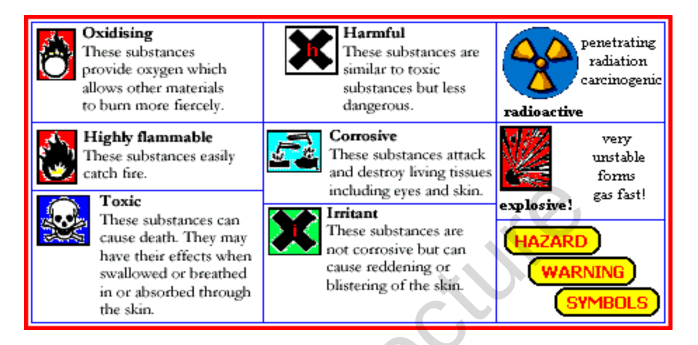
### **CHEMICAL TESTS**



INORGANIC TESTS			
TEST FOR	TEST METHOD	OBSERVATIONS	TEST CHEMISTRY
hydrogen gas H <sub>2</sub>		<b>squeaky pop!</b> (might see condensation on test tube)	$2\mathbf{H}_{2(g)} + O_{2(g)} ==> 2H_2O_{(I)} + $ energy!
CO <sub>2</sub>	bubble into limewater (aqueous calcium hydroxide solution)	milky white precipitate of calcium carbonate	$Ca(OH)_{2(aq)} + CO_{2(g)} ==> CaCO_{3(s)} + H_2O_{(l)}$
oxygen gas O <sub>2</sub>	glowing splint or spill	re-ignites it - flame	$C_{\text{(in wood)}} + \mathbf{O}_{2(g)} ==> CO_{2(g)}$
chloride gas	(ii) drop of silver		(i) Strongly acid gas, (ii) in water forms chloride ions - hence precipitate with silver nitrate.
bromide HBr and Hydrogen iodide	they are		as above - combination of acid and halide ion tests

dioxide gas SO <sub>2</sub>	potassium dichromate(VI) paper	orange to green	the dichromate(VI) ion, $Cr_2O_7^{2-}_{(aq)}$ is reduced to the green $Cr^{3+}_{(aq)}$ ion
NH <sub>2</sub>	odour*, (i) red	(ii) white clouds with HCl fumes.	(i) only common alkaline gas and (ii) forms fine ammonium chloride crystals with HCl (*volatile organic aliphatic amines give the same result, and smell more fishy)
das Cla	drop silver nitrate on the end of a glass rod	litmus turns red and	(i) non-metal, is acid in aqueous solution and a powerful oxidising agent, (ii) forms chloride ion in water
×		(i) purple vapour, (ii) blue black colour with starch solution	
	no simple relatively unambiguous test	nasty brown gas	strong oxidising agent
	anhydrous		(i) blue hydrated copper(II) crystals or solution formed, (ii) hydrated cobalt ion formed $[\text{Co}(\text{H}_2\text{O})_6]^{2^+}$
(or hydrogencarbonate HCO3 <sup>-</sup> )	strong acid to the suspected carbonate - if	limewater milky cloudy (see above CO <sub>2</sub> )	carbonate/hydrogencarbonate + acid ==> salt + water + <b>carbon dioxide</b> , then white precipitate with limewater.
ion [sulphate(VI)] SO <sub>4</sub> <sup>2-</sup>	to a solution of the suspected sulphate add dilute hydrochloric acid and a few drops of barium chloride or nitrate solution	barium sulphate	Ba <sup>2+</sup> (aq) + <b>SO<sub>4</sub><sup>2-</sup>(aq)</b> ==> <b>BaSO<sub>4(s)</sub></b> any soluble barium salt + any soluble sulphate ==> barium sulphate
[sulphate(IV)] <b>SO<sub>3</sub><sup>2-</sup></b>	any gas evolved with fresh	sulphur dioxide gas formed, (ii) the dichromate	(i) sulphite salt + hydrochloric acid ==> chloride salt + sulphur dioxide, (ii) the sulphur dioxide reduces the dichromate(VI) to chromium(III). Note: sulphites do not give ppt. with acidified barium chloride/nitrate because sulphites dissolve in acids.

	dichromate(VI)		
	paper		
Sulphid e ion  S <sup>2-</sup> for (ii)  dangerous hydrogen sulphide formed	<ul><li>(i) If soluble, add a few drops lead(II) ethanoate solution.</li><li>(ii) If solid, add dil. HCl(aq) acid, test</li></ul>	sulphide. (ii) Rotten egg smell	(i) $Pb^{2+}_{(aq)} + S^{2-}_{(aq)} => PbS_{(s)}$ (ii) $MS_{(s)} + 2H^{+}_{(aq)} => M^{2+}_{(aq)} + H_{2}S_{(q)}$ (e.g. M = Pb, Fe, Cu, Ni etc.) Then reaction (i) above occurs.
Chloride ion Cl <sup>-</sup>	conc. sulphuric acid, warm if necessary then test gas as for HCl	precipitate of silver chloride soluble in	(i) $Ag^+_{(aq)} + Cl^{(aq)} ==> AgCl_{(s)}$ , any soluble silver salt + any soluble chloride ==> silver chloride precipitate, (ii) $Cl^{(s)} + H_2SO_{4(l)} ==> HSO_4^{(s)} + HCl_{(q)}$ , then $Ag^+_{(aq)} + Cl^{(aq)} ==> AgCl_{(s)}$
Bromide ion Br <sup>-</sup>	conc. sulphuric	precipitate of silver bromide, only soluble	(i) $Ag^+_{(aq)} + Br^{(aq)} ==> AgBr_{(s)}$ any soluble silver salt + any soluble bromide ==> silver bromide precipitate, (ii) bromide ion is oxidised to bromine and the sulphuric acid is reduced to sulphur dioxide
Iodide ion I	sulphuric acid, (ii) get purple fumes	precipitate of silver iodide insoluble in concentrated	(i) $Ag^+_{(aq)} + I^{(aq)} ==> AgI_{(s)}$ , any soluble silver salt + any soluble iodide ==> silver iodide precipitate, (ii) iodide ion is oxidised to iodine and the sulphuric acid is reduced to hydrogen sulphide, (iii) insoluble lead(II) iodide formed, $Pb^{2+}_{(aq)} + 2I^{(aq)} ==> PbI_{2(s)}$
Nitrate ion [or nitrate(V)] NO <sub>3</sub>	and fine aluminium powder (Devarda's Alloy)	contain ammonia, which turns red litmus blue, see ammonia	<ul> <li>(i) the aluminium powder is a powerful reducing agent and converts the nitrate ion, NO<sub>3</sub>-, into ammonia gas, NH<sub>3</sub></li> <li>(ii) NO complex of iron(II) formed</li> </ul>

Nitrite ion [or nitrate(III)] NO <sub>2</sub> -  Ammonium ion NH <sub>4</sub> +	(ii) Add iron(ii) sulphate solution and then conc. sulphuric acid (the 'brown ring' test)  No simple test, (i) decolourises (purple iodine from acidified powder/NaOH <sub>(aq)</sub> ar no smell at first, add COLD sodium hydroxide solution to the suspected ammonium salt	e ==> colourless) acidif d potassium iodide solut nd gives 'brown ring' tes smelly	imposes to give nasty brown fumes of NO <sub>2</sub> , (ii) it fied potassium manganate(VII), (iii) it liberates tion, (iv) forms ammonia with hot Al st - see nitrate tests above.  ammonia gas is evolved: NH <sub>4</sub> <sup>+</sup> (aq) + OH̄(aq) ==> NH <sub>3(g)</sub> + H <sub>2</sub> O <sub>(l)</sub>
Hydrogen ion ie acids! H <sup>+</sup> or	universal indicator or pH meter, (ii) add a little sodium hydrogencarbonate powder	variety of colours with univ. ind. strong - red, weak - yellow/orange,	(i) pH meter gives a value of less than 7, the lower the pH number the stronger the acid, the higher the $H^+$ concentration, (ii) $HCO_3^-$ (aq) $+ H^+$ (aq) $==> H_2O_{(I)} + CO_{2(g)}$
ie an alkali	universal indicator or pH meter, (ii) add ammonium salt	variety of colours univ. ind. dark green -	(i) pH meter gives a value of more than 7, the higher the pH number the stronger the alkali, the higher the OH $^-$ concentration, (ii) ammonia gas is evolved: $NH_4^+_{(aq)} + OH^{(aq)} ==> NH_{3(g)} + H_2O_{(I)}$
flame tests	The metal salt or other compound is mixed with concentrated hydrochloric acid and a sample of the mixture is heated strongly in a bunsen flame on	lithium Li <sup>+</sup> crimson sodium Na <sup>+</sup> yellow potassium K <sup>+</sup> lilac calcium Ca <sup>2+</sup> brick red	All colours are due to electronic excitation to a higher level. You see the light emitted as the electron returns to its lower more stable level. This is the basis of atomic emission and absorption spectroscopy. Aluminium, magnesium, iron and zinc do not produce a useful identifying flame colour.
cations via sodium	hydroxide	<ul> <li>The ppt. is not solub</li> </ul>	$_{(a)}$ + 30H $_{(aq)}$ ==> <b>Al(OH)</b> $_{3(s)}$ white precipitate ble in excess of the weak alkali ammonia, but alkali sodium hydroxide: Al(OH) $_{3(s)}$ + 30H $_{(aq)}$ hphoteric behaviour)

	•	
		calcium ion: $Ca^{2+}_{(aq)} + 2OH_{(aq)}^{-} ==> Ca(OH)_{2(s)}$ white ppt. * The
solutions (both alkalis,		ppt. is not soluble in excess of NH <sub>3</sub> or NaOH.
giving hydroxide ions,		magnesium ion: $Mg^{2+}_{(aq)} + 2OH_{(aq)} ==> Mg(OH)_{2(s)}$ white ppt. *
On . In their solutions)	i	The ppt. is not soluble in excess of NH <sub>3</sub> or NaOH. You could distinguish
	offect of excess	Mg from Ca with a <u>flame test</u> .
	alkali ara	copper(II) ion: $Cu^{2+}_{(aq)} + 2OH_{(aq)} ==> Cu(OH)_{2(s)}$
	important	***blue/turquoise ppt this does dissolve in excess ammonia to
	observations.	give a deep blue solution.
		iron(II) ion: $Fe^{2+}_{(aq)} + 2OH_{(aq)} = > Fe(OH)_{2(s)}$ dark green ppt.*
	Ali precipitates	The ppt. is not soluble in excess of NH <sub>3</sub> or NaOH.
		iron(III) ion: $Fe^{3+}_{(aq)} + 3OH_{(aq)}^{-} => Fe(OH)_{3(s)}$ brown ppt.* The
		ppt. is not soluble in excess of NH <sub>3</sub> or NaOH.
		zinc ion: $Zn^{2+}_{(aq)} + 2OH_{(aq)}^{-} ==> Zn(OH)_{2(s)}$ white ppt. The ppt.
		dissolves in both excess sodium hydroxide or ammonia to give a clear
	nature.	colourless solution.
	The test sam be	
	The test can be repeated with	
	aqueous	
	ammonia	
	solution	
	(sometimes	
	wrongly called	
	'ammonium	V)
	hydroxide'). The	
	observations are	
	usually, but not	
	always, similar.	
	ppt. = precipitate.	70 - 7 2
MISCELLANEOUS		(i) $Pb^{2+}_{(aq)} + 2I^{-}_{(aq)} = > PbI_{2(s)} lead(II)$ iodide ppt.
CATION TESTS:	iodide solution =>	
	yellow precipitate	
(i) Lead(II) ion		
Metal Carbonates	Sometimes heating	copper(II) carbonate==> copper(II) oxide + carbon dioxide: CuCO <sub>3(s)</sub>
	a metal carbonate	$==> CuO_{(s)} + CO_{2(q)}$
	strongly to	
	decompose it	[green] ==> [black] + [colourless gas, test with limewater, white
	provides some	precipitate]
	clues to its	
	identity. Adding acid => CO <sub>2</sub> and	zinc carbonate==> zinc oxide + carbon dioxide
	the colour of the	
	resulting solution	$ZnCO_{3(s)} ==> ZnO_{(s)} + CO_{2(g)}$
	(eg blue $Cu^{2+}_{(ag)}$ ,	
	may also provide	[white] ==> [yellow hot, white cold] +[colourless gas, test with
	clues. The metal	limewater, white precipitate]
	ion solution might	
	also give a flame	
	colour or a	

	hydroxide precipitate with sodium hydroxide eg copper.		
ORGANIC TESTS			
TEST FOR	TEST METHOD	OBSERVATIONS	TEST CHEMISTRY
alkyne any other non-aromatic unsaturated	through, or add liquid to, a solution of <b>bromine</b> in hexane or water	bromine, decolourises,	$RC \equiv CR + 2Br_2 ==> Br_2RC-CRBr_2$ $R = H$ , alkyl or aryl