [−] A low concentration of H ⁺ means that the pH is higher.	The cathod electrode.	le is the negative	The elec
strong acids	weak acids	positive ion cathode and	as are attracted to the d are called <i>cations</i>	negati the an <i>anion</i>
hydrochloric nitric sulfuric It is possible to have c strong and concentrated	ethanoic citric carbonic in acid that is: weak and concentrated	di periori	Half Equ the disc Na ⁺ + e ⁻ the disc C ⁺ - e ⁻ BUT chlo atoms, s	ations harge c → Na harge c → Cl rrine is c o: 2Cl
ତି ବିଶିତି ତି ତି ବିତ ତି ବି ବି ତି ତି ତି ତି ବି ତି ବି ତି ବି ତି ତି ବି ତି ତି ବି ବି ତି ତି ବି ବି ତି ତି ବି ତି ବି ତି ବି ବି ତି ତି ତି ବି ତି ବି ତି ତି strong and dilute	မာ မာ မာ မာ မာ မာ မာ မာ မာ မာ မာ မာ မာ မ	Electro Is n bro Co	olysis of molter nade up of two omide) In conduct ele	<u>n com</u> o ions ectricit
6 6 6 6 6 6 6 6 6 6 6 6 6 6 6 6 7 6 6 6 7 6 6 6	60 6 6 6 6 6 6 6 6 6 6 6 6 6 6 6 6 6 6	me	elted (ions can d.c. power	supply
 (HT) pH, Neutralisation and Titration Curves The pH scale is related to the concentration of H⁺ ions Strong acids have a lower pH 			dions (P)	•••
Low pH = higher concentration H+	High pH = lower concentration H+	negative ele	ectrode	
As the pH decreas concentration incr	At the Electron	e cathode	Eleci	
10 Neutralisation in a titro	ation curve:	Electrolyte	Half equation at cat	thode I
14 4 13 alka	, pH high	KC1	$2K^+ + 2e^- \rightarrow 2K$	2
12 - 11 -	acid has reacted with some of the	CuCl ₂	$\mathrm{Cu}^{2^+} + 2\mathrm{e}^- \to \mathrm{Cu}$	2
9-	alkali, pH falls	PbI ₂	$Pb^{2+} + 2e^- \rightarrow Pb$	2
在 7 - 6 - 5	all the alkali is neutralised, pH=7	$Al_2 O_3$	$2Al^{3+} + 6e^- \rightarrow 2Al$	6
4 - 3 - 2 - 1 -	excess acid, pH is below 7	Electro At the	olysis of aqueous cathode: 2H ⁺ + 20	solution $e^{-} \rightarrow H_{2}$

4 8 12 16 20 24 28 32 36 40 44 48 52

volume of acid in cm³

process of passing direct Electrolysis to extract metals ugh a solution or melted Metals found towards the top of the reactivity to move ions apart and series are too reactive to be extracted by carbon, so must be extracted by electrolysis. pound down and Large amounts of energy are used to melt the of the elements at the compounds and to produce an electric current, so these metals can be expensive. The anode is the positive Aluminium is extracted in this way. It is very *i*e expensive to extract which is why aluminium electrode. recycling is encouraged. negative ions are attracted to to the the anode and are called Electrolysis of aluminium ions anions Aluminium ore is **bauxite** (Al_2O_3) . • It is mixed with cryolite to lower its If Equations discharge of a sodium ion melting temperature. + + e⁻ → Na The electrolysis cell has carbon discharae of a chlorine ion: electrodes. - e⁻ → Cl chlorine is a molecule of 2 oms, so: 2Cl⁻ - 2e⁻ \rightarrow Cl₂ olten compounds carbon lining as cathode f two ions (eg. lead steel tank lined with electricity only when heat-resistant can move) bricks power supply molten aluminium collects at the bottom Half equations $2AI_2O_3 \rightarrow 4AI + 3O_2$ At the cathode: Molten lead(II) bromide At the anode: 00 6O²⁻ + 12e⁻ → 3O₂ 05 Negative bromide ions attracted to the positive $3C + 3O_2 \rightarrow 3CO_2$ electrode At the cathode At the anode H⁺ is discharged in preference to Zn²⁺ Al³ Mg²⁺ Ca²⁺ Na⁺ and K⁺ Electrons are given up Cu2+ and Ag+ are discharged in preference Cl⁻ and Br⁻ are discharged in preference to to H+ (as Cu2+ and Ag+ are less reactive than H⁺) at cathode Half equation at anode H⁺ produce: 2K $2Cl^{-} - 2e^{-} \rightarrow Cl_{2}$ The e Cu $2Cl^{-} - 2e^{-} \rightarrow Cl_{2}$ To the cath Copper ions Pb $2I^- - 2e^- \rightarrow I_2$ Hydrogen io $6O^{2-} - 12e^- \rightarrow 3O_2$ 2A1 Hydrogen io electrolyte. eous solutions Copper ions

At the anode: $4OH^- - 4e^- \rightarrow O_2 + 2H_2O$

carhon anodes molten aluminium oxide $4AI^{3+} + 12e^{-} \rightarrow 4AI (AI^{3+} + 3e^{-} \rightarrow AI - per ion)$ The oxygen reacts with the electrodes: Preferential discharge of ions At the anot OHT is discharged in preference to SO42-

0	OH	
s hydrogen gas	OH [−] prod	luces oxygen gas
electrolysis	of coppe	<u>er sulphate</u>
ođe	To the anode	
(Cu ²⁺) ns (H ⁺) (from the water)	Sulfate ions (SO ₄ ²⁻) Hydroxide ions (OH ⁻) (from the water)	6 d.c. power supply
ns stay in the are discharged <i>in</i> nd make copper metal.	Sulfate ions stay in the electrolyte as hydroxide ions are more easily discharged. Hydroxide ions discharge <i>in</i> <i>preference</i> and form oxygen gas.	negative positive electrode

preference a