CHAPTER 10: Chemical Periodicity

- 10.1 Periodicity in Physical Properties
- 10.2 Periodicity in Chemical Properties
- 10.3 Period 3 Oxides
- 10.4 Period 3 Chlorides

Learning outcomes:

- (a) describe qualitatively (and indicate the periodicity in) the variations in atomic radius, ionic radius, melting point and electrical conductivity of the elements (see the Data Booklet).
- (b) explain qualitatively the variation in atomic radius and ionic radius.
- (c) interpret the variation in melting point and in electrical conductivity in terms of the presence of simple molecular, giant molecular or metallic bonding in the elements.
- (d) explain the variation in first ionisation energy.
- (e) describe the reactions, if any, of the elements with oxygen (to give Na₂O, MgO, Al₂O₃, P₄O₁₀, SO₂, SO₃), chlorine (to give NaCl, MgCl₂, Al₂Cl₆, SiCl₄, PCl₅) and water (Na and Mg only).
- (f) state and explain the variation in oxidation number of the oxides and chlorides in terms of their valence shell electrons.
- (g) describe the reactions of the oxides with water.

 [treatment of peroxides and superoxides is not required]
- (h) describe and explain the acid/base behaviour of oxides and hydroxides including, where relevant, amphoteric behaviour in reaction with sodium hydroxide (only) and acids.
- (i) describe and explain the reactions of the chlorides with water.
- (j) interpret the variations and trends in (f), (g), (h), and (i) in terms of bonding and electronegativity.
- (k) suggest the types of chemical bonding present in chlorides and oxides from observations of their chemical and physical properties.
- (l) predict the characteristic properties of an element in a given group by using knowledge of chemical periodicity
- (m) deduce the nature, possible position in the Periodic Table, and identity of unknown elements from given information about physical and chemical properties.

10.1 Periodicity in Physical Properties

What is periodicity?

1) Periodicity is the recurrence of similar properties at regular intervals when the elements are arranged in increasing atomic number.

s- bi	lock												p	- block	ŀ		Noble
	$\overline{}$			1	2	l						_				$\overline{}$	Gas
· I	П.			Н	He							Ш	IV	V	VI	VII '	0
3	4	1				'						5	6	7	8	9	10
Li	Be	J										В	C	N	0	F	Ne
11	12	d- block							13	14	15	16	17	18			
Na	Mg		a- viock								Al	Si	P	S	Cl	Ar	
19	20	21	22	23	24	25	26	27	28	29	30	31	32	33	34	35	36
K	Ca	Sc	Ti	V	Cr	Mn	Fe	Co	Ni	Cu	Zn	Ga	Ge	As	Se	Br	Kr
37	38	39	40	41	42	43	44	45	46	47	48	49	50	51	52	53	54
Rb	Sr	Y	Zr	Nb	Mo	Te	Ru	Rh	Pd	Ag	Cd	In	Sn	Sb	Te	I	Xe
55	56	57	72	73	74	75	76	77	78	79	80	81	82	83	84	85	86
Cs	Ba	La	Hf	Ta	W	Re	Os	Ir	Pt	Au	Hg	T/	Pb	Bi	Po	At	Rn
87	88	89	89														
Fr	Ra	Ac				f- bl	lock										
						<i>J- Di</i>	OLK									_	
		58	59	60	61	62	63	64	65	66	67	68	69	70	71		
		Ce	Pr	Nd	Pm	Sm	Eu	Gd	Tb	Dy	Но	Er	Tm	Yb	Lu		
		90	91	92	93	94	95	96	97	98	99	100	101	102	103		
		Th	Pa	U	Np	Pu	Am	Cm	Bk	Cf	Es	Fm	Md	No	Lr		

Variation in size of atoms

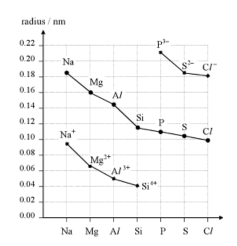
- 1) Covalent radius is half the internuclear distance between two like atoms bonded by a single covalent bond. $\frac{d_1}{2}$ = covalent radius
- between two adjacent non-bonded atoms.
- 2) Van der Waal's radius is half the average distance $\frac{d_2}{2}$ = van der Waals radius
- 3) For example, the covalent radius of Cl2 is 0.099 nm while the van der Waal's radius of Cl2 is 0.180 nm. Note: van der Waal's radius is always larger than covalent radius.
- 4) Metallic radius is half the distance between two like metal atoms bonded by metallic bond.
- 5) All these measurable quantities can be given a general name called 'atomic radii'.

- 6) Across Period 3, the atomic radius <u>decreases</u> gradually. This is because the **nuclear charge increases** while the **shielding effect remains constant**. The outer electrons are more attracted towards the nucleus, making the atoms smaller.
- 7) For comparison, metallic radii are used for Na, Mg and Al, covalent radii are used for Si, P, S and Cl. For argon, van der Waal's radius is used(argon do not form any bonds)

Note: This trend excludes argon because comparing van der Waal's radius with covalent and metallic radius is not fair.

Variation in ionic radius

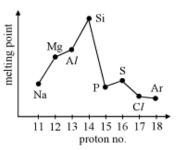
- 1) Cations are formed when an atom loses electron(s). In Period 3, Na, Mg, Al and Si form cations by losing electron(s) to achieve stable octet electronic configuration. The ions formed are Na⁺, Mg²⁺, Al³⁺ and Si⁴⁺ respectively.
- 2) Cations are <u>smaller</u> than their respective atoms because a whole layer of electrons are lost. The remaining electrons are attracted more strongly towards the centre by the same nuclear charge.



- 3) Anions are formed when an atom gains electron(s). In Period 3, P, S and Cl form anions by gaining electron(s) to achieve stable octet electronic configuration. The ions formed are P³⁻, S²⁻ and Cl⁻ respectively.
- 4) Anions are <u>bigger</u> than their respective atoms because they have more electrons than protons. The electrons are held less strongly by the nucleus. Besides, a repulsion is created between the electrons when a new electron is introduced and this causes the ion to expand.
- 5) **Anions are bigger than cations** because anions have one more shell of electrons compared to cations.
- 6) In the isoelectronic series(from Na⁺ to Si⁴⁺ and P³⁻ to Cl⁻), the ionic radius <u>decreases</u> gradually. This is because the same number of electrons are attracted more strongly by the increasing nuclear charge.

Variation in melting and boiling points

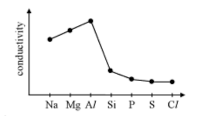
- 1) Across a Period,
 - i. melting point **increases from Na to Al** because the strength of the metallic bond increases.
 - ii. melting point of Si is **highest** because Si has a **giant covalent structure**, a lot of energy is required to overcome the strong covalent bonds.



- iii. melting points of P, S, Cl and Ar are **lower** because these have **simple molecular structures**, only weak van der Waal's forces of attraction exist between them.
- 2) Melting point of S > P > Cl > Ar because these elements exist as S₈, P₄, Cl₂ and Ar respectively. S₈ contains the **most number of electrons**, followed by P₄, Cl₂ and Ar. Van der Waal's forces get stronger with increasing number of electrons.

Variation in electrical conductivity

1) Across the Period, the elements change from metals(Na to Al) to semi-metal(Si) and then to non-metals(P to Ar).



- 2) Electrical conductivity is highest is metals, lower in semi-metals and lowest in non-metals (Most non-metals do not conduct electricity at all).
- 3) The electrical conductivity of Period 3 elements:
 - i. **increases from Na to Al** because the number of electrons contributed by per atom to the sea of delocalised electrons increases from one in Na, two in Mg and three in Al. There are more electrons to conduct electricity.
 - ii. **decreases from Al onwards**. Si is a semi-metal therefore it is a semi-conductor. The remaining elements do not conduct electricity because there are no mobile electrons.

Variation is first ionisation energy

(Refer Chapter 3)

10.2 Periodicity in Chemical Properties

Reaction with oxygen gas, O2

1) **Sodium** burns on heating with an **orange-yellow flame** to form white sodium oxide.

$$4Na(s) + O_2(g) \rightarrow 2Na_2O(s)$$

2) **Magnesium** burns on heating with a **brilliant** white flame to form white magnesium oxide.

$$2Mg(s) + O_2(g) \rightarrow 2MgO(s)$$





3) An oxide layer will form on the **aluminium** when it is exposed to air, this oxide layer prevents aluminium from reacting. However, if powdered aluminium is used, it burns on heating with **white flames** to form white aluminium oxide.

$$4Al(s) + 3O_2(g) \rightarrow 2Al_2O_3(s)$$

4) **Silicon** burns slowly at red heat to form silicon(VI) oxide or silicon dioxide.

$$Si(s) + O_2(g) \longrightarrow SiO_2(s)$$

5) **Phosphorus** burns on heating with a **white flame** to form clouds of white covalent oxides, phosphorus(III) oxide and phosphorus(V) chloride.

$$P_4(s) + 3O_2(g) \rightarrow P_4O_6(s)$$
; if oxygen gas is limited $P_4(s) + 5O_2(g) \rightarrow P_4O_{10}(s)$; if oxygen gas is in excess

6) **Sulfur** burns on heating with a **blue flame** to form sulfur dioxide gas.

$$S(s) + O_2(g) \rightarrow SO_2(g)$$

Under suitable conditions, sulfur dioxide can be converted to sulfur trioxide.(See also the Contact process)

$$2SO_2(g) + O_2(g) \rightleftharpoons 2SO_3(g)$$

- 7) **Chlorine** forms several oxides(Cl₂O and Cl₂O₇), but it will not react directly with oxygen.
- 8) **Argon** does not react with oxygen to form any oxides.
- 9) Going across Period 3, the **reactivity towards oxygen decreases** because the reducing power(tendency to be oxidised) of the elements decreases.

Reaction with chlorine gas, Cl2

1) **Sodium** burns on heating in chlorine gas with an **orange-yellow flame** to form white sodium chloride.

$$2Na(s) + Cl_2(g) \rightarrow 2NaCl(s)$$

2) **Magnesium** burns on heating in chlorine gas with a **brilliant white flame** to form white magnesium chloride.

$$Mg(s) + Cl_2(g) \rightarrow MgCl_2(s)$$

3) **Aluminium** burns on heating to form ionic aluminium chloride.

$$2Al(s) + 3Cl_2(g) \rightarrow 2AlCl_3(s)$$

At temperature about 180 °C, aluminium chloride converts to a molecular form, Al₂Cl₆, a dimer of covalent AlCl₃. At even higher temperature, Al₂Cl₆ breaks into simple AlCl₃ molecules.

$$2Al(s) + 3Cl_2(g) \rightarrow Al_2Cl_6(g)$$

4) **Silicon** burns slowly in chlorine gas at red heat to form covalent silicon(IV) chloride or silicon tetrachloride, a colourless liquid which vaporises.

$$Si(s) + Cl_2(g) \rightarrow SiCl_4(l)$$

5) **Phosphorus** burns in chlorine gas to produce a mixture of two chlorides, phosphorus(III) chloride, PCl₃ and phosphorus(V) chloride, PCl₅. In excess chlorine gas, PCl₅ is the major product.

$$\begin{array}{c} P_4(s) + 6Cl_2(g) \rightarrow 4PCl_3(l) & ; \mbox{if chlorine gas is limited} \\ P_4(s) + 10Cl_2(g) \rightarrow 4PCl_5(s) & ; \mbox{if chlorine gas is in excess} \\ PCl_3 \mbox{is a fuming liquid while } PCl_5 \mbox{is an off-white solid.} \end{array}$$

6) **Sulfur** burns in chlorine gas to produce disulfur dichloride, an orange, evilsmelling liquid.

$$2S(s) + Cl_2(g) \rightarrow S_2Cl_2(l)$$

- 7) **Chlorine** obviously does not react with chlorine gas.
- 8) **Argon** does not react with chlorine gas to form any chlorides.

Reaction with water, H₂O

1) **Sodium** catches fire in cold water and a violently exothermic reaction occurs to form sodium hydroxide and hydrogen gas.

$$2Na(s) + 2H_2O(l) \rightarrow 2NaOH(aq) + H_2(g)$$

2) **Magnesium** reacts very slowly with cold water, taking several days to collect a test tube of hydrogen gas and a weakly alkaline magnesium hydroxide solution.

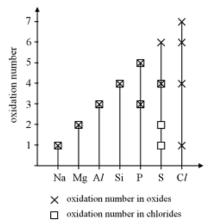
$$Mg(s) + 2H_2O(l) \rightarrow Mg(OH)_2(aq) + H_2(g)$$
; very slow

However, it reacts rapidly with steam to produce magnesium oxide and hydrogen gas.

$$Mg(s) + H_2O(g) \rightarrow MgO(s) + H_2(g)$$
; very fast

Variation in oxidation number of Period 3 oxides and chlorides

- 1) Oxidation number of a Period 3 oxide or chloride corresponds to the **number of electrons used for bonding**. It is always positive because oxygen is more electronegative than any of the element.
- 2) The maximum oxidation number is the **same as Group number**. This corresponds to the total number of valence electrons.
- 3) i. In the oxides, the maximum oxidation number increases from +1 in Na to +6 in S.
 - ii. In the chlorides, the maximum oxidation number increases from +1 in Na to +5 in P.
- 4) Phosphorus and sulfur show several oxidation numbers because they can expand their octet through the excitation of electrons to the empty 3d orbitals.



- 5) For example:
 - i. In SO₂, S has oxidation number +4 because only four electrons are used for bonding.
 - ii. In SO₃, S has oxidation number +6 because all six electrons are used for bonding.

10.3 Period 3 Oxides

Summary of the properties of Period 3 oxides.

	Na ₂ O	MgO	AI_2O_3	SiO ₂	P ₄ O ₆ / P ₄ O ₁₀	SO ₂ /SO ₃	
structure	* Al ₂ O ₃ is io	Ionic nic with covale	ent character.	Giant covalent	Simple molecular		
melting point		high bonds must b nelting to occur		high (strong covalent bonds must be broken)	low (only weak dipole- dipole forces need to be broken)		
acid/base nature	ba	sic	amphoteric	(inert)	acidic		
effect of water on oxide	alkaline solution (pH ≈ 13)	dissolves slightly (pH≈9)	(high lattic	in water ce energies tion difficult)	solı	y acidic ution [≈2)	

Reaction with water, H₂O

1) **Sodium oxide** reacts exothermically with cold water to form sodium hydroxide. A **strongly alkaline** solution of sodium hydroxide is produced. $Na_2O(s) + H_2O(l) \rightarrow 2NaOH(aq)$; pH = 13

2) **Magnesium oxide** reacts **slightly** with water to the extent that it is almost insoluble. A **weakly alkaline** solution of magnesium hydroxide is produced. MgO(s) + H₂O(l) → Mg(OH)₂(aq) ; pH = 9

- 3) **Aluminium oxide** does not react or dissolve in water due to its high lattice energy.
- 4) **Silicon dioxide** does not react or dissolve in water due to the strong covalent bonds.
- 5) Phosphorus oxides react with water to form acidic solutions(pH = 2). Phosphorus(III) oxide reacts with water to form phosphorous acid. P₄O₆(s) + 6H₂O(l) → 4H₃PO₃(aq)
 Phosphorus(V) oxide reacts with water to form phosphoric(V) acid. P₄O₁₀(s) + 6H₂O(l) → 4H₃PO₄(aq)
- 6) Sulfur oxides react with water to form **acidic solutions**(pH = 2). **Sulfur dioxide** reacts with water to give sulfurous acid or sulfuric(IV) acid. $SO_2(g) + H_2O(l) \rightarrow H_2SO_3(aq)$ **Sulfur trioxide** reacts violently with water to form a mist of sulfuric acid. $SO_3(g) + H_2O(l) \rightarrow H_2SO_4(aq)$

Acid-base behaviour of Period 3 oxides

- 1) Going across Period 3, the nature of the oxide changes from **basic(Na₂O, MgO)** to **amphoteric(Al₂O₃)** then to **acidic(SiO₂, P₄O₆/P₄O₁₀, SO₂/SO₃)**. The **acidity of the oxides increases** across the Period.
- 2) **Sodium and magnesium oxides** are **basic oxides**, they react with acid to give the corresponding **salts and water**.

$$Na_2O(s) + 2HCl(aq) \rightarrow 2NaCl(aq) + H_2O(l)$$

 $MgO(s) + 2HCl(aq) \rightarrow MgCl_2(aq) + H_2O(l)$

3) **Aluminium oxide** is **amphoteric**, it can react with **both acid and base**. Aluminium oxide reacts with hot and concentrated acids to give salt and water

$$Al_2O_3(s) + 6HCl(aq) \rightarrow 2AlCl_3(aq) + 3H_2O(l)$$

Aluminium oxide reacts with hot and concentrated sodium hydroxide to give sodium aluminate.

$$Al_2O_3(s) + 2NaOH(aq) + 3H_2O(l) \rightarrow 2NaAl(OH)_4(aq)$$

sodium aluminate

4) **Silicon dioxide** is an **acidic oxide**, it reacts with hot and concentrated sodium hydroxide to give a colourless solution of sodium silicate.

$$SiO_2(s) + 2NaOH(aq) \rightarrow Na_2SiO_3(aq) + H_2O(l)$$

sodium silicate

5) Phosphorus oxides are **acidic oxides**, they react with alkalis to form salts and water.

Phosphorus(III) oxide reacts with sodium hydroxide to form sodium phosphate(III) and water.

$$P_4O_6(s) + 12NaOH(aq) \rightarrow 4Na_3PO_3(aq) + 6H_2O(l)$$

Phosphorus(V) oxide reacts with sodium hydroxide to form sodium phosphate(V) and water.

$$P_4O_{10}(s) + 12NaOH(aq) \rightarrow 4Na_3PO_4(aq) + 6H_2O(l)$$

6) Sulfur oxides are **acidic oxides**, they react with alkalis to form salt and water. **Sulfur dioxide** reacts with sodium hydroxide to form sodium sulfate(IV) and water.

$$SO_2(g) + 2NaOH(aq) \rightarrow Na_2SO_3(aq) + H_2O(l)$$

Sulfur trioxide reacts with sodium hydroxide to form sodium sulfate(VI) and water.

$$SO_3(g) + 2NaOH(aq) \rightarrow Na_2SO_4(aq) + H_2O(l)$$

10.4 Period 3 Chlorides

Summary of the properties of Period 3 chlorides

	NaC <i>l</i>	$MgCl_2$	AICI ₃	$SiCI_4$	PCl ₃ /PCl ₅		
structure	Giant (composed o charged ions he strong electro	f oppositely eld together by	Simple molecular (composed of small discrete molecules held together by weak van der Waals' forces)				
melting point	hig (strong ionic b broken for mel	onds must be	sublimes # (at 180 °C)	low (only weak van der Waals' forces need to be broken) * Exception: PCI ₅ solid			
effect of water on chloride	dissolve (neutral solution)	readily (slightly acidic)	hydrolyse to give fumes of HCl gas; acidic solution				
	(pH≈7)	(pH ≈ 6.5)	(pH ≈ 3)	(pH ≈ 2)			

Reaction with water, H₂O

1) **Sodium chloride** dissolves in water to form a **neutral solution** of sodium chloride.

$$NaCl(s) + aq \rightarrow Na^{+}(aq) + Cl^{-}(aq)$$
; pH = 7

2) **Magnesium chloride** dissolves in water with slight hydrolysis to form a solution of magnesium chloride.

$$\begin{split} MgCl_2(s) + 6H_2O &\rightarrow [Mg(H_2O)_6]^{2+}(aq) + 2Cl^{-}(aq) \\ &\stackrel{hexaaquamagnesium \ ions}{[Mg(H_2O)_6]^{2+}(aq)} + H_2O(l) \rightleftharpoons [Mg(H_2O)_5(OH)]^{+}(aq) + H_3O^{+}(aq) \quad ; \ pH = 6.5 \end{split}$$

3) **Aluminium chloride**, AlCl₃ hydrolyses in water to give an acidic solution, white fumes of hydrogen chloride gas are formed.

$$\begin{array}{l} AlCl_3(s) + 6H_2O(l) \longrightarrow \ [Al(H_2O)_6]^{3\text{+}}(aq) + 3Cl^{\text{-}}(aq) \\ [Al(H_2O)_6]^{3\text{+}}(aq) + H_2O(l) \rightleftharpoons [Al(H_2O)_5(OH)]^{2\text{+}}(aq) + H_3O^{\text{+}}(aq) \end{array} ; pH = 3 \end{array}$$

4) **Silicon tetrachloride** undergoes complete hydrolysis in water to form a strongly acidic solution, white fumes of hydrogen chloride gas are formed.

$$SiCl_4(l) + 2H_2O(l) \rightarrow SiO_2(s) + 4HCl(g)$$
; pH = 2

5) **Phosphorus(III) chloride** reacts violently with water in a hydrolysis reaction to give a strongly acidic solution and fumes of hydrogen chloride gas.

$$PCl_3(l) + 3H_2O(l) \rightarrow H_3PO_3(aq) + 3HCl(g)$$
; $pH = 2$

Phosphorus(V) chloride is an off-white ionic solid at room temperature and sublimes at 163 °C to give phosphorus(III) chloride and chlorine gas.

$$PCl_5(s) \rightleftharpoons PCl_3(l) + Cl_2(g)$$

It reacts violently with water in a hydrolysis reaction to give a strongly acidic solution and fumes of hydrogen chloride gas.

$$PCl_5(s) + 4H_2O(l) \rightarrow H_3PO_4(aq) + 5HCl(g)$$
; pH = 2

6) The **acidity of the chlorides increases** across the Period as the nature of the chlorides changes from ionic to covalent.