UNIT 3: INORGANIC CHEMISTRY

Recommended Prior Knowledge: Unit 1 (Theoretical Chemistry) and Unit 2 (Physical Chemistry) should have been studied before this unit. The unit builds on the ideas of atomic and molecular structure and bonding developed in Unit 1, and those concerning enthalpy changes, equilibrium, kinetics and

redox developed in Unit 2.

Context: This self-contained unit can be studied either before or after or in between the two Organic Chemistry Units 4 and 5.

Outline: The unit covers the whole of the Inorganic part of the AS course. Concepts developed in Units 1 and 2 (see prior knowledge above) are applied to Period 3

and Groups II and VII of the Periodic Table, as well as to aspects of nitrogen and sulfur chemistry.

AO	Learning Outcomes	Suggested Teaching Activities	Learning Resources
9.1 (a)	[Across Period 3] Describe qualitatively	Graphs could be plotted of atomic radius, ionic radius and melting point from Na to Ar,	C+H 15.1-15.3
	(and indicate the periodicity in) the	(and also from Li to Ne), to show periodicity. Conductivity increases from Na to Al, then	M 87
	variations in atomic radius, ionic radius,	falls to a low at Si, then zero for the rest.	site 4 (period 3)
	melting point and electrical conductivity of		site 7 (periodic
	the elements.		trends)
			site 17 (the periodic
			table)
9.1 (b)	Explain qualitatively the variation in	Increasing nuclear charge across periodic pulls the electron shells in more. In the same	C+H 15.3
	atomic radius and ionic radius	period anions contain one more shell than cations.	M 88
			R+N 9.4
9.1 (c)	Interpret the variation in melting point and	Melting point should be related to structure (see sections 3k and 4e). Conductivity should	R+N 9.5
	in electrical conductivity in terms of the	be related to (increasingly more) delocalised e in Na-Al; conduction bands in Si; localised	
	presence of simple molecular, giant	electrons in covalent bonds for the rest.	
	molecular or metallic bonding in the	A graphic illustration of the semiconduction of Si is showing the increase in conduction on	
	elements.	heating a block of Si over a Bunsen (use a battery + bulb or ammeter circuit in a	
		demonstration of this).	
0.4.4.10		See also section 2i. Increasing nuclear charge but little extra shielding across period	5 11 6 6
9.1 (d)	Explain the variation in first ionisation	causes general increase in I.E. Slight decreases at Mg-Al, and P-S due to new sub-shell,	R+N 9.6
	energy	and repulsion between electrons sharing the same orbital, respectively.	
9.1 (e)	describe the reactions, if any, of the	Na, Mg, Al, P and S should all be burned in gas jars filled with oxygen or air. To avoid	C+H 15.4-15.5
9.1 (6)	elements with oxygen (to give Na ₂ O,	contamination by metallic combustion spoons, a good technique with Na is to heat a small	M 89
	MgO, Al_2O_3 , P_4O_{10} , SO_2 , SO_3), chlorine	piece of Na on a dry brick with a Bunsen until it starts to burn, and then to place over it an	R+N 9.7
	(to give NaCl; MgCl ₂ ; Al ₂ Cl ₆ ; SiCl ₄ ; PCl ₅),	inverted gas jar of O ₂ . This also works with Cl ₂ , producing a white powder of NaCl.	P(AS) 17, 18, 19
	and water (Na and Mg only)	Revise work done in Unit 2 on oxidation numbers. Maximum O.N. equals the number of	F (AS) 17, 18, 19
9.1 (f)	state and explain the variation in oxidation	electrons in the outer shell.	R+N 9.8
3.1 (1)	number of the oxides and chlorides	dictions in the outer shell.	10110 3.8
9.1 (g)	describe the reactions of the above	Add universal indicator solution to the oxides prepared above. The trend is from strong	C+H 15.4
3 (3)	oxides with water [treatment of peroxides	alkali (NaOH), through very weak alkali (Mg(OH) ₂) to neutral (Al ₂ O ₃ and SiO ₂) to acidic.	M 89
	and superoxides is not required]	Students should work out the full balanced equations for all reactions.	R+N 9.9
		This could include some titrations, but also practical work on the dissolving on MgO in	site 3 (the nature of
9.1 (h)	describe and explain the acid/base	acids, and the precipitation and redissolving of Al(OH) ₃ in NaOH(aq). Sodium aluminate	oxides)
' '	behaviour of Period 3 oxides and	can be represented as either NaAIO or NaAI(OH)	site 16 (periodicity)

	hydroxides, including, where relevant, amphoteric behaviour in reaction with sodium hydroxide (only) and acids		
9.1 (i)	describe and explain the reactions of the above chlorides with water	Show by experiment that NaCl and MgCl $_2$ merely dissolve (to form pH 7 solutions containing dissociated ions) whereas the rest undergo hydrolysis. With a small amount of water AlCl $_3$ gives Al $_2$ O $_3$ + HCl(g), but with an excess a solution containing [Al(H $_2$ O) $_5$ (OH)] $^{2+}$ + H $^+$ + Cl $^-$ ions is formed.	C+H 15.5 M 89 R+N 9.11 <u>site 3</u> (aluminium chloride + water)
9.1 (j)	interpret the variations and trends described above in terms of bonding and electronegativity	Electropositive metals such as Na and Mg form ionically bonded oxides which are basic, and ionically bonded chlorides which are neutral. As the electronegativity of the element increases and becomes similar to that of O or CI, covalently bonded oxides and chlorides are formed, which are acidic. Al is on the borderline – the structure of the solid is ionic but with a large degree of polarisation, and the oxide is amphoteric.	C+H 15.1 M 88-89 R+N 9.12
9.1 (k)	suggest the types of chemical bonding present in chlorides and oxides from observations of their chemical and physical properties	These generalisation can be applied to the oxides and chlorides of other elements such as Ga, Ge, As.	R+N 9.13
9.1 (I) 9.1 (m)	predict the characteristic properties of an element in a given group by using knowledge of chemical periodicity deduce the nature, possible position in	Properties such as metallic/non-metallic; macro or simple covalent; reactive or non-reactive, forming ionic or covalent oxides and chlorides, m.pt. and b.pt. Similar generalisations to those above. Also include Group trends – metals becoming more reactive down theirs groups, but non-metals less reactive.	C+H 15.1 M 89 R+N 9.14, 9.15
9.1 (III)	the Periodic Table, and identity of unknown elements from given information of physical and chemical properties	A classroom team-game of "predict the properties" of a series of elements drawn out of a hat encourages the application of knowledge.	
9.2 (a)	[For Group II] describe the reactions of the elements with oxygen and water	The reactions become more vigorous as the Group is descended. Show Mg ribbon reacting with steam and Mg powder reacting slowly with water (the hydrogen can be collected using an inverted funnel + inverted test tube)	C+H 16.1-16.2 M 93 R+N 10a.5 <u>site 7</u> (periodic trends)
9.2 (b)	describe the behaviour of the Group II oxides with water	As the solubility of the hydroxides increases from Mg to Ba, so the solutions formed become increasingly alkaline. The exothermicity of the CaO + H_2O reaction (the "slaking" of lime) should be demonstrated and contrasted with the virtual non-reaction of MgO + H_2O .	C+H 16.2 M 93 R+N 10a.7 P(AS) 20
9.2 (c)	describe the thermal decomposition of the nitrates and carbonates of Group II	Decomposition becomes more difficult down the Group, as the polarising power of the cation decreases with increasing radius. Heating samples of the (anhydrous) nitrates will show the relative ease of production of NO_2	C+H 16.4 M 93 R+N 10a.8 P(AS) 20 site 16 (periodicity)
9.2 (d)	interpret, and make predictions from, the trends in physical and chemical properties of the Group II elements and their compounds	m. pts. and b. pts. of elements show no trend, but atomic and ionic radii increase, and E ^e values become more negative. Lattice energies of the oxides, and hence their m.pts., decrease down the Group.	C+H 16.3-16.5 M 93 site 8 (Group II)
9.2 (e)	explain the use of magnesium oxide as a refractory lining material and calcium	Due to its high m. pt. and low thermal conductivity. Used in, e.g. blast furnaces (where its basic pature is also an advantage). Students could search their local areas for examples of	

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	carbonate as a building material	limestone or marble being used in buildings or for statues.	
9.2 (f)	describe the use of lime in agriculture	Describe the production of CaO and Ca(OH) ₂ from CaCO ₃ . Both CaCO ₃ and Ca(OH) ₂ are used, to decrease the acidity of the soil, and hence to enhance the uptake of essential cations by plants. Investigate local sources of "garden lime" and find out whether it is CaCO ₃ or Ca(OH) ₂ , and what it is used for.	
9.4 (a) 9.4 (b)	describe the trends in volatility and colour of chlorine, bromine and iodine interpret the volatility of the elements in terms of van der Waals' forces	Deepening colour (from pale green to orange-brown to purple) is due to the decreasing energy of the n - π^* transition. Prepare solutions of the halogens in hexane to demonstrate these colours. Increased van der Waals' forces are due to the increasing number of (polarisable) electrons in the clouds around the molecules.	C+H 17.1 M 101 R+N 12.1
9.4 (c)	describe the relative reactivity of the Group VII elements as oxidising agents	Practical examples could include X_2 + NaBr, NaI etc.; X_2 + Na $_2$ S $_2$ O $_3$. Other reactions include Cl $_2$ + Fe (\rightarrow FeCl $_3$) and I $_2$ + Fe (\rightarrow FeI $_2$)	C+H 17.4 M 101 R+N 12.2 P(AS) 21, 23
9.4 (d) 9.4 (e)	describe and explain the reactions of the Group VII elements with hydrogen (i) describe and explain the relative thermal stabilities of the hydrides, (ii) interpret these relative stabilities in terms of bond energies	The explosive nature of the $H_2 + CI_2$ mixture with light; the steady burning of a H_2 jet in a gas jar of CI_2 ; the Pt-catalysed combination of $H_2 + Br_2$ or I_2 . The steady decrease in thermal stability of HX (e.g. demonstrate the effect of a hot nichrome wire plunged into a test tube of the gas), explained in terms of the weakening H-X bond. Use of bond energies to calculate ΔH^{θ} for the reaction.	C+H 17.2 M 101 R+N 12.3, 12.4 site 21a
9.4 (f)	describe and explain the reactions of halide ions with (i) aqueous silver ions followed by aqueous ammonia, (ii) concentrated sulfuric acid	Practical work can involve preparing and noting the colours of AgX (white, cream and yellow) and whether they dissolve or not in dil or conc NH $_3$. Equations (full and ionic) and complex formation (\rightarrow [Ag(NH $_3$) $_2$] $^+$. Simple acid-base reaction with Cl $^-$; increasing oxidation of X $^-$ to X $_2$ with Br $^-$ (\rightarrow SO $_2$ as byproduct) and with I $_2$ (\rightarrow H $_2$ S and SO $_2$ byproducts) Students should use oxidation numbers to construct balanced equations for these reactions.	C+H 17.3-17.4 M101 R+N 12.5, 12.6 P(AS) 22 <u>site 21a</u>
9.4 (g)	outline a method for the manufacture of chlorine from brine by a diaphragm cell	See Unit 2. Revise the conditions, the overall equation, and the main uses of the three products.	C+H 20.4 M 84 R+N 6a.2 site 3 (the membrane cell)
9.4 (h)	describe and interpret in terms of changes of oxidation number the reaction of chlorine with cold, and with hot, aqueous sodium hydroxide	Introduce the idea of disproportionation. The preparation of KCIO ₃ from the reaction with hot KOH can be attempted, but NaClO ₃ is too soluble for easy separation. Students should use oxidation numbers to construct balanced equations for these reactions.	C+H 17.2, 17.4 M 101 R+N 12.7
9.4 (i)	explain the use of chlorine in water purification	The equilibrium $Cl_2 + H_2O = HCl + HOCl$ produces too great an acidity, so an alkali is added at the same time. Visit the local waterworks.	R+N 12.8
9.4 (j)	recognise the industrial importance and environmental significance of the halogens and their compounds, {e.g. for bleaches; pvc; halogenated hydrocarbons as solvents, refrigerants and in aerosols)	Mention the discovery by atmospheric chemists of the CFC-caused problem in the O_3 layer, and the replacement of CFCs by hydrocarbons and freons. The potential problems with the incineration of pvc waste; the phasing-out of chlorinated solvents; DDT and BHC residues in the environment. (see also Unit 4). Internet searches for other uses	C+H 17.1 R+N 12.9



9.6 (a)	explain the lack of reactivity of nitrogen	Mention that the strength of the N≡N affects both the kinetic inertness (large E _a) and the	M 97
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9.6 (c)	describe the Haber process for the	See Unit 2 (AO 6b). Revise the conditions, and the reasons for using them, the overall	C+H 20.2
(-)	manufacture of ammonia from its	equation, and the main uses of the ammonia produced.	M 83
	elements, giving essential operating		R+N 7a.7
	conditions, and interpret these conditions		site 16 (manufacture
	(qualitatively) in terms of the principles of		of ammonia)
	kinetics and equilibria		
9.6 (b)	describe: (i) the formation, and structure,	NH ₃ as a Bronsted base. Neutralisation, titration. NH ₄ ⁺ isoelectronic with CH ₄ , hence	C+H 3.12, 6.3
	of the ammonium ion	same shape.	M 97
	(ii) the displacement of ammonia	Warming with NaOH(aq) and testing for NH ₃ with moist red litmus paper – a good test for	R+N 14.2
/ 1)	from its salts	NH_4^+ .	2 11 22 2
9.6 (d)	recognize the industrial importance of	Ammonium salts, urea and NH ₃ itself as fertilisers (students could calculate the %N in	C+H 20.2
	ammonia and nitrogen compounds	each). Nitric acid, and its use in making nitrates, explosives, dyes, polyamides,	M 97
	derived from ammonia	polyurethanes, and pharmaceuticals. Internet search for the formulae and uses of these	R+N 14.3
0.0 (a)	recoming the environmental	products.	D.N.44.4
9.6 (e)	recognize the environmental	Eutrophication of rivers and lakes, especially in areas of high rainfall, due to algal blooms and subsequent decomposition using up dissolved oxygen. Discuss local examples.	R+N 14.4
	consequences of the uncontrolled use of nitrate fertilisers	and subsequent decomposition using up dissolved oxygen. Discuss local examples.	
9.6 (f)	describe and explain the occurrence, and	The production of NO from $N_2 + O_2$ in internal combustion engines. The role of Pt as a	C+H 15.4
3.0 (I)	catalytic removal, of oxides of nitrogen (in	heterogeneous catalyst. Key reactions that occur in the catalytic converter are NO + CO	R+N 14.5
	IC engines and power stations)	$\rightarrow \frac{1}{2}N_2 + CO_2$; CO + $\frac{1}{2}O_2 \rightarrow CO_2$;	114.5
	lo origines and power stations,	$h/c + O_2 \rightarrow CO_2 + H_2O$.	
9.6 (g)	explain why atmospheric oxides of	Formation of peroxyacetylnitrate (PAN) in smog. The homogeneous catalysis that occurs	
-1- (3)	nitrogen are pollutants, including their	when SO ₂ and NO ₂ interact.	
	catalytic role in the oxidation of		
	atmospheric sulfur dioxide		
9.6 (h)	describe the formation of atmospheric	The increasing use of flue gas desulfurisation (FGD) in power stations, and low-sulfur road	R+N 14.6
	sulfur dioxide from the combustion of	fuels in cars and lorries.	
	sulfur contaminated carbonaceous fuels		
9.6 (i)	state the role of sulfur dioxide in the	$SO_2 + \frac{1}{2}O_2 + H_2O \rightarrow H_2SO_4$. Catalysed by sunlight or NO_2 .	C+H 15.4
	formation of acid-rain and describe the	Damage to stonework and mortar on buildings; aggravation of asthma; acidifying lakes	R+N 14.7
	main environmental consequences of	and rivers. Investigate local examples.	
	acid-rain .		
9.6 (j)	state the main details of the Contact	See Unit 2. Revise the conditions, the equations for each stage, and the main uses of the	C+H 20.2
	process for sulfuric acid production	sulfuric acid produced. (detergents, dyes, paints, car batteries, "pickling" metals etc)	M 83
9.6 (k)	recognize the industrial importance of		R+N 7a.7, 14.8
0.0.0	sulfuric acid	White wine, dried vegetables and fruit. Extension project could be a study of food labels or	D. N. C. C.
9.6 (I)	describe the use of sulfur dioxide in food	an internet search.	R+N 14.9
	preservation		

